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# Critical issues in the determination of the bentonite cation exchange capacity for the assessment of the macroscopic density of the solid electric charge

By

Dominijanni A.\*, Fratalocchi, E., Guarena N., Manassero M., Mazzieri F.

#### Abstract

The swelling pressure and transport properties of bentonites are controlled by the electric charge density of solid particles, which is commonly estimated from the laboratory measurement of the cation exchange capacity (CEC). However, the standard ammonium displacement method for CEC determination does not take into account the fabric changes that occur in bentonites under exposure to high salt concentration solutions. A series of laboratory tests was conducted to assess the relevance of such a critical issue, by varying the concentration of the extracting KCI solution with respect to that of the standard test. The obtained results show that the release of the adsorbed ammonium cations depends on the bentonite fabric, which is controlled by the KCI concentration. As a consequence, the ammonium displacement method may provide an unrepresentative estimate of the CEC of bentonites. The methylene blue titration method, despite its apparently more limited accuracy, instead seems to provide a more reliable estimation of the CEC, as the bentonite fabric is maintained dispersed during the test.

#### Keywords chosen from the ICE Publishing list

Geosynthetic application, Landfills, Waste management & disposal

#### List of notations

α	empirical coefficient of the fabric boundary surface equation
β	empirical coefficient of the fabric boundary surface equation
ρ <sub>sk</sub>	solid-phase density
bn	average half-distance between the platelets in the tactoid
<b>C</b> 0	reference molar concentration in the fabric boundary surface equation
Cs	salt molar concentration
$\overline{\mathbf{C}}_{sk,0}$	molar concentration per unit solid volume of the solid skeleton electric charge
CEC	cation exchange capacity
$\mathbf{d}_{d}$	dstern/bn
dStern	thickness of the Stern Layer around the external surface of the tactoid
е	total void ratio
em	micro-void ratio
f <sub>Stern</sub>	fraction of cations adsorbed in the Stern layer
Н	height of the bentonite layer in the filtration apparatus of the ammonium displacement
	method
k	hydraulic conductivity of the bentonite layer in the filtration apparatus of the ammonium
	displacement method
<b>N</b> I,AV	average number of lamellae per tactoid
NI,AV0	empirical coefficient of the fabric boundary surface equation

- S total specific surface of bentonite
- usw swelling pressure

#### 1 Introduction

The transport properties and mechanical behaviour of clay soils with a high specific surface, such as bentonites, are governed by the microscopic interactions that occur between the electrically charged solid particles and the ions that are contained in the pore solution (Lambe, 1960; Groenevelt and Bolt, 1969; Sridharan and Rao, 1973; Mitchell, 1991; Moyne and Murad, 2002; Mitchell and Soga, 2005; Guimarães et al., 2013; Musso et al., 2017; Revil, 2017a, 2017b; Delage, 2019).

8 Bentonites can have either a dispersed fabric, in which clay particles are present as well 9 separated montmorillonite units, or an aggregated structure that consists of packets of particles, 10 or tactoids, within which several clay platelets are in a parallel array. The formation of tactoids is 11 determined by a reduction in the electrical repulsive forces among the clay particles, which is 12 mainly induced by an increase in the salt concentration or a decrease in the solvent dielectric 13 constant of the pore solution.

The expected performances of bentonites can be assessed in field applications through physics-based models, which relate the macroscopic constitutive parameters, including hydraulic conductivity, chemico-osmotic efficiency, diffusion coefficient and swelling pressure, to microscopic fabric parameters, such as the average number of montmorillonite lamellae per tactoid, N<sub>I,AV</sub> (Dominijanni and Manassero, 2012; Shackelford et al., 2019).

Manassero (2017) described a physics-based model that was obtained by volume averaging the equations that govern the electric potential distribution, the water flow and the ion transport at the pore scale, and by imposing the condition of macroscopic thermodynamic equilibrium between the pore solution and the external bulk solutions in contact with the bentonite at its boundaries (Dominijanni and Manassero, 2005, 2012; Dominijanni et al., 2006, 2013, 2018).

