Time-resolved porosity changes at cement-clay interfaces derived from neutron imaging

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Abstract

Interfaces between cement and clay are common in engineered barrier systems for the underground disposal of radioactive waste. The geochemical contrast between clay and cement leads to alterations of both materials following dissolution and precipitation reactions. Porosity is a key parameter for, and a key outcome of, such reactive transport. Here we report on the evolution of porosity within small samples made from ordinary portland cement and Na montmorillonite. Water contents, a proxy for porosity in saturated samples, were derived from neutron imaging. Specific calibration procedures were developed to account for multiple scattering effects. An increase in cement porosity was observed propagating ~ 2 mm away from the interface within 600 days, and a considerable decrease in clay porosity within ~ 2 mm. Our experiments provide unique information on the dynamics of the processes up to nearly two years. The data set provides valuable input to better constrain reactive transport models.

Keywords: neutron imaging, cement-clay interaction, radioactive waste (E), porosity, water content, reactive transport, image analysis (B), interfacial transition zone (B), transport properties (C), cement paste (D), portland cement (D), waste management (E)

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1 1. Introduction

Cement and clay are two key components of the barrier systems of radioactive waste repositories. In the multi-barrier design of such repositories, intended to mechanically reinforce open galleries and to enclose waste canisters, large amounts of cementitious material will be used in direct contact with the clay formations or with clays of the engineered barrier systems (EBS). One of the factors affecting the EBS performance is the long-term development of the contact zone between cement and clay. Differences in chemical compositions between these two materials cause diffusive mass flux across their interface, possibly followed by mineral dissolution and precipitation reactions which will alter the composition and structure of both materials.

Several laboratory and field experiments [e.g., 1–11] as well as evidences from 12 natural analogues [e.g., 12] provide information on possible alterations at the 13 material interfaces. For instance, the cement-clay interaction (CI) field experi-14 ment in the Mont Terri rock laboratory was initialized to investigate interactions 15 between Opalinus Clay and various types of concretes and cements [2–4]. Alter-16 ations within clay or cement matrix were investigated with various techniques. 17 The presence of aggregate grains within the concrete made the application of 18 these techniques difficult, but it was possible to delineate various reaction zones 19 based on SEM/EDX element mapping and XRD measurements. Porosity val-20 ues derived from impregnation with a traced resin followed by autoradiography 21 were reported by Jenni et al. [2]. They reported a reduction in porosity in the 22 first $\sim 1 \text{ mm}$ of cement compared to the cement porosity further away, which 23 they related to carbonation. Porosities in both cement regions were, however, 24 larger than expected based on the used water to cement ratio. 25

Autoradiography and petrophysical measurements of Gaboreau et al. [7] on cement-clay interface samples from the Tournemire underground research laboratory (France) after 15 years of interaction showed a clogging of the porosity in the claystone, likely due to precipitation of secondary phases (e.g., C-S-H), and a porosity increase in the cement. They observed heterogeneous zones of the porosity evolution on both sides of the interface, with extents on the order of centimeters (3.5 cm in the cement and 1.5-2 cm in the claystone). In another study at the same location, Techer et al. [8] confirmed a limited spatial extent of the interactions in the clay-rich rock, with the alteration extending over a few centimeters. In an experiment, also at the Tournemire site, Lalan et al. [10, 11] contacted ordinary portlandite heated to 70°C for one year with the argillite. They found a decrease of porosity on both sides of the interface.

Cruz Alonso et al. [5] and Fernández et al. [6] reported on the interaction 38 of concrete with bentonite in the FEBEX experiment at the Grimsel Test Site. 39 After 13 years of interaction, pronounced changes were observed within the 40 first centimeter of a concrete plug in contact with bentonite. The complete 41 disappearance of portlandite in this first centimeter was noted along with a 42 massive ettringite formation, with the sulfate very likely originating from the 43 bentonite. A limited impact on the bulk mineralogy was found on the clay 44 side, with only small changes (some dissolution of primary minerals and some 45 precipitation of secondary minerals) in the first few millimeters, but a somewhat 46 more extended redistribution of exchangeable cations. 47

Various numerical studies were performed to predict mineral alteration path-48 ways when cementitious material interacts with clay. In a study on the inter-49 action of cement with different clay host rocks, Kosakowski and Berner [13] 50 estimated for instance the extents of alteration zones for different transport 51 scenarios in a set of long-term predictive calculations. After ~ 2000 years, the 52 alterations were limited to a comparably narrow zone around the interface (< 553 cm), with the porosity tending to zero in the clay, which stopped a further 54 development in the diffusive transport scenario. In another numerical study, 55 Traber and Mäder [14] obtained an altered zone at an Opalinus Clay-concrete 56 interface with an extent of less than 20 cm. The uncertainties of numerical 57 studies are rather large, among other things because kinetic reaction rates and 58 the feedback of mineral alterations on the further diffusive transport are largely 59 unknown [15, 16]. 60

