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Coupled chemical-hydraulic-mechanical behaviour of bentonites

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Bentonites are clay soils characterised by a high specific surface and a permanent negative electric charge on their solid skeleton. Their common use as hydraulic and contaminant barriers for landfill and soil remediation applications, including the final disposal of nuclear waste, needs to be supported by adequate theoretical modelling of their mechanical behaviour and transport properties, in order to assess the expected performance in the long term. To this end, a theoretical approach has been proposed in order to derive constitutive equations for their coupled chemical-hydraulic-mechanical behaviour. The phenomenological parameters that govern the transport of electrolyte solutions through bentonites – that is, the hydraulic conductivity, the reflection coefficient (which is also called the chemico-osmotic efficiency coefficient), and the osmotic effective diffusion coefficient – have been measured through laboratory tests on a bentonite with porosity of 0.81, over a range of sodium chloride concentrations in the pore solution that varied from 5 mM to 100 mM. The relevance of the osmotic phenomena has been shown to decrease when the salt concentration increases. The obtained results have been interpreted by assuming that the microscopic deviations of the pore solution state variables from their average values are negligible. In this way, it is possible to interpret the macroscopic behaviour on the basis of the physical and chemical properties of the bentonite mineralogical components.

KEYWORDS: chemical properties; clays; constitutive relations; expansive soils; laboratory tests; pore pressures

INTRODUCTION

The term ‘bentonite’ is commonly used to indicate a clay soil with a high content (>70%) of montmorillonite, a mineral of the smectite group. Montmorillonite particles are thin lamellae that are characterised by a high specific surface (defined as the surface per unit weight) and a permanent negative electric charge. Bentonite is used in hydraulic and contaminant barriers, because of its low hydraulic conductivity, k, to permeation with water and dilute aqueous solutions (k typically ≈ 3 × 10⁻¹¹ m/s). Geosynthetic clay liners (GCLs), which consist of a thin layer of bentonite (~5–10 mm thick) sandwiched between two geotextiles, are examples of such barriers. GCLs are currently used in bottom and cover landfill barriers to limit water infiltration and contaminant migration.

Owing to its physical and chemical properties, bentonite’s macroscopic mechanical behaviour and transport properties cannot be modelled through classical soil mechanics approaches (Manassero & Dominijanni, 2003). In fact, bentonite swells or shrinks in response to changes in the chemical composition of the pore solution. Moreover, when a hydraulic pressure gradient, or a solute concentration gradient, is applied to it, both a volumetric flux and a diffusive solute mass flux occur, as in semi-permeable membranes. For this reason, a theoretical approach that accounts for the electric interaction that occurs between the solid skeleton of the bentonite and the ions contained in the pore solution is proposed in the first part of this paper. The experimental determination of the swelling pressure and the transport properties of a natural sodium bentonite is described in the second part. The obtained results have been interpreted with the proposed theoretical model, in which the macroscopic properties of the bentonite have been linked to the observed macroscopic behaviour.

THEORY

Montmorillonite lamellae are characterised by a negative electric charge, owing to the isomorphic substitution of a portion of the tetravalent silicon (Si⁴⁺) and the trivalent aluminium (Al³⁺) in their crystalline structure, with metals, such as magnesium (Mg²⁺), which have a lower valence. This electric charge per unit solid volume can be expressed as \( F \cdot e_{\text{tot}} \), where \( F \) is Faraday’s constant (96 485 C/mol) and \( e_{\text{tot}} \) is the molar concentration per unit solid volume of the solid skeleton electric charge, which is assumed to have unit valence (i.e. \( e_{\text{sk}} = -1 \)). \( e_{\text{tot}} \) represents the moles of solid skeleton electric charge per volume of solids and, in order to be compared with the ion concentrations of the pore solution, it needs to be divided by the void ratio, \( e \), which represents the pore volume per volume of solids. Dominijanni & Manassero (2012b) have shown that \( e_{\text{sk}} \) is proportional to the effective specific surface of the solid particles, and decreases when the montmorillonite lamellae aggregate to form the so-called tactoids.

If the pore solution contains a single salt that is completely dissociated with the stoichiometric reaction

\[
(\text{Cation})_{z_1}^{z_1}(\text{Anion})_{z_2}^{z_2} \rightarrow v_1(\text{Cation})^{z_1} + v_2(\text{Anion})^{z_2}
\]

where \( z_1 \) and \( z_2 \) are the electrochemical valences of the cation and the anion, and \( v_1 \) and \( v_2 \) are the stoichiometric coefficients of the cation and the anion respectively, then the following condition has to be satisfied in order to preserve electroneutrality within a saturated porous medium, even in the presence of the solid skeleton electric charge

\[
z_1 \xi_1 + z_2 \xi_2 = \frac{e_{\text{sk},0}}{e}
\]

(1)
where \( c_1 \) and \( c_2 \) are the molar concentrations of the cation and the anion respectively, and \( e \) is the void ratio.

As a consequence, the solid skeleton electric charge influences the distribution of the ions contained in the pore solution. This phenomenon is known as the ion partition effect, and is expected to be more relevant for porous media characterised by higher solid skeleton charge concentrations.

**Equilibrium conditions**

When an electrically charged porous medium is placed in contact with an external bulk solution that contains the same ions as are present in the pore solution, a thermodynamic equilibrium condition is reached, after a certain period of time, in which the water chemical potential and the ion electrochemical potentials between the two solutions are equal. The external bulk solution can be considered as a ‘chemical thermometer’ in order to evaluate the equilibrium conditions of the porous medium (Coussy, 2004). The electroneutrality condition in the external solution is given by

\[
z_1 c_1 + z_2 c_2 = 0
\]

where \( c_1 \) and \( c_2 \) are the molar concentrations of the cation and the anion that are contained in the bulk solution.

It is convenient to define the salt concentration, \( c_s \), of the external solution as

\[
c_s = \frac{c_1}{\nu_1} + \frac{c_2}{\nu_2}
\]

Using equation (3), the electroneutrality condition, equation (2), provides the relation between the electrochemical valences and the stoichiometric coefficients as

\[
z_1 \nu_1 + z_2 \nu_2 = 0
\]

As a result, the equilibrium condition can be characterised by the following state variables of the external bulk solution: the absolute temperature, \( T \); the hydraulic pressure (referenced to the atmospheric pressure, as is usual in soil mechanics), \( u \); and the salt concentration, \( c_s \). The corresponding variables of the pore solution can be evaluated from the following conditions

\[
T = T
\]

\[
\mu_w = \mu_w
\]

\[
\mu^{\text{ec}}_i = \mu^{\text{ec}}_i \quad i = 1, 2
\]

where \( T \) and \( T \) are the absolute temperature in the pore solution and in the external bulk solution respectively; \( \mu_w \) and \( \mu_w \) are the water chemical potential in the pore solution and in the external bulk solution respectively; and \( \mu^{\text{ec}}_i \) and \( \mu^{\text{ec}}_i \) are the electrochemical potentials of the \( i \)th ion in the pore solution and in the external bulk solution respectively.

The water chemical potential, \( \mu_w \), and the ion electrochemical potentials, \( \mu^{\text{ec}}_i \), of the external solution can be related to the hydraulic pressure, \( u \), and the salt concentration, \( c_s \), for a dilute solution according to (Katchalsky & Curran, 1965; Dominijanni & Manassero, 2012a)

\[
\mu_w = \mu_w(T) + \frac{u - \Pi}{c_w}
\]

\[
\mu^{\text{ec}}_i = \mu_i + z_i F \phi = \mu_i(T) + RT \ln \left( \frac{c_i}{c_s} \right) + z_i F \phi \quad i = 1, 2
\]

where \( \mu^{\text{ec}}_i \) and \( \mu^{\text{ec}}_i \) are integration constants that only depend on the absolute temperature \( T \); \( c_w \) is the water molar concentration; \( \Pi = RT \sum_{i=1}^{\infty} \nu_i c_i \) is the osmotic pressure; \( \mu_i \) is the chemical potential of the \( i \)th ion; \( R \) is the universal gas constant \((= 8.314 \text{ J/(mol K)})\); and \( \phi \) is the electric potential.