On the basis of such a theoretical approach, the microscopic (pore scale) properties of bentonites are taken into account through a single fundamental parameter, that is the molar concentration per unit solid volume of the solid skeleton electric charge,  $\bar{c}_{sk,0}$ . This parameter is related to the cation exchange capacity, CEC, as follows (Dominijanni and Manassero, 2012):

28

$$29 \qquad \overline{c}_{sk,0} = \frac{1 - f_{Stern}}{N_{I,AV}} \cdot CEC \cdot \rho_{sk}$$
(1)

31 where  $\rho_{sk}$  is the solid-phase density ( $\approx 2700 \text{ kg/m}^3$ ) and  $f_{Stern}$  is the fraction of cations 32 immobilized in the so-called Stern layer ( $\approx 0.75-0.95$ ).

Although CEC determination alone is not sufficient to allow an evaluation of  $\bar{c}_{sk,0}$ , it plays a fundamental role in relating the physical and chemical properties of bentonite at the pore scale to the macroscopic constitutive parameters. For this reason, a series of experimental tests has been conducted to assess the reliability of the most commonly used CEC measurement methods, including the ammonium displacement method and the methylene blue titration (MB) method.

39

#### 40 2. Materials and methods

41

#### 42 2.1 Bentonite

The powdered bentonite used in this study comes from the same lot and was subjected to the same cyclic-squeezing procedure as in Puma et al. (2015) to remove soluble salts prior to further testing. The main properties of the bentonite are listed in Table 1.

46

#### 47 **2.2** Testing procedures

As far as the ammonium displacement method is concerned, the CEC was determined according to the ASTM D7503-18 procedure, except for the KCl solutions that were used in the final extraction step: in addition to the 1 M KCl solution (standard procedure), extracting solutions with different KCl concentrations (4.5 M, 0.1 M, 0.05 M, 0.025 M, 0.01 M, 0.0025 M, 0.001 M) were used together with distilled water (DW) to investigate the effect of changes in the bentonite fabric on the CEC results. The tests were generally performed in duplicate for each extracting solution.

The MB test was carried out following the procedure outlined in EUBA (2002). The method is a rapid qualitative procedure that is used in industry for routine quality controls, which provides a measure of the accessible anionic sites in a condition of enhanced dispersion of the clay. This dispersion is obtained by bringing the bentonite suspension to the boil and subsequently titratingthe suspension with an anionic dye (methylene blue).

60

#### 61 3. Test results

62 The CEC measurement results are listed in Table 2 and plotted in Figure 1. The solid symbols 63 in Figure 1 represent the CEC values obtained from the ammonium release measurements. The 64 values obtained using DW are plotted at 0.0001 M KCI concentration. Considering the average 65 of the measurements for each extracting solution concentration, the maximum value of CEC 66 measured with ammonium was 75.3 meq/100g, which was obtained for the 0.1 M KCI solution 67 (the value of 103 meg/100g for the third replicate at 0.1 M KCI was considered to be an outlier 68 of the dataset and, therefore, neglected). As a result, the maximum CEC value obtained with 69 ammonium was lower than the values obtained with the MB tests (97 and 104 meg/100g; for the 70 sake of simplicity the mean value is plotted in Figure 1). The results suggest that the measured 71 CEC depends on the aggregation state of the bentonite particles. The MB test seems to provide 72 an upper bound of the measured CEC due to the disperse state of the bentonite, which 73 enhances the accessibility of exchange sites with respect to the state of aggregation obtained in 74 the standard test. Despite some scatter in the results, the average CEC versus KCI 75 concentration tends to be practically constant for KCI > 0.1 M, thus suggesting a similar 76 aggregation state of bentonite particles. As the KCI concentration decreased (< 0.1 M), the CEC 77 calculated from the released ammonium also decreased. It should be considered that the 78 amount of K cations available for exchange also decreased. The maximum theoretically 79 measurable CEC, as calculated from the available potassium, is also shown in Figure 1. The 80 measured CEC values for low (< 0.02 M) to practically zero (DW) KCI concentrations are in the 81 13.9 - 23.9 meq/100g range, that is higher than the theoretically predicted values based on the 82 available potassium. A blank test (performed without soil and using a 1 M KCl solution) showed 83 that the ammonium residue in the apparatus at most accounts for 1.3 meg/100g. Therefore, the 84 measured CEC suggests a release of ammonium from the clay that is not related to the 85 adsorption of potassium. According to the test method, washing the clay with isopropanol 86 should remove the excess unbound ammonium acetate; however, the test results suggest that 87 some residual ammonium remains in the clay. A replicate test was performed using 0.0025 M