⁶¹ Even though clogging might eventually be favorable to delay the migration

of radionuclides to the surrounding clay formation, it may have a negative effect 62 on the re-saturation of the clay formation after closure of a repository, reducing 63 its chemical and mechanical buffer capacity as well as the overall functionality 64 of the repository. It could also impact the release of gases at a later stage. De-65 veloping methodologies to provide experimental data on these alterations and 66 their coupled feedback on the repository's transport processes at different time 67 scales is an essential task in order to refine the current assumptions used in pre-68 dictive transport simulations and to increase the reliability of these predictions. 69 Porosity is a central input parameter in reactive transport models, but also a 70 central outcome. Accordingly, knowing its evolution with time is particularly 71 important. 72

Within the scope of the present study, a laboratory experiment has been 73 designed and performed to investigate the above mentioned alterations under 74 well controlled conditions, in a non-destructive way, and over a time period 75 of months to several years. The experimental cell design makes it possible 76 to benefit from a broad range of non-destructive analyses. Here we present 77 the results from the first three years of monitoring of water contents at an 78 ordinary portland cement (OPC) - Na-montmorillonite interface as obtained 79 from neutron imaging. 80

⁸¹ 2. Materials and methods

82 2.1. Sample preparation

The samples of hardened OPC paste, with 5 mm diameter and about 5 mm length, were drilled out of a 14 years old CEM I 52.5 N HTS plug, which was prepared in 2001 [17–20], then stored in a sealed container and, before sample preparation, brought into contact with the artificial cement pore water, as described by Tits et al. [19]. For the clay samples, a conditioned Namontmorillonite powder [21] was used. The powder was pre-saturated within a desiccator that contained humid air, then compacted to the final 5 mm diameter and 5 mm length in a compaction cell. Details on the origin of the material are
given in Shafizadeh et al. [22], including the chemical compositions.

The cement and clay plugs were then brought into direct contact in the 92 diffusion cell designed to host the samples and at the same time to allow in-situ 93 experimental techniques to be applied (Fig. 1). In detail, the samples were first 94 introduced into the inert PTFE holders which was then inserted in the aluminum 95 cell. The PEEK porous filters were added to both sides. The two PEEK end 96 caps (screwed to the aluminum body) kept the samples, the PTFE and the 97 filters in place while providing a connection to an external fluid reservoir. Cell 98 type II was designed after the first measurement campaign with cell type I. The 99 improved cell design led to a better signal to noise ratio and caused less artifacts 100 in neutron radiographs, owing to the thinner aluminum part and the elimination 101 of interfering joints. The cell geometry was also changed to a cylindrical shape 102 for future tomography measurements. Also, the new cell has a smaller outer 103 dimension compared to type I and thus allows for having an open beam section 104 within the image, which lowers the uncertainty in data evaluation. 105

Table 1 summarizes the cement-clay interface samples prepared for this study 106 as well as the properties of both cement and clay materials, prior to their contact. 107 Samples were kept in the diffusion cell while the left and right ends were in 108 contact with the corresponding pore water solutions. A peristaltic pump was 109 used to continuously circulate the solution between sample and reservoir. The 110 samples were kept in a glove box environment to avoid CO_2 contaminations and 111 carbonation of the cement. The corresponding pore water compositions for the 112 OPC and the Na-montmorillonite are reported in Table 2. 113

For the neutron radiography measurements, the cells were disconnected from the pump and the solution reservoir, and openings were closed with dead end screws to ensures the cell tightness.

117 2.2. Neutron imaging measurements

The high interaction probability of neutrons with hydrogenous compounds makes neutron imaging a powerful non-destructive technique to study water dis-



Figure 1: The two designs of the reaction cells used in this study. The components of both reaction cells are (1) Cement sample, (2) clay sample, (3) O-rings, (4) PTFE sample holder, (5) PEEK frits, (6) solution reservoirs within PEEK caps, (7) PEEK caps, (8) aluminum holder and (9) screws. The black square shows the field of view (FOV, 27.5 mm x 27.5 mm) for the neutron images. Outer cell dimensions (without screws): 83 mm x 40 mm x 25 mm (type I), 53 mm x 22 mm Ø (type II)

tribution inside porous materials [24]. At present, it allows measurements with 120 a spatial resolution down to $\sim 10 \ \mu m$ and a temporal resolution of minutes to 121 hours [24-32]. Neutron radiography measurements of the cement-clay interface 122 samples were performed at the ICON beamline (Imaging with Cold Neutrons) 123 at the Paul Scherrer Institut (PSI) [33]. Technical specifications of the measure-124 ments have been given earlier in Shafizadeh et al. [22]. The prepared interface 125 samples were radiographed at different times in order to obtain the evolution of 126 water content profiles across the interface. 127