The state variables in the external bulk solution can be measured easily, whereas it is very difficult to determine the corresponding variables in the pore solution. Moreover, the relations obtained by linking the chemical potentials to the state variables of the pore solution are more uncertain, owing to the interaction with the solid skeleton charge, which alters the ion concentration distribution near the solid particles. The simplest assumption that can be adopted involves using relations analogous to equations (8) and (9), as they are also considered valid for the pore solution. This assumption, which was first proposed by Donnan (1911), neglects the microscopic deviations of the ion concentrations from their average values that are induced by the electric potential distribution within the pores. If this approximation is accepted, then the water chemical potential, \( \mu_w \), and the ion electrochemical potentials, \( \mu^{\text{ec}}_i \), of the pore solution can be expressed as

\[
\mu_w = \mu^{\text{ec}}_w(T) + \frac{\hat{u} - \Pi}{c_w}
\]

\[
\mu^{\text{ec}}_i = \mu_i + z_i F \hat{\phi} = \mu_i^{\text{ec}}(T) + RT \ln \left( \frac{c_i}{c_s} \right) + z_i F \hat{\phi} \quad i = 1, 2
\]

where \( \mu^{\text{ec}}_w \) and \( \mu^{\text{ec}}_w \) are integration constants that depend only on the absolute temperature, \( T \); \( c_w \) is the molar concentration of the water in the pore solution, which can be taken equal to the molar concentration of the water in the external bulk solution, that is, \( c_w \cong c_w \); \( \Pi = RT \sum_{i=1}^{\infty} \nu_i c_i \) is the osmotic pressure of the pore solution; \( \mu_i \) is the chemical potential of the \( i \)th ion in the pore solution; and \( \phi \) is the electric potential in the pore solution.

The hydraulic pressure of the pore solution, \( \hat{u} \), and the ion partition factor, \( \Gamma_i \), defined as the ratio between the ion concentration of the pore solution and the ion concentration of the external bulk solution, can therefore be expressed on the basis of equations (6) and (7), and using equations (8)–(11)

\[
\hat{u} = u + (\Pi - \Pi)
\]

\[
\Gamma_i = \frac{c_i}{c_i} - \frac{c_i}{c_i} = \exp \left( -\frac{z_i F \psi}{RT} \right) \quad i = 1, 2
\]

where \( \psi = \hat{\phi} - \phi \) is the electric potential of the porous medium, which is also called Donnan’s potential.

On the basis of this approach, the hydraulic pressure of the pore solution is different from the hydraulic pressure of the external solution that is in equilibrium with it. The pressure difference between the pore solution and the external solution is called the swelling pressure, \( u_w \), and is given by

\[
u_w = \Pi - \Pi
\]

Equations (12) and (13) for \( i = 1, 2 \), together with equation (1), constitute a set of four equations that can be solved to find the four unknown variables: the hydraulic pressure, \( u \), the ion concentrations, \( c_i \) for \( i = 1, 2 \), and the electric potential, \( \psi \).

When the ion electrochemical valences are both unitary, such as for sodium chloride, equation (13) implies that

\[
\Gamma_i = \Gamma_i^{-1}
\]

Inserting equation (15) into equation (1) results in the equation

\[
\Gamma_i^{-1} - \Gamma_i - \frac{c_{\text{sk},0}}{c_{es}} = 0
\]

which has a positive solution of the form
\[
\Gamma_2 = -\frac{\alpha_{k,0}}{2\varepsilon_0} + \sqrt{\left(\frac{\alpha_{k,0}}{2\varepsilon_0}\right)^2 + 1} \tag{17}
\]

and

\[
\Gamma_1 = \Gamma_2^{-1} = \frac{\alpha_{k,0}}{2\varepsilon_0} + \sqrt{\left(\frac{\alpha_{k,0}}{2\varepsilon_0}\right)^2 + 1} \tag{18}
\]

On the basis of equations (17) and (18), the swelling pressure can be expressed as

\[
u_{sw} = RTc_s(\Gamma_1 + \Gamma_2 - 2) = 2RTc_s\left[\sqrt{\left(\frac{\alpha_{k,0}}{2\varepsilon_0}\right)^2 + 1} - 1\right] \tag{19}
\]

**Change in equilibrium**

When the hydraulic pressure and/or the salt concentration are changed in the external bulk solution, a new equilibrium condition is restored in the porous medium after a sufficiently long period of time. To evaluate this new condition, the change in the free energy per unit initial (undeformed) volume of the porous medium, \(\Delta G\), can be determined by assuming that it is a function of the macroscopic strain tensor, \(\varepsilon\), and the concentration of each fluid component (Dormieux et al., 2003). A second assumption that can be made refers to the reversible mechanical behaviour of the solid constituents; in such a case, the intrinsic dissipation, due to the solid skeleton strains, is zero (Dormieux et al., 2003; Dominijanni & Manassero, 2012a). On the basis of these assumptions, the increment in free energy per unit initial volume, under isothermal conditions, can be expressed as

\[
d\Delta G = \sigma : d\varepsilon + \sum_{k=w,i} \mu_k \frac{d(e_k)}{1 + \varepsilon_0} \tag{20}
\]

where \(\sigma\) is the total stress tensor and \(\varepsilon_0\) is the initial void ratio.

In equation (20), the chemical potentials of the pore solution, \(\mu_k\), can be substituted by the electrochemical potentials, \(\mu^*_k\), since the adding term given by

\[
F = \frac{d(e_{(\varepsilon_1, \varepsilon_2)}/(1 + \varepsilon_0))}{1 + \varepsilon_0} = F \frac{d\alpha_{k,0}}{1 + \varepsilon_0}
\]

is null, if the solid skeleton charge, \(\alpha_{k,0}\), is assumed constant.

Then, applying equations (6) and (7), the water chemical potential and the ion electrochemical potentials in the pore solution can be substituted by the corresponding potentials in the external solution. If the electroneutrality condition, equation (2), is taken into account, the ion electrochemical potentials of the bulk solution can be substituted by the ion chemical potentials. As a result, \(d\Delta G\) can be expressed using the readily available chemical potentials of the external bulk solution, instead of the chemical potentials of the pore solution. If the analysis is restricted to a unidimensional geometry, equation (20) becomes

\[
d\Delta G = \sigma : d\varepsilon + \sum_{k=w,i} \mu_k \frac{d(e_k)}{1 + \varepsilon_0} \tag{21}
\]

The free energy of the solid skeleton, which accounts for the interaction with the fluid phase, \(\Delta G^s\), can be derived by subtracting, from \(\Delta G\), the free energy of the fluid phase that is given as a function of the specific free energies of the components of the external bulk solution, \(F_k (k = w, 1, 2)\), to give

\[
\Delta G^s = \Delta G - \frac{e}{1 + \varepsilon_0} \sum_{k=w,1} \left(\tilde{\varepsilon}_k F_k\right) \tag{22}
\]

The free energy change, due to the interaction between the solid skeleton charge and the ions in the pore solution, is included in \(\Delta G^s\) because the specific free energy of the equilibrium bulk solution is subtracted from \(\Delta G\), instead of the specific free energy of the pore solution.