88 KCI solution and a double wash with isopropanol (240 ml instead of 120 ml), but no significant 89 difference in the measured CEC was observed (Table 2). Therefore, the test results suggest 90 that, even after washing with isopropanol, some unbound ammonium remains entrapped in the 91 bentonite layer. Filtration with low concentration (< 0.02 M KCI) solutions induces a change in 92 bentonite fabric during the test, namely swelling and reorganisation, whereby the release of 93 ammonium is favoured. This is reflected, at the macroscopic scale, by an increase in the 94 average height of the bentonite film that forms in the filtration apparatus, H, at the end of the 95 tests, and a corresponding decrease in the hydraulic conductivity, k. The k value was estimated 96 from the observed flow rate of the filtrate during the addition of the final 50 ml of the extracting 97 solution and from the hydraulic gradient that was established across the bentonite film during 98 filtration. H was equal to 2.5 mm and the average k was estimated to be about 10<sup>-7</sup> m/s with the 99 1 M KCI solution (standard procedure), whereas H was 8 mm and k was about 10<sup>-9</sup> m/s with 100 DW.

101

#### 102 4. Discussion

The observed changes in the hydraulic conductivity of bentonite suggest that the CEC variation, as a function of the extracting KCI solution concentration, can be related to fabric modifications. Manassero et al. (2016, 2018), Dominijanni et al. (2017) and Manassero (2017) modelled bentonite fabric modifications through a fabric boundary surface (FBS), whereby the average number of lamellae per tactoid, N<sub>I,AV</sub>, is related to the salt concentration, c<sub>s</sub>, and the micro-void ratio, e<sub>m</sub>, which in turn is obtained by subtracting the void space between the platelets of the tactoids from the total void space.

110 A first phenomenological formulation of FBS was proposed by Manassero et al. (2016):

111

112 
$$\mathbf{N}_{\mathrm{I,AV}} = \mathbf{N}_{\mathrm{I,AV0}} + \frac{\alpha}{\mathbf{e}_{\mathrm{m}}} \cdot \left(\frac{\mathbf{c}_{\mathrm{s}}}{\mathbf{c}_{\mathrm{0}}} + 1\right) + \beta \cdot \mathbf{e}_{\mathrm{m}} \cdot \left[1 - \exp\left(-\frac{\mathbf{c}_{\mathrm{s}}}{\mathbf{c}_{\mathrm{0}}}\right)\right]$$
 (2)

113

114 where  $c_0$  represents the reference molar concentration (= 1 M) and N<sub>I,AV0</sub>,  $\alpha$  and  $\beta$  are non-115 dimensional empirical parameters. 116 The micro-void ratio, e<sub>m</sub>, in Eq. 2 can be derived from the total void ratio, e, through the 117 following equation:

119 
$$e_{m} = \frac{e \cdot N_{l,AV} - S \cdot \rho_{sk} \cdot b_{n} \left( N_{l,AV} + d_{d} - 1 \right)}{N_{l,AV}}$$
(3)