The neutron beam is collimated from the Swiss spallation neutron source (SINQ, PSI) through vacuum flight tubes towards the sample (collimation ratio 343 m/m). Once it reaches the sample, it can be transmitted through the sample or interact with it, which results in attenuation. This attenuation happens when neutrons are absorbed or scattered by the nuclei of the atoms. The transmitted neutrons hit a Gd-based scintillator (with 20 μ m thickness) with a total field of

	Clay			Cement			
Cell No.	Dry bulk density	Porosity ^a	Length	Dry bulk density	Porosity	Length	Interface contact date
(-)	$(\mathrm{gcm^{-3}})$	$(m^3 m^{-3})$	(mm)	$(\mathrm{gcm^{-3}})$	$(\mathrm{m}^3\mathrm{m}^{-3})$	(mm)	
C1	1.67	0.39	5.5	0.77	0.63	4.5	14.03.2013
C5	1.44	0.49	5.1	0.77	0.63	4.9	25.09.2013

Table 1: Overview on the properties of the interface samples used in this study.

^{*a*} From dry bulk density with grain density of 2.76 g cm⁻³ (C1) and 2.8 g cm⁻³ (C5) Common parameters:

Sample diameter is 5 mm

Cement initial pore water pH is 13.3 [19] and clay initial pore water pH is 8.7

view of 27×27 mm². The scintillator converts the neutron signals into visible 134 light photons. The photons are then guided via a mirror and a lens system 135 to the cooled CCD camera (with 2048×2048 pixels), where they are digitally 136 recorded and displayed as a 16-bit gray value image. The exposure time of 137 90 s was needed to obtain a sufficiently high signal to noise ratio to allow a 138 quantitative analysis of the image data. It is also important in order to reach 139 the highest possible resolution which is reported to be 27 μ m × 27 μ m [33]. 140 In own independent test measurements with the Siemens star periods around 141 50 μ m could be resolved. The obtained gray value images were corrected (flat 142 field, dark noise, dose, geometry, [22] and Appendix C) and then processed to 143 obtain water content profiles across the interface sample. 144

The physical relation between the incident neutron beam I_0 and transmitted neutrons I_T is described by the Beer-Lambert law,

$$\frac{I_T}{I_0} = e^{-\sum_{i=1}^n \mu_i d_i}$$
(1)

where μ_i is the attenuation coefficient of each individual material present in the

	OPC	Initial gradient	Montmorillonite
	$({\rm mol}\ L^{-1})$		$(mol L^{-1})$
Na	0.11	\longleftrightarrow	0.30
К	0.21	\rightarrow	-
Ca	0.023	\rightarrow	1.0×10^{-6}
Mg	1.0×10^{-6}	←	-
$\mathbf{C}^{\mathbf{IV}}$	1.1×10^{-4}	\rightarrow	-
Cl	1.7×10^{-5}	←	0.30
S^{VI}	-	←	0.019
OH^-	0.306	\rightarrow	5.1×10^{-6}
pН	13.3	\rightarrow	8.7
IS	0.3		0.3

Table 2: Aqueous total concentrations, OH⁻ concentrations, pH, and ionic strength (IS) of pore waters in OPC and montmorillonite, calculated by GEMS [23].

sample and d_i the distance that a beam travels through that material. The attenuation coefficient μ of a material is a measure of how strong neutrons interact with the matter. The high attenuation coefficient of hydrogen atoms (present mainly in water), relative to other elements in our samples, makes neutron attenuation most sensitive to the amount of pore water in the cementclay interface samples.

Ideally, the thickness of an individual material (e.g., d_w of water) in a sample composed of several different materials can be calculated from Eq. 1 with known μ_i and thicknesses $d_{i\neq w}$ of all other materials. The contribution of all other materials can be obtained from the image $I_{T,dry}/I_{0,dry}$ of the dry sample, leading to:

$$d_w = -\frac{1}{\mu_w} \left(\ln \left[\frac{I_T}{I_0} / \frac{I_{T,dry}}{I_{0,dry}} \right] \right) .$$
⁽²⁾

The water content θ is then given as $\theta = d_w/d_T$, where d_T is the total thickness of the sample. In reality, however, obtaining an image of the dry sample may not be feasible, and the attenuation of neutrons through water is not fully linear, making calibration not so simple. The details of our calibration procedure are given in the next subsection.