Using the thermodynamic relations

\[
\mu_k = F_k + \frac{u_k}{c_k} \quad k = w, 1, 2 \tag{23}
\]

\[
d\mu_k = \frac{du_k}{c_k} \quad k = w, 1, 2 \tag{24}
\]

\[
dF_k = -u_k \frac{d\left(\frac{1}{c_k}\right)}{c_k} \quad k = w, 1, 2 \tag{25}
\]

where \(u_k\) represents the partial pressure of the \(k\)-th component of the bulk solution, and \(\varepsilon^s\) can be expressed as

\[
d\Delta G^s = \sigma : d\varepsilon + \sum_{k=w,1} \frac{u_k}{1 + \varepsilon_0} d\left(\frac{e_k}{c_k}\right) \tag{26}
\]

Observing that \(\tilde{e}_k \equiv c_k\) and

\[
u_{sw} = u - \Pi \tag{27}
\]

where the osmotic pressure represents the sum of the ion partial pressures

\[
\Pi = \sum_{i=1}^{2} u_i \tag{28}
\]

the increment \(d\Delta G^s\) can be expressed as

\[
d\Delta G^s = \sigma : d\varepsilon + (u - \Pi) \frac{d\varepsilon}{1 + \varepsilon_0} + \sum_{i=1}^{2} \frac{u_i}{1 + \varepsilon_0} d\left(\frac{e_{c_i}}{c_i}\right) \tag{29}
\]

For infinitesimal strains, assuming that the solid component is incompressible

\[
de = -\frac{d\varepsilon}{1 + \varepsilon_0} \tag{30}
\]

and, as a consequence, equation (29) can be expressed as

\[
d\Delta G^s = [\sigma - (u - \Pi)]de + \sum_{i=1}^{2} \frac{u_i}{1 + \varepsilon_0} d\left(\frac{e_{c_i}}{c_i}\right) \tag{31}
\]

In order to derive the mechanical constitutive equations, it is convenient to work with the Gibb's free energy, \(G^s\), which is the following Legendre transform of \(\Delta G^s\)

\[
G^s = [\sigma - (u - \Pi)]e + \sum_{i=1}^{2} \frac{e_{c_i}}{1 + \varepsilon_0} u_i - \Delta G^s \tag{32}
\]

Taking into account equations (31) and (24), the infinitesimal increment in \(G^s\) is given by

\[
dG^s = e[d\sigma - (du - d\Pi)] + \sum_{i=1}^{2} \frac{e_{c_i}}{1 + \varepsilon_0} du_i - [e(d\varepsilon_1 + d\varepsilon_2)] \frac{1}{\nu_1 + \nu_2} dv_s \tag{33}
\]
where $d\mu_s$ is the salt chemical potential increment, which is defined as

$$d\mu_s = n_1 d\mu_1 + n_2 d\mu_2 = \frac{1}{c_s} d\Pi$$

(34)

The function $G^\phi$ can be considered to depend on the variables $[\alpha - (u - \Pi)]$ and $\mu_s$, therefore the simplest constitutive equations that can be inferred are

$$d\varepsilon = \beta_c [\varepsilon_0 - (du - d\Pi)] + \beta_v d\mu_s$$

(35)

$$d\rho_c \varepsilon_0 = \frac{\beta_c \varepsilon_0}{(v_1 + v_2)(1 + v_0)} = \beta_v [\varepsilon_0 - (du - d\Pi)] + \beta_w d\mu_s$$

(36)

Symmetry of the coefficients (i.e. $\beta_{cv} = \beta_{vc}$) can be demonstrated by considering $G^\phi_c$ as a continuous function of the variables $[\alpha - (u - \Pi)]$ and $\mu_s$ (Dominijanni & Manassero, 2012a).

Equation (35) can also be expressed in the form

$$d\varepsilon = du - d\varepsilon_sw = M de$$

(37)

where $M = 1/\beta_c$ is the unidimensional elastic modulus of the porous medium and $d\varepsilon_sw$ represents the swelling pressure increment, which is given by

$$d\varepsilon_sw = -\sigma d\Pi$$

(38)

where $\sigma = 1 + (\beta_{cv}/\beta_{vc} c_s)$ is the swelling coefficient (Dominijanni & Manassero, 2012a).

Since it is known, from experimental observations, that the swelling pressure tends to zero when $c_s \to \infty$, the swelling pressure, $\varepsilon_sw$, can be obtained as

$$\varepsilon_sw = \int_0^\infty \sigma d\Pi$$

(39)

The effective stress principle can be restored, on the basis of equation (37), if the effective stress increment, $d\sigma'$, is defined as

$$d\sigma' = du - d\varepsilon_sw$$

(40)

The classical definition of effective stress, $d\sigma' = du - d\varepsilon$, corresponds to the particular case for which $d\varepsilon_sw = 0$, that is, $\sigma = 0$ and $\beta_{cv} = -\beta_{vc} c_s$.

Adopting Donnan's assumptions and using equation (14), the swelling coefficient can be expressed as

$$\sigma = 1 - \frac{d\Pi}{d\Pi}$$

(41)

Dominijanni & Manassero (2012b) have demonstrated that equation (41) can also be expressed in the form

$$\sigma = 1 - \frac{v_1 + v_2}{v_1 \Gamma_1 + v_2 \Gamma_2}$$

(42)

Transport equations

If clay is interposed between two reservoirs with different hydraulic pressures or ion concentrations, a pore solution volumetric flux, $q$, and an ion mass flux, $J_i$, relative to the solid skeleton, are generated. In order to derive appropriate flux equations, the linear momentum balance equations of the fluid components can be developed on the basis of a number of assumptions. If only unidimensional problems are considered and inertial effects are neglected, these equations can be expressed as (Ehlers, 2002; Dominijanni & Manassero, 2010)

$$-\frac{\partial (\rho_w u_w)}{\partial x} + f_w = m_w$$

(43)

$$-\frac{\partial (\rho_i u_i)}{\partial x} + f_i = m_i$$

(44)

where $\rho_i$ is the porosity, $x$ is the spatial coordinate, $f_w$ is the external force per unit volume acting on the water, $m_w$ is the momentum supply of water, $f_i$ is the external force per unit volume acting on the $i$th ion, and $m_i$ is the momentum supply of the $i$th ion.

The external forces are gravity, which can be considered only to act on the solvent (i.e. water) in the $x$ direction, and the electric force, which is proportional to the electric potential gradient.

$$f_w = \rho_w g$$

(45)

$$f_i = -n_{w,sk} F \frac{\partial \phi}{\partial x}$$

(46)

where $\rho_w$ is the water density and $g$ is the gravitational acceleration.

The momentum supply of the fluid components can be expressed as

$$m_w = -u_w \nabla n + m_k^E$$

(47)

where $m_k^E$ represents the extra-momentum supply of the $k$th fluid component, which can be associated with the friction between the porous medium components that move with different velocities.

If the extra-momentum supply is associated with the frictional force per unit volume exchanged with the other components of the porous medium, it can be assumed to be equal to the sum of the binary interactions

$$m_w^E = m_w^E + \sum_{k=1}^{2} m_{w,j}^E$$

(48)

$$m_i^E = m_{i,j}^E + m_{i,sk}^E$$

(49)

where $w$ is the water, sk the solid skeleton, and $i$ and $j$ are the $i$th and $j$th ions. The friction between the ions and between the ions and the solid skeleton can be considered negligible for a dilute solution compared with the friction between the ions and the solvent

$$m_{i,j}^E, m_{i,sk}^E \ll m_{i,w}^E$$

(50)

Moreover, on the basis of the assumption of binary interaction, it can be assumed that

$$m_{w,j}^E = -m_{i,w}^E$$

(51)

For the friction forces per unit volume, the following constitutive equations can be adopted.

$$m_{w,sk}^E = \alpha (v_w - v_{sk}) = \alpha q$$

(52)

$$m_{i,j}^E = -m_{i,w}^E = \beta_i n_{w}(v_i - v_w)$$

(53)

where $\alpha$ and $\beta_i$ are friction coefficients, and $v_w, v_{sk}$ and $v_i$ are the water, solid skeleton and $i$th ion velocities respectively.