120

121 where  $b_n$  is the average half-distance between the platelets in the tactoid ( $\approx 0.4$  nm), S is the 122 total specific surface ( $\approx$  700 m<sup>2</sup>/g) and d<sub>d</sub> is the thickness of the Stern Layer divided by b<sub>n</sub> ( $\approx$  4). 123 Inserting Eq. 3 into Eq. 2 the number of lamellae per tactoid is related to the total void ratio and

124 the salt concentration through a cubic equation, which can be solved analytically or numerically 125

126 Although a sufficient number of experimental data is not available for the tested bentonite to 127 determine the FBS parameters, a qualitative analysis was conducted by using the calibration

for given values of the parameters N<sub>I,AV0</sub>,  $\alpha$ ,  $\beta$ , S,  $\rho_{sk}$ ,  $b_n$  and  $d_d$ .

128 performed by Manassero (2017) on the hydraulic conductivity experimental results obtained by 129 Petrov and Rowe (1997) on a needle-punched geosynthetic clay liner (GCL), which provided 130 N<sub>LAV0</sub> = 1.56,  $\alpha$  = 8.82,  $\beta$  = 10.01. A plot of the corresponding FBS in the space of the variables 131 NLAV, em and cs is shown in Figure 2. Such an FBS can be regarded as suitable for the analysis 132 of the obtained laboratory data, as the features of the bentonite in the GCL are similar to the 133 ones of the tested bentonite (Table 1) and the GCL hydraulic conductivity is not expected to be 134 influenced by the presence of needle-punched fibres, at least in the range of low salt 135 concentrations (< 0.1 M KCl solution) (Puma et al., 2015).

136 The ability of the proposed FBS to accurately model microstructural changes was verified using 137 a series of experimental results from the literature that included direct measurements of NI,AV, on 138 bentonites with similar properties to the ones of the bentonite tested in the present study (Table 139 1). The values of  $N_{LAV}$ , which were estimated directly from the ratio of the intra-tactoid to the 140 inter-tactoid pore-space by means of Small Angle X-Ray Scattering Spectroscopy (Muurinen et 141 al., 2013) and Nuclear Magnetic Resonance techniques (Muurinen et al., 2013; Ohkubo et al., 142 2016), are shown to be in good agreement with the FBS predictions in Figure 3, for a salt 143 concentration  $c_s = 0.1$  M.

The total void ratio was estimated, during the washing phase with isopropanol, from the measurement of the bentonite layer thickness to be about 4 and, as a result,  $N_{I,AV}$  is calculated through the FBS equation to be about 4.74, assuming  $c_s = 0$  as a consequence of the complete removal of the pore aqueous solution.

148 The relation between N<sub>I,AV</sub> and the KCI concentration, c<sub>s</sub>, provided by the FBS for the values of 149 the bentonite void ratio that were estimated at the end of the ammonium displacement tests 150 (Table 2) is shown in Figure 4. The value of the KCI concentration corresponding to  $N_{I,AV} = 4.74$ 151 is equal to about 0.027 M. As a result, the bentonite is expected to swell during the KCI solution 152 filtration phase and assume a more dispersed fabric when the KCI concentration is lower than 153 0.027 M, while it is expected to flocculate and assume a more aggregated fabric when the KCI 154 concentration is higher than 0.027 M. Such a theoretical threshold value of the KCI 155 concentration is very close to the experimentally found value of 0.02 M, below which the 156 released ammonium overcomes the available potassium.

157 This qualitative result suggests an interpretation of the CEC data that were obtained from the 158 ammonium release measurements. After the washing phase with isopropanol, a portion of 159 mobile ammonium was not removed because of the presence of pores that are less accessible 160 to the advection flux. When the KCI concentration of the extracting solution was lower than 161 about 0.02 M, the dispersion of the bentonite allowed the mobile ammonium ions to be released 162 by opening such pores. Instead, when the KCI concentration of the extracting solution was 163 higher than about 0.02 M, the bentonite flocculated and created additional less accessible pore 164 voids. The potassium cations only had access to a limited portion of the available pores for the 165 highest values of the KCI concentration (≥ 1 M), and, as a result, the exchanged ammonium ion 166 measurements underestimated the effective bentonite CEC.