157 2.3. Calibration procedure

As our samples had to remain saturated throughout the whole experiment, 158 the reference image of the dry sample was synthesized from separate image parts 159 with a dried cement and a dried clay [22]. This introduces additional uncertainty 160 especially with respect to single pixel values, but the following calibration can 161 compensate for a possible average bias. To derive the 2D water content field 162 in a radiograph, first the cylindrical geometry (variable d_T) had to be taken 163 into account according to $d_T(y) = 2\sqrt{2yr - y^2}$, where r is the radius of the 164 sample and y the lateral position in the radiograph with the sample extending 165 from y = 0 to y = 2r (Fig. 2). Then, the water attenuation coefficient (μ_w) is 166 needed. This 2D matrix can be obtained in principle from the theoretical values 167 for hydrogen and oxygen at the beam energy peak of the ICON instrument; 168

or, by performing a calibration measurement with pure water. Both of these methods provide a single value for μ_w but result in water contents that are not matching with true water contents of our samples (measured for the cement and calculated from the bulk density for the clay). Therefore, the contribution of multiple scattering on the recorded radiographs as well as other image artefacts need to be corrected for in a more detailed manner.

From calibration measurements with pure water (Fig. A.1a,b in the appendix), we observed that the effective attenuation coefficient μ_w decreased with increasing sample thickness, which coincides here with increasing distance from the lateral boundaries of the sample (Fig. A.1c) due to multiple scattering [34]. Whereas using a single value for μ_w will cause both under- and over-estimation of water contents (Fig. A.1d, the green and red lines) in some regions of the sample, a linear relation

$$\mu_{w,\text{eff}} = \mu_0 + \beta \, d_w \tag{3}$$

(Fig. A.1c, the blue line) will provide correct water contents in the sample (Fig. A.1d, the dashed line). Using such a linear relation in Eq. 2 allows then estimating the water content from μ_0 and β according to

$$\theta = \frac{d_w}{d_T} = -\frac{1}{d_T} \left[\left(\frac{\mu_0}{2\beta} \right) + \sqrt{\left(\frac{\mu_0}{2\beta} \right)^2 + \frac{1}{\beta} \ln \left[\frac{T_{dry}}{T} \right]} \right]$$
(4)

It is clear that the empirical linear relation for μ_w will be only approximately 175 correct for cylindrical samples, even if they have homogeneous water contents 176 within their circular cross section, because effects of multiple scattering depend 177 not only on the local amount (thickness) of water but also on the distance 178 between sample and detector [34], which is a function of y for cylindrical samples. 179 Nevertheless, this approximate correction works well for samples of our size 180 having radially homogeneous water content, as was demonstrated for the pure 181 water sample. Accordingly, we expect it to lead to good results also for our 182 clay-cement samples, where no (or no large) radial gradients in water content 183 are expected. 184

In Shafizadeh et al. [22], it was suggested to use two internal parts of the

sample, denoted as 1 and 2, with known water contents θ_1 and θ_2 (e.g., one in clay and one in cement) to derive the two required calibration parameters μ_0 and β in Eq. 4. By substituting known θ_1 and θ_2 , d_{T_1} and d_{T_2} from sample geometry, and A_1 and A_2 from the neutron radiographs where $A_i = \ln \left[\frac{T_{dry}}{T_i}\right]$, Eq. 4 can be solved for μ_0 and β ,

$$\mu_{0} = \frac{A_{2} \cdot \theta_{1}^{2} \cdot d_{T_{1}}^{2} - A_{1} \cdot \theta_{2}^{2} \cdot d_{T_{2}}^{2}}{d_{T_{1}} \cdot d_{T_{2}} \cdot \theta_{1} \cdot \theta_{2} (d_{T_{1}} \cdot \theta_{1} - d_{T_{2}} \cdot \theta_{2})}$$

$$\beta = \frac{-A_{2} \cdot \theta_{1} d_{T_{1}} + A_{1} \cdot \theta_{2} d_{T_{2}}}{d_{T_{1}} \cdot d_{T_{2}} \cdot \theta_{1} \cdot \theta_{1} (d_{T_{1}} \cdot \theta_{1} - d_{T_{2}} \cdot \theta_{2})}.$$
(5)

It was expected that these parameters, obtained from internal standards, can then be used with Eq. 4 to obtain the water content (θ) values across the whole sample domain.

This approach worked for one given sample, regardless of the detailed positioning (extent, lateral location) of the calibration regions within the unaltered clay or cement. However, this was not the case for most of the other samples. In these cases Eq. 5 could not be solved for μ_0 and β , which shows that a single calibration valid for both cement and clay did not exist. Therefore, μ_0 and β parameters were separately calibrated for cement and clay.