The factors $n$ and $(v_w - v_{sk})$ in equation (52), as well as $n_{w}$ and $(v_i - v_w)$ in equation (53), have been picked out as an indication of the fact that $m_{w,sk}^E = 0$ if $n = 0$ or
\( \nu_w - \nu_h = 0 \), and \( m^w \) = 0 if \( n^w_1 = 0 \) or \( \nu_i - \nu_w = 0 \). However, this does not mean that the friction coefficients \( \alpha \) and \( \beta_i \) are independent of \( n \), \( c \), and the relative velocities.

Taking into account these constitutive assumptions, and using the thermodynamic relation that relates the chemical potentials to the partial pressure

\[
d\mu_k = \frac{d\mu_k}{c_k} \quad k = w, \ 1, \ 2
\]

the following flux equations can be derived from equations (43) and (44).

\[
q = n(v_w - v_h)
\]

\[
= -\frac{n}{\alpha} \left( \frac{\partial \mu_w}{\partial x} - \rho_w \kappa + 2 \sum_{i=1}^{2} \frac{\partial \mu^c}{\partial x} \right)
\]

\[
J_i = n \sigma \left( v_i - v_h \right)
\]

\[
= q \sigma_i - n \frac{D_i}{RT} \sigma_i \frac{\partial \mu^c}{\partial x} \quad i = 1, \ 2
\]

where \( D_i = RT/\beta_i \) represents the macroscopic diffusion coefficient of the \( i \)th ion.

At this point, it is convenient to introduce the concept of a virtual external bulk solution that is in thermodynamic equilibrium with the pore solution at the generic position \( x \) within the porous medium (Dormieux et al., 1995; Yaroshchuk, 1995). The virtual solution coincides with the real bulk solutions in contact with the porous medium at the boundaries. The thermodynamic potentials in equations (55) and (56) can be substituted with the corresponding potentials of this virtual solution, using equations (6) and (7). This leads to a formulation that does not depend on the determination of the pore solution variables. Therefore, equations (55) and (56) can be expressed as

\[
q = -\frac{n}{\alpha} \left( \frac{\partial \mu}{\partial x} - \frac{\partial \Pi}{\partial x} - \rho \kappa + RT \sum_{i=1}^{2} \frac{\partial \sigma_i}{\partial x} + F \frac{\hat{e}_{w,0}}{e} \frac{\partial \phi}{\partial x} \right)
\]

\[
J_i = q \sigma_i c_i - n \frac{D_i}{RT} \sigma_i \frac{\partial \sigma_i}{\partial x} - z_i \sigma_i n \frac{D_i}{RT} F \frac{\partial \phi}{\partial x} \quad i = 1, \ 2
\]

If an external electric current is not applied to the porous medium, the electric potential derivative in equations (57) and (58) can be eliminated by means of the condition of electric current, \( I_e \), equal to zero.

\[
I_e = F \sum_{i=1}^{2} z_i J_i = 0
\]

The resulting flux equations can be expressed as follows

\[
q = -\frac{k}{\gamma_w} \left( \frac{\partial \mu}{\partial x} - \frac{\partial \Pi}{\partial x} \right)
\]

\[
J_s = (1 - \omega)qc_s - nD_w \frac{\partial c_s}{\partial x}
\]

where

\[
k = \frac{n^\gamma_w}{\alpha + \frac{RT}{\nu_1} (\Gamma_1 - 2 \nu_2 c_1^2)}
\]

\[
\omega = 1 - \frac{\nu_1 D_2 + \nu_2 D_1}{\nu_1 D_2 + \nu_2 D_1} \Gamma_1 \Gamma_2
\]

\[
J_s = \frac{J_1}{v_1} = \frac{J_2}{v_2}
\]

\[
D_w = (1 - \omega)D_s
\]

\[
D_s = \frac{(v_1 + v_2) D_1 D_2}{v_1 D_2 + v_2 D_1}
\]

where \( k \) is the hydraulic conductivity, \( \omega \) is the reflection coefficient, \( J_s \) is the salt molar flux, \( D_w^s \) is the osmotic effective diffusion coefficient, and \( D_s \) is the macroscopic salt diffusion coefficient.

Dominijanni & Manassero (2012) have demonstrated that, if the microscopic deviations of the variables from their average values are assumed to be negligible, the macroscopic ion diffusion coefficients, \( D_i \), are equal to the ion effective diffusion coefficients, \( D^e_i \)

\[
D_i = D^e_i = \tau_m D_{i,0} \quad i = 1, \ 2
\]

\[
D_s = D^e_s = \frac{(v_1 + v_2) D_1 D_2}{v_1 D_2 + v_2 D_1}
\]

\[
= \tau_m \frac{(v_1 + v_2) D_1 D_2}{v_1 D_2 + v_2 D_1}
\]

where \( \tau_m \) is the dimensionless matrix tortuosity factor, which accounts for the tortuous nature of the actual diffusive pathway through the porous medium (Malusis & Shackelford, 2002b); \( D_{i,0} \) is the free (aqueous) solution diffusion coefficient of the \( i \)th ion; \( D^e_i \) is the salt effective diffusion coefficient; and \( D_{i,0} \) is the free-solution diffusion coefficient of the salt.

An interesting observation is that the reflection coefficient, \( \omega \), results as equal to the swelling pressure coefficient, \( \hat{\sigma} \), when the ion free-solution diffusion coefficients are equal.

When the solid skeleton electric charge is equal to zero, the ion partition coefficients, \( \Gamma_i \), are equal to 1, and equations (60) and (61) reduce to the Darcy equation and the classical advective diffusion equation respectively.

The osmotic effective diffusion coefficient, \( D^e_o \), results as related to the reflection coefficient, \( \omega \), through equation (65), so that \( D^e_o = 0 \) when \( \omega = 1 \). As a result, the condition \( \omega = 1 \) implies a null salt flux through the porous medium, which in this case can be said to act as a ‘perfect’ or ‘ideal’ barrier.

If equation (65) is compared with the expression for \( D^e_o \) proposed by Malusis & Shackelford (2002b) and Malusis et al. (2012)

\[
D^e_o = \tau_i D^e_s
\]

where \( \tau_i \) is the restrictive tortuosity factor, the result is that \( \tau_i \) is given by

\[
\tau_i = (1 - \omega) \frac{D_s}{D_s}
\]

Moreover, if the hypotheses implied by equation (68) are adopted, the expression for \( \tau_i \) reduces to

\[
\tau_i = 1 - \omega
\]

The coefficient \( k \) can be measured, under steady-state conditions, using traditional permeameters. Malusis et al. (2001) developed a testing apparatus to determine \( \omega \) and \( D^e_o \). This apparatus is able to impose the condition of no volumetric flux (\( q = 0 \)) through a soil sample in contact with two external solutions, maintained at constant salt concentrations, so that the global or averaged values of the coefficients can be measured. The global values of \( \omega \) and \( D^e_o \) are defined as (Auclair et al., 2002)
where \( c_1 \) and \( c_b \) represent the salt concentration at the top and the bottom boundaries of the clay sample respectively, and \( \Delta c_{s} = c_t - c_b \) is their difference. These coefficients can be determined by means of the following relations under steady-state conditions.

\[
\omega_0 = \left( \frac{\Delta \Pi}{\Delta \Pi} \right)_{q=0}
\]

\[
D^* = \left( \frac{L}{n} \right)_{q=0}
\]

where \( \Delta u = u_t - u_b \) and \( \Delta \Pi = \Pi_t - \Pi_b \) represent the differences between the hydraulic pressure and the osmotic pressure at the boundaries of the clay sample, and \( L \) is the length of the sample.