167

#### 168 **5. Conclusions**

The tests performed by varying the KCI solution concentrations showed that the CEC measurements based on ammonium release are influenced by bentonite fabric modifications and may provide an unreliable estimation of the effective density of the exchangeable sites of the bentonite. For this reason, the ammonium displacement method does not seem to be

173 sufficiently accurate to assess the fundamental fabric parameters of coupled hydro-chemo-174 mechanical models, such as the one proposed by Manassero et al. (2016). The experimental 175 swelling pressure data obtained by Dominijanni et al. (2013) for a saturated sodium bentonite at 176 a void ratio of 4.26 are compared in Figure 5, by way of example, with the theoretical predictions 177 that are obtained by determining  $\overline{c}_{_{sk,0}}$  from the measurement of CEC through Eq. 1, with a 178 constant  $N_{I,AV}$  value ( $N_{I,AV} = 3$ ) for the investigated range of low salt concentrations (Manassero, 179 2017). The theoretical curve derived from the average CEC provided by the standard 180 ammonium displacement method underestimates the experimental data to a great extent, 181 whereas an acceptable fitting is obtained when the average CEC value provided by the 182 methylene blue titration method is used. As a result, the methylene blue titration method, which 183 is used in industry for routine quality controls, seems to be able to provide a more reliable 184 estimation of CEC, despite its apparently more limited accuracy, as the bentonite fabric is 185 maintained dispersed during the test.

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- 264
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266	Table captions
267	
268	Table 1. Main properties of the bentonite used in this study and comparison with the data of
269	similar bentonites from the literature (Petrov and Rowe, 1997; Muurinen et al., 2013; Ohkubo et
270	al., 2016) (DW = distilled water).
271	
272	Table 2. CEC values obtained in the study.
273	
274	
275	Figure captions
276	
277	Figure 1. CEC values determined with the ammonium acetate method using different KCI
278	solutions in the final stage of the test and comparison with MB results.
279	
280	Figure 2. Plot of the fabric boundary surface (FBS) in the three-dimensional space of the
281	variables: average number of lamellae per tactoid, $N_{I,AV}$ , micro-void ratio, $e_m$ , and salt molar
282	concentration, c <sub>s</sub> .
283	
284	Figure 3. Comparison between the average number of lamellae per tactoid provided by the
285	Fabric Boundary Surface (continuous line) and the experimental results taken from the
286	literature.
287	
288	Figure 4. Average number of lamellae per tactoid of the bentonite as a function of the
289	concentration of the extracting KCI solution. The arrows indicate the KCI concentration ( $c_s$ =
290	0.027 M) that corresponds to $N_{I,AV} = 4.74$ , i.e. the average number of lamellae per tactoid after
291	the washing phase with isopropanol ( $c_s = 0$ ).
292	
293	Figure 5. Comparison between the swelling pressure of bentonite, as theoretically predicted on
294	the basis of the average CEC value derived from the methylene blue titration method [curve (a)
295	- CEC = (97.4 + 104)/2 = 100.7 meq/100g] and the standard ammonium displacement method
296	[curve (b) - CEC = $(73.2 + 60.7)/2 = 66.9 \text{ meq}/100g$ ], and the experimental data obtained by
297	Dominijanni et al. (2013) (closed circles).

1 Table 1. Main properties of the bentonite used in this study and comparison with the data of

2 similar bentonites from the literature (Petrov and Rowe, 1997; Muurinen et al., 2013; Ohkubo et

3 al., 2016) (DW = distilled water).

