

1 Predictive surface complexation model of the calcite-aqueous solution 2 interface: the impact of high concentration and complex composition of brines

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14 **Abstract**

15 Electrochemical interactions at calcite-water interface are characterized by the zeta potential and
16 play an important role in many subsurface applications. In this work we report a new physically
17 meaningful surface complexation model that is proven to be efficient in predicting calcite-water zeta
18 potentials for a wide range of experimental conditions.

19 Our model uses a two-stage optimization for matching experimental observations. First, equilibrium
20 constants are optimized, and the Stern layer capacitance is optimized in the second stage. The
21 model is applied to a variety of experimental sets that correspond to intact natural limestones
22 saturated with equilibrated solutions of low-to-high salinity, and crushed Iceland Spar sample
23 saturated with NaCl at non-equilibrium conditions.

24 The proposed linear correlation of the Stern layer capacitance with the ionic strength is the main
25 novel contribution to our surface complexation model without which high salinity experiments cannot
26 be modelled. Our model is fully predictive given accurately known conditions. Therefore, the reported
27 parameters and modelling protocol are of significant importance for improving our understanding of
28 the complex calcite-water interfacial interactions. The findings provide a robust tool to predict
29 electrochemical properties of calcite-water interfaces, which are essential for many subsurface
30 applications including hydrology, geothermal resources, CO₂ sequestration and hydrocarbon
31 recovery.

32 **1. Introduction**

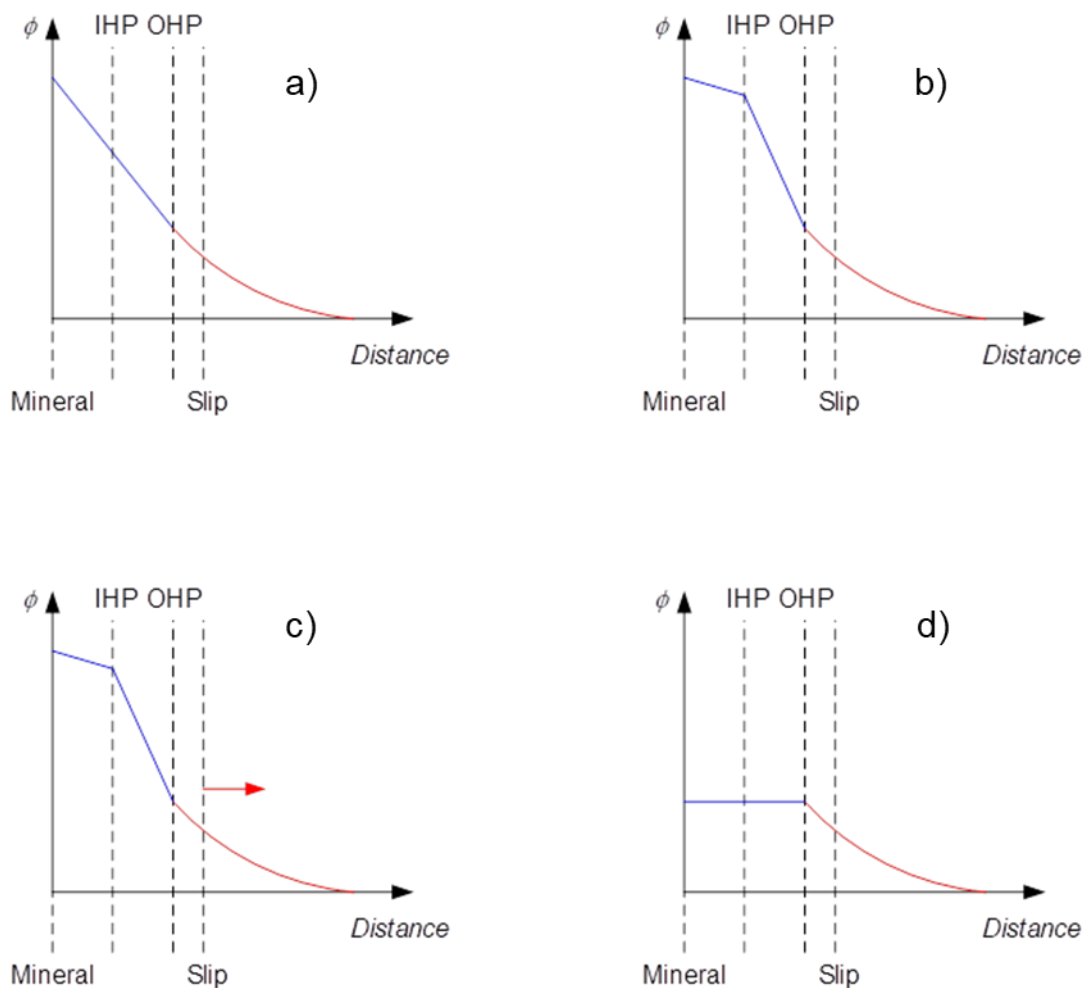
33 Calcite is a very common mineral on the Earth's surface comprising approximately 4% by weight of
34 Earth's crust. Physico-chemical processes that take place in porous calcite-aqueous solution (water
35 for simplicity) systems are of great importance for a variety of scientific and technological areas.
36 Properties of the calcite mineral surface and the interface between calcite and aqueous solution are
37 important for a broad range of applications including geological storage of nuclear waste and CO₂
38 [1], freshwater aquifer management [2,3], hydrocarbon production [4], paper production [5].
39 However, electrochemical properties of the interfaces between calcite and aqueous solutions
40 (specifically, electrical surface charge and zeta potential), especially under conditions relevant to
41 natural subsurface systems (i.e., water chemical composition, temperature, pressure) remain poorly
42 understood because of their microscopic nature, the high reactivity of the calcite mineral, and the
43 lack of relevant measurements and models [6].