It is interesting to observe that the relationship between \( D^* \) and \( \omega_0 \) is also maintained between their corresponding global values; in fact, inserting equation (65) into equation (73) with \( D_s = D^* \) leads to

\[
D^*_{ag} = (1 - \omega_0)D^*_k
\]

\[
= (1 - \omega_0)\tau_nD_{s,0}
\]

For a salt constituted by monovalent ions, inserting equation (63) into equation (72) and using equations (17) and (18), the following expression for \( \omega_0 \) is obtained

\[
\omega_0 = 1 + \frac{c_{\text{eq},0}}{2\Delta c_{s}} [Z_2 - Z_1 - (2t_1 - 1) \ln \left( \frac{Z_2 + 2t_1 - 1}{Z_1 + 2t_1 - 1} \right)]
\]

where

\[
t_1 = \frac{D_{1,0}}{D_{1,0} + D_{2,0}}
\]

\[
Z_1 = \left( 1 + \frac{2\Delta c_{s} e}{c_{\text{eq},0}} \right)^2
\]

\[
Z_2 = \left( 1 + \frac{2\Delta c_{s} e}{c_{\text{eq},0}} \right)^2
\]

and \( t_1 \) is the cation transport number.

MATERIALS AND METHODS

Materials

The powdered bentonite tested in this study is an Indian sodium bentonite that is used for the production of a needle-punched GCL. The bentonite is characterised by a cation exchange capacity (CEC, measured using the methylene blue adsorption method) of 105 meq/100 g. The mineralogical composition, evaluated through X-ray diffraction analysis, indicates a bentonite that is composed primarily of smectite (> 98%), with traces of calcite, quartz, mica and gypsum.

The bentonite is characterised by a liquid limit (LL) of 525% and a hydraulic conductivity of \( 8 \times 10^{-12} \) m/s, measured at a 27-5 kPa confining effective stress using de-ionised water (DW) as the permeant liquid.

Sodium solutions were prepared with sodium chloride (ACS reagent, purity > 99%) and DW. The sodium solutions were prepared at different molarity values in the range 5–100 mM, with the aim of investigating the effect of the monovalent cations on the osmotic behaviour of the bentonite. The DW (pH = 6.95; electrical conductivity (EC) at 20°C = 0.6 mS/m) consisted of tapwater processed through a series of activated carbon filters, a reverse osmosis process and, finally, an ultraviolet (UV) lamp (Elix Water Purification system). Moreover, the DW was de-aerated prior to use. The EC measured at 20°C for the sodium chloride solutions ranged from 60.5 mS/m to 1.1 mS/m.

Bentonite preparation. Prior to the osmotic property determination, the bentonite was submitted to a process with the aim of removing the soluble salts (mainly sodium) that are naturally present inside the material, owing to its marine origin. The treatment prevents soluble salts from interfering with the determination of the osmotic properties.

Previous studies (Malusis et al., 2001; Malusis & Shackelford, 2002a, 2002b; Shackelford & Lee, 2003; Yeo et al., 2005; Kang & Shackelford, 2009; Di Emidio, 2010) have used the ‘flushing’ method to remove soluble salts. This method consists of an initial permeation phase, performed under back-pressure, which requires a long period of time (i.e. from months to a year), because of the low bentonite hydraulic conductivity.

In this study the ‘squeezing’ method has been used, with the aim of reducing the salt removal time. The squeezing method consists of a series of consecutive phases of powder bentonite hydration with DW, at a higher water content than the LL, and drained consolidation, performed in a consolidometer under a maximum load of 500 kPa. Moreover, the drained solution is sampled daily, and the EC is monitored to evaluate the soluble salt concentration in the bentonite pore water. After the squeezing process, the material is oven-dried at 105°C and pulverised once again. When a 51 consolidometer is used, the above procedure can produce about 500 g of squeezed dry powder bentonite, characterised by an EC value lower than 50 mS/m, in 40–50 days. The results of the EC monitoring during the squeezing process are reported in Fig. 1.

The specimen for the chemico-osmotic test is prepared by rehydrating the squeezed bentonite with DW at a lower water content than the LL value, and then by statically compacting the material in a compaction mould, while allowing the excess water to be released.

Dry powder bentonite is required for the swelling pressure test.

Testing apparatus and procedures

Chemico-osmotic test. The testing apparatus used to measure the global reflection coefficient and the global osmotic effective diffusion coefficient is described in detail in Malusis et al. (2001). The main components of the apparatus, shown in Fig. 2, are the osmotic cell, the flow pump system, the pressure transducer, which is used to measure the differential pressure that develops across the specimen during the test, and the data acquisition system.

The cell consists of a modified rigid-wall permeameter, in which the top piston and the bottom pedestal are equipped with three ports each: two enable the different solutions to circulate through the top (sodium chloride solution) and the bottom (DW) porous stones, with the aim of establishing a constant concentration gradient across the specimen. The third port is installed in both the top piston and the bottom pedestal to allow the differential pressure across the specimen to be measured.

The flow pump system, which consists of a dual-carriage
syringe pump and two stainless steel accumulators (Model 33 twin syringe pump, produced by Harvard, Holliston, MA), prevents volumetric flux through the specimen by simultaneously injecting into and withdrawing from the porous stones the same volume of solution. In order to obtain this result, the syringes have to move at the same rate.

The test was performed according to the procedure proposed by Malusis et al. (2001): a solution containing a known electrolyte concentration (sodium chloride) was circulated in the top porous stone, while DW was circulated in the bottom porous stone. The concentration difference across the specimen was maintained constant by continuously infusing the two liquids at the boundaries of the specimen.

As the specimen was first squeezed with DW to remove the soluble salts, the EC of the electrolyte solutions in the flux exiting from the porous stones at the steady state was induced solely by the contributions of Cl\(^-\) and Na\(^+\) ions. Moreover, the calibration reported in Fig. 3 shows that the relation between the EC and solution molarity was linear over the concentration range examined in the study. As a consequence, the EC of the withdrawn fluxes (i.e. from the top and bottom porous stones) was monitored by sampling the solution contained in the pistons, and the sodium chloride molar concentration was derived using a linear relation.

Since the volumetric flux through the specimen was hindered, the global reflection coefficient could be calculated using equation (74).

The diffusive solute flux through the specimen was calculated for the \(n\)th sampling interval according to

\[
f_s^n = \frac{\sum_{m=1}^{n} (c_m^n \Delta P_m^n)}{A_S \Delta \rho_s^n} = \frac{\Delta Q_s^n}{\Delta \tau^n}
\]  

(81)
where \( c_{n}^{s} \) is the solute molar concentration measured by sampling the solution coming out from the bottom porous stone, \( \Delta V^m \) is the volume of the solution circulating in the porous stones in the interval \( \Delta r^m \), \( A_S \) is the cross-section of the specimen, and \( \Delta Q^m \) is the cumulative salt molar mass per unit area that passed through the specimen. The global osmotic effective diffusion coefficient, \( D_{avg}^e \), is calculated at the steady state as

\[
D_{avg}^e = \frac{\Delta Q}{\Delta r} n \frac{L}{n(c_{top,avg} - c_{bot,avg})} \quad (82)
\]

where \( c_{top,avg} \) and \( c_{bot,avg} \) are the average top and bottom salt concentrations respectively.