Property	This Study	Petrov and Rowe (1997)	Muurinen et al. (2013)	Ohkubo et al. (2016)
Smectite content (%)	> 98	91	84	> 98
Prevalent adsorbed cation	Na⁺	Na⁺	Na <sup>+</sup>	Na⁺
Liquid Limit to DW (%)	525	530	-	-
CEC (meq/100g)	97 - 104 <sup>(a)</sup>	85.8 <sup>(b)</sup>	80 - 88 <sup>(c)</sup>	-
Hydraulic conductivity to DW (m/s)	8.0×10 <sup>-12 (d)</sup>	1.2×10 <sup>-11 (e)</sup>	5.0×10 <sup>-12 (f)</sup>	-

4 <sup>(a)</sup> measured through the methylene blue titration method

5 <sup>(b)</sup> measured through the Ag-Thiourea exchange for Na<sup>+</sup> and K<sup>+</sup>, the KCI exchange for Mg<sup>2+</sup> and

6 Ca<sup>2+</sup>

7 <sup>(c)</sup> measured through the Cu(II)-Triethylentetramine exchange method

8 <sup>(d)</sup> measured at a 27.5 kPa confining effective stress

9 <sup>(e)</sup> measured at a 35 kPa confining effective stress

10 <sup>(f)</sup> measured at a bulk dry density equal to 517 kg/m<sup>3</sup>

11

		Test 1 (meq/100 g)	Test 2 (meq/100 g)	Test 3 (meq/100 g)	Note
ASTM D7035 (standard and modified procedure)	Estimated bentonite void ratio, e <sup>(*)</sup>	( - 1 3/	/	(	
ĎW	14	17.5	20.4	13.9	
0.001 M	12	18.9	-	-	
0.0025 M	12	21.4	17.7	-	
0.0025 M	12	21.5	-	-	Double wash with isopropano
0.01 M	10	23.9	18.4	-	
0.025 M	8	42.0	45.4	-	
0.05 M	7	60.0	63.4	-	
0.1 M	6	76.2	74.4	103	
1 M (standard)	4	73.2	60.7	-	
1 M (standard)	-	1.3	-	-	Blank test (no soil)
4.5 M	3	63.0	76.3	-	
<u>Methylene Blue</u> (EUBA)					
	-	97.4	104	-	

13 Table 2. CEC values obtained in this study.

14 (\*) total void ratio estimated from the detection of the bentonite layer thickness within the filtration

15 apparatus at the end of the ammonium displacement tests

16

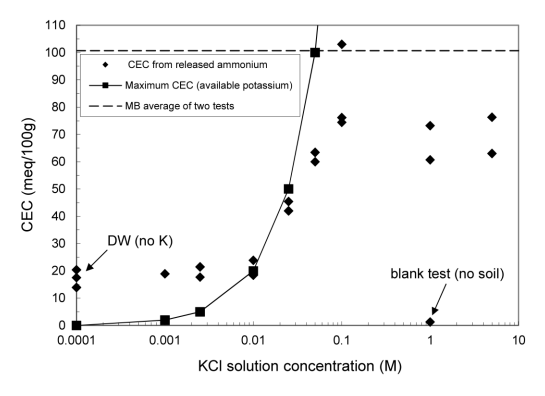


Figure 1. CEC values determined with the ammonium acetate method using different KCIsolutions in the final stage of the test and comparison with MB results.