For this purpose, a line in a region of the sample domain far from the interface and with the least alterations was chosen for each material. To improve the statistics, instead of just two-points [22], multiple points were selected (e.g., averaged over 10 × 10 pixels) along the calibration line. The transmissivities $A = \ln \left[\frac{T_{dry}}{T}\right]$ in these calibration regions represent projections of the circular cross sections, obtained for different water thicknesses d_w . Effective water attenuation coefficients $\mu_{w,\text{eff}}(d_w)$ were then calculated using:

$$\mu_{w,\text{eff}}(d_w) = \frac{1}{\theta \, d_T} \ln\left[\frac{T_{dry}}{T}\right] \tag{6}$$

for each calibration region. In this way, one obtains $\mu_{w,\text{eff}}(d_w)$, i.e., the effective scattering coefficients as a function of the total water thickness in each calibration region (Fig. 2). Finally, the μ_0 and β valid for the clay or cement sample domain were estimated by fitting the scattered data linearly. The upper and lower parts of the sample domain with the smallest sample thickness were excluded from the calibration fit, as they showed generally much larger scatter.
That is, only the inner part of the sample, shown by the red rectangle in Fig. 2,
was used for the fit.

As a further problem, it turned out that even though the images were cor-202 rected for the variable average neutron inensity (dose), a calibration was only 203 valid for measurements taken around the calibration time, but not for images 204 taken at another time. This is probably related to variations in the beam in-205 tensity histograms for images taken at different times or days. Such variations, 206 which were documented (Appendix B), may lead to differences in multiple scat-207 tering effects, which then preclude the application of a unique calibration ac-208 cording to our procedure. As a consequence, calibration had to be done for 209 each image. The obtained μ_0 and β served then as the calibration parame-210 ters in Eq. 4 for each measurement time. These parameters account for all the 211 variables affecting the gray values of the image. 212

The calibration procedure allowed finally deriving the 2D water content field 213 from the water transmission image. The 2D water content field was then aver-214 aged in y direction, perpendicular to the cylinder axis, in order to present the 215 results as 1D profiles along the x direction across the material interface. Our 216 samples were always connected to fluid reservoirs on each side, except during 217 the neutron imaging experiments, where the samples were disconnected from 218 the reservoirs but sealed. We thus expect them to be saturated, such that the 219 volumetric water contents represent porosity. 220

221 3. Results and Discussion

Fig. 3 shows the time series of the obtained volumetric water content profiles across the interfaces of two samples (Table 1). As it can be seen from the general trend of the time-series profiles, the porosity (equal to volumetric water content) on the cement side of the interface tended to increase with increasing reaction time, accompanied by a porosity reduction on the clay side of the



Figure 2: Internal calibration procedure: $\mu_{w,\text{eff}}$ values were derived from transmission values A according to Eq. 6 using the known water content (cement: measured; clay: from bulk dry density) and the total sample thickness along the parallel lines shown in the bottom left figure. A linear fit provided μ_0 (intercept) and β (slope) for Equation 4, which was used to derive the water content image.

interface. Fig. 4 shows a closer view of the interface for the same samples
at select reaction times. Cement and clay zones near the interface changed
differently over time, very likely due to dissolution and precipitation reactions.
The results are discussed more in detail in the following.

231 3.1. Amount and extent of porosity changes

The porosity in the cement near the interface increased by ~0.05 m³ m⁻³ at most, whereas in the clay near the interface it was reduced by up to ~0.12 m³ m⁻³ (sample C1) and ~0.14 m³ m⁻³ (sample C5). Although there is a trend in reduction of clay porosity, a residual porosity clearly larger than zero was preserved at the given spatial resolution of about 30–50 μ m. The extent of the zone where porosity was altered is on the order of 2 mm on each side of the interface in sample C1 within 20 months (599 d) of interaction time, and ~1 mm



Figure 3: Evolution of profiles of water content with reaction time (in days) across two interface samples (top: sample C1, bottom: sample C5). The regions blanked out for sample C1 in the top figure are artefacts originating from the joints of the aluminum holder of cell type I; no such artefacts were present anymore with cell type II used for sample C5 in the bottom figure.





Figure 4: Details of the evolution of the water content profiles across sample C1 (top row) and sample C5 (bottom row) for the cement parts next to the interface (left plots), and for the clay parts next to the interface (right plots).