### RESULTS

#### Chemico-osmotic test results

The chemico-osmotic test was performed using the oven-dried squeezed material, rehydrated with DW and then

**Swelling pressure test.** The swelling pressure apparatus, shown in Fig. 4, consists primarily of a stainless steel oedometer cell, a sodium chloride solution supply tank placed above the pressure panel, a displacement transducer connected to the cell top piston, which is used to measure the axial strains of the specimen, a load cell, and a data acquisition system.

The swelling pressure apparatus consists of a rigid cell that confines the sample (i.e. the oedometer), which allows access to the water through both porous stones. The cell is connected to a pressure panel that allows the specimen to be back-pressurised. The rigid piston above the upper porous stone is connected to the load cell, which measures the pressure that has to be applied in order to hinder the axial strain of the specimen.

The test procedure requires a known amount of dry material to be dusted inside the oedometer ring, the cell to be assembled, and a sodium chloride solution to be supplied. The specimen, which is characterised by an initial dry height of 5 mm, is allowed to swell to 10 mm. The piston is then blocked, the sample is back-pressurised, and the steady-state swelling pressure is recorded after a short transitional phase.

Since the bentonite that is initially dusted inside the oedometer is dry, the pressure increases for a number of days during the hydration phase, and the steady-state swelling pressure is reached when hydration has been completed.

---

**Fig. 3.** Calibration of sodium chloride (NaCl) concentration with electrical conductivity. \( R^2 \) is coefficient of determination of regression line

**Fig. 4.** Schematic view of the swelling pressure apparatus
statically compacted, in a drained compaction mould, at a porosity $n$ equal to 0.81 ($e = 4.26$). After the preparation phase, the 17 mm thick specimen was transferred to the cell for the osmotic test.

After assembling the cell, DW was circulated through the top piston and the bottom pedestal for 2 weeks in order to establish a steady baseline differential pressure, before a concentration gradient was applied to the specimen. A source concentration of sodium chloride was then injected into the top porous stone, while DW was continuously circulated in the bottom porous stone.

A multiple-stage chemico-osmotic test was performed by sequential circulation of chemical solutions containing 5.16, 10.27, 20.24, 51.94 and 109.31 mM sodium chloride concentrations at a constant flow rate of 0.05 ml/min.

The EC values of the salt mass fluxes withdrawn from the top and the bottom porous stones, measured during the testing stages, are shown in Fig. 5. The measured values depend on the sodium chloride concentrations imposed at the boundaries of the specimen: the EC values progressively increase during the test as the sodium chloride concentration of the injected solution in the top porous stone rises. The trends of the electrical conductivity of the flux withdrawn from the top porous stone, $EC_{t,exit}$, and of that withdrawn from the bottom porous stone, $EC_{b,exit}$, both show that a steady state has been reached for each stage.

Moreover, the difference between the EC values measured in the flux withdrawn from the top porous stone ($EC_{t,exit}$) and those of the solutions injected into the same stone ($EC_{t,ref}$) is due to the loss in sodium chloride mass induced by the diffusion through the bentonite from the top to the bottom boundary.

The global reflection coefficient values, $\omega_g$, obtained during the multiple-stage chemico-osmotic test, are shown in Fig. 6 as a function of time. The $\omega_g$ values are determined using equation (74), on the basis of the differential pressure, $\Delta u$, measured during the test with a time step of 10 min, and the osmotic pressure, $\Delta\Pi$, calculated from the average of the top and bottom sodium chloride concentrations. The steady-state values of the variables are reported in Table 1 for each concentration stage. The sodium chloride concentrations were derived from the measured EC using the linear calibration reported in Fig. 3.

As far as the EC measurements are concerned, the trend of the global reflection coefficient shows that a steady state has been reached for each stage. The steady-state $\omega_g$ values tend to decrease as the salt concentration in the top porous stone increases. The recorded values range from 68%, for a

![Fig. 5. Electrical conductivity of salt flux withdrawn from (a) top porous stone and (b) bottom porous stone as a function of time during multiple-stage chemico-osmotic test](image-url)
5.16 mM sodium chloride source concentration, to 5%, for a 109.31 mM sodium chloride source concentration. The global reflection coefficient can be assumed approximately null for higher molarities.

The cumulative molar mass per unit area, $Q$, of the sodium chloride that migrated through the specimen during the multiple-stage test is reported in Fig. 7. The values of the global osmotic effective diffusion coefficient, $D_{\text{avg}}^*$, which

Table 1. Steady-state values of variables involved in multiple-stage chemico-osmotic test

<table>
<thead>
<tr>
<th>$c_{t,\text{ref}}$: mM</th>
<th>$c_{l,\text{exit}}$: mM</th>
<th>$c_{b,\text{exit}}$: mM</th>
<th>$c_{t,\text{avg}}$: mM</th>
<th>$c_{b,\text{avg}}$: mM</th>
<th>$\Delta u$: kPa</th>
<th>$\Delta \Pi$: kPa</th>
<th>$\omega_g$</th>
<th>$D_{\text{avg}}^*$: m$^2$/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.16</td>
<td>5.12</td>
<td>0.83</td>
<td>5.14</td>
<td>0.42</td>
<td>15.65</td>
<td>23.02</td>
<td>0.68</td>
<td>2.54 x 10$^{-10}$</td>
</tr>
<tr>
<td>10.27</td>
<td>9.61</td>
<td>0.85</td>
<td>9.94</td>
<td>0.43</td>
<td>26.87</td>
<td>46.33</td>
<td>0.58</td>
<td>3.52 x 10$^{-10}$</td>
</tr>
<tr>
<td>20.24</td>
<td>18.93</td>
<td>1.45</td>
<td>19.58</td>
<td>0.72</td>
<td>30.32</td>
<td>91.89</td>
<td>0.33</td>
<td>4.19 x 10$^{-10}$</td>
</tr>
<tr>
<td>51.94</td>
<td>47.39</td>
<td>4.39</td>
<td>49.67</td>
<td>2.19</td>
<td>32.38</td>
<td>231.30</td>
<td>0.14</td>
<td>4.60 x 10$^{-10}$</td>
</tr>
<tr>
<td>109.31</td>
<td>97.18</td>
<td>9.78</td>
<td>103.24</td>
<td>4.89</td>
<td>23.96</td>
<td>479.21</td>
<td>0.05</td>
<td></td>
</tr>
</tbody>
</table>

The cumulative molar mass per unit area, $Q$, of the sodium chloride that migrated through the specimen during the multiple-stage test is reported in Fig. 7. The values of the global osmotic effective diffusion coefficient, $D_{\text{avg}}^*$, which

Fig. 6. Global reflection coefficient as a function of time during multiple-stage chemico-osmotic test

Fig. 7. Cumulative molar mass of sodium chloride per unit area as a function of time during multiple-stage chemico-osmotic test:
(a) $c_{t,\text{ref}}$ = 10.27 mM; (b) $c_{t,\text{ref}}$ = 20.24 mM; (c) $c_{t,\text{ref}}$ = 51.94 mM; (d) $c_{t,\text{ref}}$ = 109.31 mM
have been obtained from the $Q$ measurements shown in Fig. 7, are reported in Table 1.

Swelling pressure test results

The swelling pressure test was performed using dry specimens, prepared with the squeezed, oven-dried bentonite and characterised by an initial dry height of 5 mm, which were allowed to swell to 10 mm during hydration. The final hydrated volume of the specimens corresponded to $n = 0.81$.

The test was performed by hydrating five specimens with five different sodium chloride solutions, characterised by increasing concentrations: 5, 10, 20, 50 and 100 mM. After hydration, the specimen volume change was inhibited, and the value of the swelling pressure was recorded after a short transitional phase.

The swelling pressure trend is reported in Fig. 8 for each test as a function of time. Since, during the tests, the load cell was unloaded until the specimen swelled to 10 mm, the initial swelling/hydration phase of the dry material (from 5 to 10 mm) was characterised by null swelling pressure values.