- ~

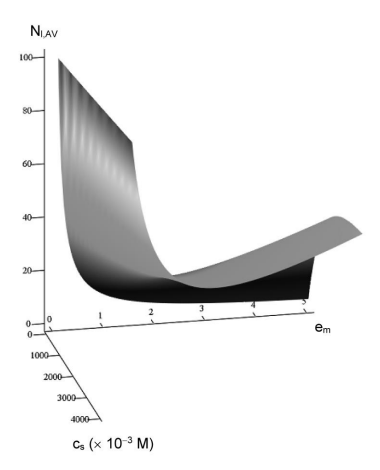
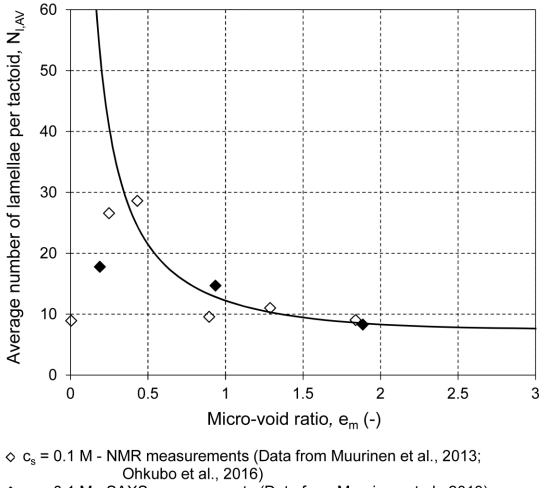


Figure 2. Plot of the fabric boundary surface (FBS) in the three-dimensional space of the
variables: average number of lamellae per tactoid, N<sub>I,AV</sub>, micro-void ratio, e<sub>m</sub>, and salt molar

- 13 concentration,  $c_s$ .



• c<sub>s</sub> = 0.1 M - SAXS measurements (Data from Muurinen et al., 2013)

Figure 3. Comparison between the average number of lamellae per tactoid provided by the
Fabric Boundary Surface (continuous line) and the experimental results taken from the
literature.

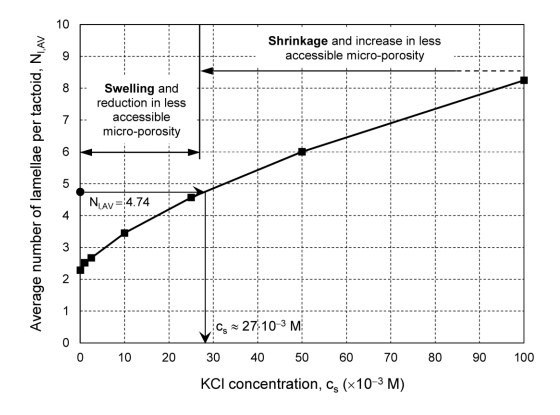


Figure 4. Average number of lamellae per tactoid of the bentonite as a function of the concentration of the extracting KCI solution. The arrows indicate the KCI concentration ( $c_s = 0.027$  M) that corresponds to N<sub>I,AV</sub> = 4.74, i.e. the average number of lamellae per tactoid after the washing phase with isopropanol ( $c_s = 0$ ).

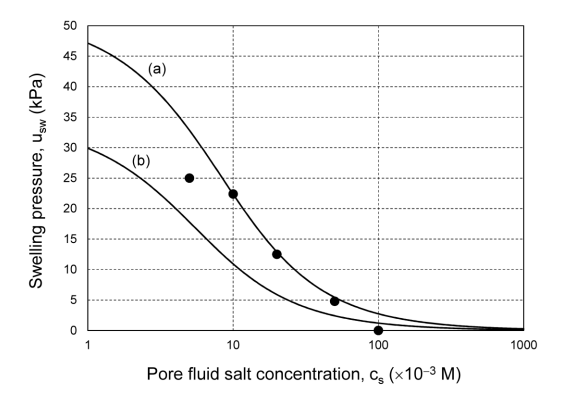




Figure 5. Comparison between the swelling pressure of bentonite, as theoretically predicted on the basis of the average CEC value derived from the methylene blue titration method [curve (a) - CEC = (97.4 + 104)/2 = 100.7 meq/100 g] and the standard ammonium displacement method [curve (b) - CEC = (73.2 + 60.7)/2 = 66.9 meq/100 g], and the experimental data obtained by Dominijanni et al. (2013) (closed circles).