The two examples represent the typical behavior as observed in most sam-240 ples, but in one sample no clear porosity changes were detected, and in another 241 sample porosity changes occurred mostly on one side only. It may be that dif-242 ferent reactions did not occur at different locations in these samples, but in 243 a mixing zone where both precipitation and dissolution happened. Reactive 244 transport simulations have shown that the extent and location of mineral trans-245 formations are very sensitive to various parameters, which may vary locally 246 (microstructural heterogeneities) or from sample to sample in our case. Also, 247 the different patterns could be related to technical limitations of radiographs 248 which flatten and average a possibly heterogeneous 3D structure into a 2D one. 249 This becomes especially crucial when the contact surface of the cement plug is 250

not perfectly flat, e.g., not at an angle of 90° with respect to the cylinder axis. Depending on the rotational position of the sample during the radiograph, a non-perpendicular interface may not be detected and lead to a mixing of clay and cement properties in the image. A tomography would be required to reveal the 3D interior of such a sample, but tomography measurements could not be obtained within the limited amount of beam time.

²⁵⁷ 3.2. Propagation of porosity front in cement and clay

The water content profiles not only reveal information about the amount and the extent of porosity changes, but also about their temporal evolution. Fig. 5 presents the propagation of the porosity fronts in cement and clay. The location of the front is defined as midpoint of the steep increase or decrease in porosity, and the error bars illustrate the approximate width of this front. The front position on the cement side increased steadily with time up to the last observation date after about 600 d. The relation between the position of the porosity front and the reaction time can be approximately described by:

$$x \approx \sqrt{D_a \cdot t} \tag{7}$$

where x is the distance between the front and the interface, D_a is an apparent 258 diffusion coefficient, and t is the reaction time since interface contact. The ap-259 parent diffusion coefficient D_a characterizes the approximate transient behavior 260 of the porosity front, but it cannot be related to a retardation coefficient, as 261 in the case of transport of a linearly sorbing tracer. Cement-clay interactions 262 involve strongly non-linear, time-dependent reactions, so using a constant D_a 263 for the porosity front evolution on the cement side during the ~ 600 d (solid 264 blue line in Fig. 5a) is certainly a strong oversimplification. Detailed reactive 265 transport simulations could give more comprehensive insight into the transient 266 behavior. In any case, if transport parameters (ε, D_p) on both sides of the inter-267 face had varied strongly with time, a change in D_a with time should have been 268 observed. Looking at the front propagation within the cement more in detail, 269 it appears that there are two major steps during the 600 d of this experiment 270

(Fig. 5a - dashed blue line). The porosity front advanced during the first 200 d,
then remained at about 1.5 mm for the next 200 d, before it advanced to 2 mm
and slowed down again. Verifying the validity of this type of behavior needs for
instance reactive transport modeling.



Figure 5: Advancement of the porosity front away from the interface as a function of time in samples C1 and C5. (a) Front position in cement, (b) front position in clay, both estimated as distance of midpoint of steep porosity change to the interface. The model curves in (a) are a strong oversimplification (see text for discussion).

For the clay side (Fig. 5b), the situation is different. The front of reduced 275 276 porosity in the clay did not advance with the same rate as the front of increased porosity in the cement. The position of the front in the clay appears to be more 277 or less stagnant during the observations. Already at the first measurement after 278 two months (64 d), the front was located about 2 mm from the interface. It 279 seems a zone with perturbed porosity had been early created as a result of the 280 interface contact, but this zone did then not further expand significantly with 281 increasing reaction time up to about 600 d, even though reactions within this 282 zone (e.g., porosity reduction) further proceeded. 283

The porosity front on the cement side moved somewhat slower in sample C5 as compared to sample C1, and the distance of the porosity front on the clay side is also somewhat smaller in sample C5 compared to sample C1. Sample

C5 had a lower initial montmorillonite density (and corresponding higher initial 287 montmorillonite porosity), but it is unclear how this could be related to this 288 lower propagation rate. As stated, small differences in local or sample-specific 289 parameters may be responsible for observed quantitative differences between 290 different samples. Reactive transport simulations can help to investigate such 291 subtle sensitivities. In turn, the present data can be very helpful in discriminat-292 ing between different parameterizations of reactive transport models, especially 293 with respect to the implementation of feedback between porosity and diffusion. 294

²⁹⁵ 3.3. Porosity changes as a function of reaction time

The temporal evolution of porosity values at three different locations within the cement and the clay, respectively, was also derived from the neutron imaging data (Fig. 6). The porosity tended to alter more in areas close to the interface compared to the areas far away from it for both cement and clay. The amount of net porosity change is higher for the clay side compared to the cement side.