All the tests show that the swelling pressure increases for approximately 15–20 h, during the controlled hydration phase (i.e. when the volumetric strain is inhibited), and that the equilibrium swelling pressure is reached when the hydration phase is completed. Moreover, in the tests with lower sodium chloride concentrations (i.e. for 5 and 10 mM sodium chloride equilibrium solutions), the specimens were successively back-pressurised to 300 kPa. The obtained results show that the swelling pressure value does not change after back-pressurisation.

In the test performed using the 100 mM sodium chloride solution, the bentonite specimen did not rise to 10 mm, as it stopped at a height of 9.5 mm, and the load cell was never loaded during the test. For this reason, the swelling pressure for this test was taken equal to zero.

INTERPRETATION OF TEST RESULTS

The experimental results can be related to the physical and chemical properties of the tested bentonite under the assumption that the microscopic deviations of the state variables from their average values are negligible. In such a case, both the global reflection coefficient and the swelling pressure depend on the solid skeleton electric charge through equations (19) and (77). Therefore, from the best-fitting of the theoretical curves with the experimental data of both tests, a value of $\bar{c}_{k,0}$ equal to 90 mM was found. The obtained theoretical curves are reported in Figs 9 and 10, together with the experimental data.

![Fig. 8. Swelling pressure as a function of time. (a) $c_s = 5$ mM; (b) $c_s = 10$ mM; (c) $c_s = 20$ mM; (d) $c_s = 50$ mM](image-url)

![Fig. 9. Global reflection coefficient, $\omega_g$, as a function of average sodium chloride (NaCl) concentration at top boundary of bentonite specimen, with best-fitting theoretical curve, obtained for $\bar{c}_{k,0} = 90$ mM in equation (77) (continuous line)](image-url)
The salt concentration at the top boundary was taken equal to \( c_{b,\text{avg}} \) to determine \( \omega_g \), and the salt concentration at the bottom boundary was considered equal to zero: that is, \( c_b = c_{b,\text{avg}} = 0 \). The sodium transport number was calculated from the sodium and chloride free-solution diffusion coefficient values \( D_{\text{Na,0}} = 13.3 \times 10^{-8} \text{m}^2/\text{s}, D_{\text{Cl,0}} = 20.3 \times 10^{-8} \text{m}^2/\text{s} \).

In Fig. 11, the experimental reflection coefficient data were also fitted with the empirical semi-log linear curve proposed by Shackelford et al. (2003) and Malusis et al. (2003),

\[
\omega_g = A + B \log(c_{t,\text{avg}})
\]

where \( A \) and \( B \) are the regression parameters. The value of the coefficient of determination \( R^2 \) close to 1 confirms the ability of this empirical curve to fit the \( \omega_g \) experimental data, as was found by Malusis et al. (2003) for the Kemper clay liner, with a bentonite porosity, \( n \), of 0.79 for different concentrations of potassium chloride (KCl). The differences in the parameters obtained from the two studies can be attributed both to the different mineralogical compositions of the tested bentonites and to the different salts contained in the pore solutions.

The theoretical linear relationship between \( D_{\text{Na}}^* \) and \( (1 - \omega_g) \) in Fig. 12 is a consequence of assuming that the pore-scale variations in the hydraulic pressure, ion concentrations and water velocity within the soil are negligible; as a result, the agreement of the experimental data with the linear relationship is an indication of the acceptability of this assumption.

The goodness of the linear fitting shown in Fig. 12 \( (R^2 = 0.9810) \) and the possibility of fitting both the global reflection coefficient and the swelling pressure data with a single value of \( c_{b,0} \) are indications of the ability of the
proposed theoretical approach to simulate the bentonite behaviour.

CONCLUSIONS

A theoretical approach that takes into account the interaction between the electric charge of the bentonite solid skeleton and the ions contained in the pore solution has been proposed. The phenomenological parameters introduced in this theoretical approach were measured for a bentonite specimen with porosity, $n$, of 0.81, over a range of sodium chloride concentrations in the pore solution varying from 5 mM to 100 mM. Both the global reflection coefficient, $\omega_g$, and the swelling pressure, $\tau_m$, were found to decrease with an increase in the salt concentration. This result is in agreement with the trends given by the proposed theoretical model, assuming that the microscopic deviations of the pore solution state variables from their average values are negligible. If this assumption is accepted, the experimental data can be used to derive the electric charge of the solid skeleton (per unit solid volume), $\xi_{k,0}$, and the tortuosity factor, $\tau_m$. The mechanical behaviour and the transport properties of bentonite can be estimated from these physical properties to evaluate its performance as a hydraulic and contaminant barrier in field applications. However, in order to verify the applicability of the proposed model under different boundary conditions and for different salts contained in the pore solution, further experimental evaluations must be conducted. Moreover, the results obtained for a single salt contained in the bentonite pore solution need to be extended to the more general problem of a solution containing an unspecified number of salts, in order to evaluate the performance of bentonites that are used as contaminant barriers for real leachates.

NOTATION

$A$, $B$ regression parameters
$A_s$ cross-section of specimen
$c_h$ salt concentration at bottom boundary of specimen
$c_{h,\text{avg}}$ average salt concentration at bottom boundary of specimen
$c_{h,\text{exit}}$ salt concentration of flux withdrawn from bottom porous stone
$c_i$ molar concentration of $i$th ion in equilibrium bulk solution
$c_k$ molar concentration of $k$th ion in pore solution
$c_{k,\text{fluid}}$ molar concentration of $k$th fluid component in equilibrium bulk solution
$c_k$ molar concentration of $k$th fluid component in pore solution
$c_s$ salt molar concentration in equilibrium bulk solution
$\Delta c_k$ salt concentration difference
$c_{k,0}$ molar concentration per unit solid volume of solid skeleton electric charge
$c_{\text{t,avg}}$ average salt concentration at top boundary of specimen
$c_{\text{t,exit}}$ salt concentration of flux withdrawn from top porous stone
$c_{\text{t,ref}}$ source salt concentration
$c_w$ water molar concentration in equilibrium bulk solution
$\bar{c}_w$ water molar concentration in pore solution
$D_i$ macroscopic diffusion coefficient of $i$th ion
$D_{i,0}$ free (aqueous) solution diffusion coefficient of $i$th ion
$D_i^g$ effective diffusion coefficient of $i$th ion
$D_{i,0}^g$ free (aqueous) solution salt diffusion coefficient
$D_i^s$ effective salt diffusion coefficient
$D_i^s$ osmotic effective diffusion coefficient
$D_{i,0}^s$ global osmotic effective diffusion coefficient
$E_{\text{C,exit}}$ electrical conductivity of flux withdrawn from bottom porous stone
$E_{\text{C,exit}}$ electrical conductivity of flux withdrawn from top porous stone
$E_{\text{C,ref}}$ electrical conductivity of source solution
$\varepsilon$ void ratio
$\varepsilon_0$ initial void ratio
$F$ Faraday’s constant (96485 C/mol)
$F_k$ free energy per unit molar mass of $k$th component of fluid phase

Table 2. Comparison between physical parameters derived from interpretation of experimental results in this work and those obtained by Dominijanni & Manassero (2012b) from interpretation of tests carried out by Malusis & Shackelford (2002a, 2002b)

<table>
<thead>
<tr>
<th>Experimental data</th>
<th>Material</th>
<th>Method for removing soluble salts</th>
<th>Tests</th>
<th>Salt in pore solution</th>
<th>Porosity, $n$</th>
<th>Solid skeleton charge concentration, $c_{k,0}$: mM</th>
<th>Tortuosity factor, $\tau_m$</th>
</tr>
</thead>
<tbody>
<tr>
<td>This study</td>
<td>Natural sodium bentonite Geosynthetic clay liner containing natural sodium bentonite</td>
<td>Squeezing Flushing</td>
<td>Chemico-osmotic test and swelling pressure test Chemico-osmotic test</td>
<td>NaCl</td>
<td>0.81</td>
<td>90</td>
<td>0.31</td>
</tr>
<tr>
<td>Malusis &amp; Shackelford (2002a, 2002b)</td>
<td>Natural sodium bentonite</td>
<td>Flushing</td>
<td>Chemico-osmotic test</td>
<td>KCl</td>
<td>0.79–0.80</td>
<td>46</td>
<td>0.14</td>
</tr>
</tbody>
</table>
\[ f_i \] \text{external force per unit volume acting on } i\text{th ion} \\
\[ f_w \] \text{external force per unit volume acting on water} \\
\[ G^k \] \text{Gibbs free energy of solid skeleton per unit volume of porous medium} \\
\[ g \] \text{gravitational acceleration} \\
\[ I_e \] \text{electric current density} \\
\[ \Xi_V \] \text{free energy per unit (undeformed) volume of porous medium} \\
\[ \Theta^k \] \text{free energy of solid skeleton per unit (undeformed) volume of porous medium} \\
\[ J_i \] \text{molar flux of } i\text{th ion relative to solid skeleton} \\
\[ J_k \] \text{salt molar flux} \\
\[ k \] \text{hydraulic conductivity} \\
\[ L \] \text{length of specimen} \\
\[ M \] \text{unidimensional elastic modulus of porous medium} \\
\[ m_i \] \text{momentum supply of } i\text{th ion} \\
\[ m^i \] \text{extra-momentum supply of } i\text{th ion} \\
\[ m_k \] \text{momentum supply of } k\text{th fluid component} \\
\[ m^k \] \text{extra-momentum supply of } k\text{th fluid component} \\
\[ m^L \] \text{extra-momentum supply exchanged between } h\text{th and } m\text{th components of porous medium} \\
\[ \left( \text{sk } = \text{solid skeleton}, \text{w } = \text{water}, 1 = \text{cation}, \quad 2 = \text{anion} \right) \] \\
\[ m_w \] \text{momentum supply of water} \\
\[ n \] \text{porosity} \\
\[ Q \] \text{cumulative salt molar mass per unit area} \\
\[ \Delta Q \] \text{increment in cumulative salt molar mass per unit area} \\
\[ q \] \text{water volumetric flux relative to solid skeleton} \\
\[ R \] \text{universal gas constant } \left( = 8.314 \text{ J/(molK)} \right) \\
\[ T \] \text{absolute temperature in equilibrium bulk solution} \\
\[ \mathcal{T} \] \text{absolute temperature in pore solution} \\
\[ t \] \text{time} \\
\[ \Delta t \] \text{time interval} \\
\[ v_1 \] \text{cation transport number} \\
\[ u \] \text{hydraulic pressure in equilibrium bulk solution} \\
\[ \Delta u \] \text{hydraulic pressure difference} \\
\[ \bar{u} \] \text{hydraulic pressure in pore solution} \\
\[ u_b \] \text{hydraulic pressure at bottom boundary of specimen} \\
\[ u_i \] \text{partial pressure of } i\text{th ion component in equilibrium bulk solution} \\
\[ \bar{u}_i \] \text{partial pressure of } i\text{th ion component in pore solution} \\
\[ u_k \] \text{partial pressure of } k\text{th fluid component in equilibrium bulk solution} \\
\[ \bar{u}_k \] \text{partial pressure of } k\text{th fluid component in pore solution} \\
\[ u_{sw} \] \text{swelling pressure} \\
\[ \bar{u}_t \] \text{hydraulic pressure at top boundary of porous medium} \\
\[ u_w \] \text{partial pressure of water in equilibrium bulk solution} \\
\[ \Delta u^w \] \text{partial pressure of water in pore solution} \\
\[ \Delta V^w \] \text{volume of solution circulating in porous stones} \\
\[ v_i \] \text{velocity of } i\text{th ion} \\
\[ v_{sk} \] \text{velocity of solid skeleton} \\
\[ v_w \] \text{water velocity} \\
\[ x \] \text{spatial coordinate} \\
\[ z_i \] \text{electrochemical valence of } i\text{th ion} \\
\[ z_{sk} \] \text{electrochemical valence of solid skeleton} \\
\[ \alpha_i, \beta_i \] \text{friction coefficients} \\
\[ \beta_{ss}, \beta_{sv}, \beta_{vv} \] \text{phenomenological parameters in chemomechanical constitutive laws} \\
\[ \Gamma_i \] \text{partial coefficient of } i\text{th ion} \\
\[ \gamma_w \] \text{water unit weight } \left( = \rho_w g \right) \\
\[ \epsilon \] \text{scalar strain in unidimensional geometry} \\
\[ \epsilon \] \text{strain tensor} \\
\[ \mu_i \] \text{chemical potential of } i\text{th ion in equilibrium bulk solution} \\
\[ \bar{\mu}_i \] \text{chemical potential of } i\text{th ion in equilibrium pore solution} \\
\[ \mu^0_w, \bar{\mu}^0_w, \bar{\mu}_w \] \text{integration constants} \\
\[ \mu^i \] \text{electrochemical potential of } i\text{th ion in equilibrium bulk solution} \\
\[ \bar{\mu}^i \] \text{electrochemical potential of } i\text{th ion in pore solution} \\
\[ \mu_s \] \text{chemical potential of } k\text{th fluid component in equilibrium bulk solution} \\
\[ \mu_k \] \text{chemical potential of } k\text{th fluid component in pore solution} \\
\[ \mu_w \] \text{water chemical potential in equilibrium bulk solution} \\
\[ \bar{\mu}_w \] \text{water chemical potential in pore solution} \\
\[ \nu_i \] \text{stoichiometric coefficient of } i\text{th ion} \\
\[ \Pi \] \text{osmotic pressure in equilibrium bulk solution} \\
\[ \Delta \Pi \] \text{osmotic pressure difference} \\
\[ \Pi_b \] \text{osmotic pressure at bottom boundary of specimen} \\
\[ \Pi_t \] \text{osmotic pressure at top boundary of specimen} \\
\[ \rho_w \] \text{water density} \\
\[ \sigma \] \text{scalar total stress in unidimensional geometry} \\
\[ \sigma' \] \text{scalar effective stress in unidimensional geometry} \\
\[ r_m \] \text{matrix tortuosity factor} \\
\[ r_r \] \text{restrictive tortuosity factor} \\
\[ \phi \] \text{electric potential in equilibrium bulk solution} \\
\[ \phi \] \text{electric potential in pore solution} \\
\[ \psi \] \text{Donnan’s electric potential} \\
\[ \omega \] \text{reflection coefficient} \\
\[ \omega_g \] \text{global reflection coefficient} \\
\[ \bar{\sigma} \] \text{swelling pressure coefficient} \\

**Indexes** \\
\[ i \] \text{index for ions } (1 = \text{cation}, 2 = \text{anion}) \\\n\[ j \] \text{index for ions } (1 = \text{cation}, 2 = \text{anion}) \\\n\[ k \] \text{index for fluid components } (w = \text{water}, 1 = \text{cation}, 2 = \text{anion}) \\

**Subscripts** \\
\[ 1 \] \text{cation} \\\n\[ 2 \] \text{anion} \\\n\[ b \] \text{bottom of specimen} \\\n\[ g \] \text{global value} \\\n\[ sk \] \text{solid skeleton} \\\n\[ t \] \text{top of specimen} \\\n\[ w \] \text{water} \\

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