The porosity trends in sample C1 seem to be more or less monotonical in 301 both, cement (increasing) and clay (decreasing), up to about day 450. Then, the 302 trend appears to be inverted on both sides, with a porosity decrease in cement 303 and a porosity increase in clay, up to about day 512. It remains unclear whether 304 these temporary inversions of the trends are real or represent a measurement 305 artefact (e.g., calibration problem). However, the inversions are strongest on the 306 cement side close to the interface, and weaker further away from the interface 307 and on the clay side, which may indicate that they are indeed related to local 308 transport and reaction processes. If real, such changes of trends suggest that 309 different reactions start to dominate at different times. While dissolution and 310 precipitation reactions may occur in parallel, our observation in terms of the 311 porosity, i.e., the sum parameter, just indicate that on the cement side, disso-312 lution mostly — with the exception around day 500 — outweighs precipitation 313 during the experiment, whereas on the clay side, precipitation mostly outweighs 314 dissolution. 315



Figure 6: Porosity changes as a function of reaction time at different positions in cement (left plots) and clay (right plots) for samples C1 (top row) and C5 (bottom row).

316 3.4. Significance of derived porosity data

As mentioned above in subsection 3.1, not all our experimental results are 317 identical and there is some variability in the observations between different 318 samples, with some samples exhibiting less reactions. The reasons are unknown, 319 but the differences may be related to small parameter variations between the 320 different samples. Anyway, the data shown here represent typical observations. 321 The materials chosen, especially in the case of the clay, are simplified analogs 322 compared to those foreseen in natural or engineered barriers or used in field ex-323 periments, as for instance in the CI experiment in the Mont Terri rock laboratory 324

[2]. This allowed us to focus on the most important interaction processes at the 325 interface and also to provide simpler, better constrained initial conditions for 326 numerical codes. The duration of the reaction between the selected cement and 327 clay materials and the size of the samples (comparably small) were also different 328 from other works on this topic. Although based on small samples, our method 329 presented here has the advantage that it can provide the time and space depen-330 dence of porosity changes, in contrast to all methods that require a destructive 331 sampling. Furthermore, it is possible to create and investigate many different 332 samples with reasonable effort. 333

We are not aware of other studies that provide the porosity evolution at 334 cement-clay interfaces as a function of time and space. In any case, the trends 335 observed during the present study are qualitatively in agreement with obser-336 vations from other in situ and laboratory experiments. For example, there is 337 a similarity between the findings from this study and the works of Read et al. 338 [1], Gaboreau et al. [7], and Bartier et al. [9]. Fig. 7 shows how the porosity 339 determined from petrophysical measurements varied as a function of distance 340 from a cement-argillite contact in the Tournemire URL after 15 years of in-341 teraction [7]. From their measurements, these authors reported "a clogging of 342 the porosity in the clay-rock while the porosity increases in the cement". The 343 porosity evolution was shown to extend to about a centimeter on both sides 344 of the interface, but "heterogeneously distributed in space as a function of the 345 fissure network and interface geometries". 346

The observed changes in porosity are most likely related to mineral alter-347 ations. The porosity increase on the cement side is probably linked to the 348 dissolution of portlandite [5], whereas the nature of the precipitating phase on 349 the clay side is yet unclear at present. Besides precipitation and dissolution 350 reactions, the interaction with cement also triggers modifications of the ex-351 changer population of a clay [e.g., 35, 36], with Na being replaced by K and 352 Ca. Such modifications, which typically propagate more quickly as compared 353 to the front(s) of dissolution or precipitation reactions, can affect the interlayer 354 distances and thus the local distribution of water between different pore en-355



Figure 7: Total connected porosity at a cement-clay interface after 15 months of interaction (modified from [7]).

vironments (e.g., interlayer, interparticle). But for our confined samples, this
redistribution of the pore water is expected to have no or only a weak effect on
the total porosity.

359 4. Conclusions and outlook

In this paper we presented time-series results of water content profiles across 360 several evolving cement-clay interfaces. The water contents, which served as 361 proxy for porosity, were evaluated from neutron radiographs with a new method-362 ology. The present work showed the need to obtain a separate calibration mea-363 surement for the neutron images for each material at every measurement time, 364 especially for investigations spanning long periods of time. Such a need for inter-365 nal standard imposes some limits to the applicability of the method to evolving 366 samples. First, a suited internal calibration region has to be available for each 367 material. Second, mineral changes within each material should remain compa-368 rably small; otherwise, the calibration may no longer be adequate. We used the 369 unaltered cement or clay material, far away from the interface, as internal cali-370

³⁷¹ bration region, and furthermore applied an approximate correction for multiple³⁷² scattering effects.

The finally observed porosity trends appear reasonable both qualitatively 373 and quantitatively. To cross check the validity of our observations, a D_2O dif-374 fusion experiment was subsequently planned and performed on several interface 375 samples. This is the topic of a forthcoming publication, but, without going 376 into details, we expect that the derived porosity changes can be confirmed by 377 the D_2O exchange experiment. Also, a new scattering correction method was 378 developed very recently, which will be applied in future experiments to improve 379 the estimation accuracy [37, 38]. Neutron imaging, which can be combined with 380 complimentary X-ray tomography in the future, offers thus unique possibilities 381 to non-destructively determine porosity changes as a function of time in reacting 382 cement or rock samples. 383

The time-series results revealed a clear evolution of water content or porosity 384 close to the interface. An increase in the porosity on the cement side of the 385 interface accompanied by a reduction of porosity on the clay side was observed. 386 These trends are broadly in agreement with observations from in-situ studies. 387 The porosity increase on the cement side is very likely related to portlandite 388 dissolution [5]. The nature of the precipitating phase(s) on the clay side has 380 still to be studied. The likely modification in the exchanger population of the 390 clay (from Na to K, Ca, [e.g., 35, 36]) is expected to have only a minor effect 391 on the total porosity in our confined samples. 392

The ultimate goal of this work was to provide quantitative porosity data 393 at reacting cement-clay interfaces as a function of time, in order to serve the 394 further development of reactive transport models, especially with respect to 395 the feedback between porosity and diffusion. Porosity is at the same time a 396 key parameter for, and a key outcome of, reactive transport. The obtained 397 porosity data set, together with forthcoming chemical analyses of the reacted 398 zones, can be examined using reactive transport codes. The comparably simple 399 initial mineralogical composition of our system certainly facilitates the further 400 interpretation and modeling of the observations. Based on the comparison of 401

experimental data and modeling results, one may have to adapt existing or introduce new model parameters and feedback mechanisms, all with the aim to eventually better simulate the evolution of cement-clay interfaces in waste repositories. The small-scale experimental observations obtained in this study may thus serve as benchmark data that assist in increasing the confidence in predictions of the long-term stability of cement and clay interfaces.

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578 Appendix A

The transmission values of a cylindrical cell fully filled with water are decreasing with increasing water thickness (Fig. A.1a,b). Scattering effects lead to decrease of the effective attenuation coefficient μ_w with increasing sample thickness (Fig. A.1c,d). This suggests using a linear calibration function for the attenuation coefficient of water in order to derive correct water contents. Such a calibration procedure was used in the present work.

585



Figure A.1: Transmission values and attenuation coefficients obtained for the cylindrical sample cell (cell type II) filled with water. (a) Transmission image, with the yellow line indicating the evaluated cross section of the radial cell. (b) Transmission values along the yellow line. (c) Water attenuation coefficients as a function of true water thicknesses along the yellow line. (d) Comparison of true water thickness (dashed line) along the yellow line, and water thickness calculated by using constant attenuation coefficients μ_w of 3.5 and 3.1 cm⁻¹, or the $\mu_{w,\text{eff}}$ that varies linearly with water thickness according to the calibration fit in (c).

586 Appendix B

Fig. B.1 shows histograms of several open beam intensities recorded over 587 a period of 3 years. The values of 16-bit images range between 0 and 65,535 588 $(2^{16}-1)$, but our open beam images had a smaller range with values between 0 589 and 10,500. The intensities were binned in intervals of 100. The histogram plots 590 show clearly that not only the recorded mean intensity of the beam varies with 591 time, but also the intensity distribution. It is likely that several parameters of 592 the beamline setup, including the incident neutron beam intensity, varied over 593 the 3 year observation time. The variations can affect the amount of neutrons 594 scattered from the sample. The lowest gray values in the histograms indicate a 595 probability of the presence of clearly lower energy neutrons in the poly-energetic 596 beam. Low energy neutrons have a higher scattering probability when passing 597 through the sample. Accordingly, any shifts or distortions in the energy spec-598 trum will affect the mean attenuation coefficient of the water in the sample, as 599 well as the probability of multiple scattering. The partly variable shape of the 600 histograms thus demonstrates that a simple dose window correction procedure is 601 not able to fully account for intensity variations. The most important influence 602 of such variations is their fingerprint on the energy dependent values of the water 603 attenuation coefficient. This highlights the necessity of an experiment-specific 604 time-dependent calibration procedure. 605



Figure B.1: Histograms of the open beam intensities over a period of 3 years as an indication of setup imposed deviations (beam energy, detector efficiency and camera system variations). The histograms present the frequencies of binned gray values. For our setup intensities of the gray values were between 0 and 10,500 (generally between 0 and 65,535 for a 16-bit image). The intensities were binned in intervals of 100, an arbitrary bin-width. The figure shows the counts in each interval for each open beam image.

606 Appendix C



Figure C.1: Image processing sequence applied on the raw data to obtain the water content in the interface samples.