44 The zeta potential is defined as the electrical potential at the shear plane when water flow occurs
45 parallel to the pore surface [7]. It is most commonly used to estimate the electrochemical properties
46 of calcite because acid-base potentiometric titration, usually employed to measure the surface
47 charge of minerals, is extremely difficult to carry out for highly reactive calcite [8]. There are many
48 studies that have reported measurement of the zeta potential of various calcite-water systems.
49 However, many reported results are contradictory (see for example Fig. 3 in Jackson et al. [9]). Most
50 of the experimental studies have reported zeta potential measured with low salinity single-salt
51 solutions (e.g., [10,11]), and there is inconsistency in the experimental conditions when some
52 experiments were conducted at equilibrium between the mineral and water while others reported the
53 zeta potential with non-equilibrium solutions (see for example discussion in Section 3 and 4 Al
54 Mahrouqi et al. [12]). The experimental studies have also employed a variety of electrokinetic
55 methods including the electrophoretic mobility (EPM [13]) and streaming potential method (SPM;
56 [14]). Each of these experimental techniques suffers from several limitations [15]. The EPM applies
57 an electrical field (of magnitude hundred Volts per meter) that mobilizes charged particles relative to
58 aqueous solution and measuring the resulting mobility. The EPM is conducted on powdered calcite
59 samples suspended in aqueous solution thus potentially exposing fresh mineral surfaces. Moreover,
60 the method is usually limited to solution concentration below 1 M ($M = \text{mol/L}$) [16]. The ratio of the
61 mineral to the solution is also very low (typically few grams per liter) and not representative of real
62 rocks. On the other hand, the SPM applies a pressure difference across the sample and measures
63 a resulting voltage, it can be carried out under equilibrium conditions, on intact rock samples at
64 salinities up to full saturation (e.g., [17,18]). However, unlike the EPM method, SPM can be time
65 consuming and very challenging for highly reactive minerals such as calcite in highly saline
66 conditions thus limiting its ability to cover a wide range of rock-solution permutations (see for
67 example typical duration of both the equilibration process and individual SPM experiment, as well as
68 the voltage stability issues at high salinity reported in Alroudan et al. [19]). Moreover, the zeta
69 potential values obtained with SPM can be specific to rock type as different carbonate samples may
70 contain different impurities such as anhydrite and dolomite (e.g., [12]).

71 A model capable of predicting the zeta potential of calcite-water systems at equilibrium as well as
72 non-equilibrium conditions, low to high salinity, and complex solution composition is crucially
73 important. Such model will not only be capable of predicting the zeta potential, but also improve our
74 understanding of the complex electrochemical processes that take place at calcite-water interface.
75 To date, there have been numerous attempts to develop such model with most of published studies
76 using the so-called surface complexation modelling (SCM) approach (e.g., [6,10,11]). In this method
77 the calcite surface is presented as an ensemble of chemically active surface sites that interact with
78 ions from the adjacent bulk electrolyte. The strength of these interactions is described by the
79 equilibrium constants (similar to chemical reactions), so that the resulting equilibrium concentration
80 of positive and negative surface complexes establishes the net surface charge and the
81 corresponding surface potential [20]. Depending on the complexity of the aqueous solution, such
82 models can be realized analytically (e.g., [21]) or by using numerical methods already available in
83 software packages such as PHREEQC (e.g., [22]).

84 Studies published to date on SCM have used different approaches to model the zeta potential
85 (**Figure 1**): Basic Stern Model (BSM, e.g., [10]), Triple Layer Model (TLM, e.g., [23]), Quad-Layer
86 Model (QLM, e.g. [24]) and Diffuse Layer Model (DLM, e.g. [25]). BSM, TLM and QLM explicitly
87 describe the Stern layer and a layered arrangement of ionic species adsorbed on the mineral surface
88 in the diffuse layer. Both the TLM and QLM distinguish between the Inner Helmholtz plane (IHP) and
89 the Outer Helmholtz plane (OHP), at which adsorption of de-hydrated and hydrated ions,
90 respectively, takes place [23]. On the other hand, BSM for calcite assumes that adsorption of all ions
91 in compact layer takes place at the OHP (e.g., [6,10,11]). All three models that explicitly describe the

92 adsorption layers (i.e., BSM, TLM, QLM) also assume that only protonation/deprotonation of
 93 hydrated calcite lattice ions take place at the mineral surface. In these models, the surface charge
 94 densities are computed at each plane, from which linear, capacitor-like variation of the electric
 95 potential between the planes is obtained. Consequently, TLM and QLM simulate separately
 96 protonation/deprotonation and salt ion adsorption reactions at three different planes (mineral
 97 surface, IHP, OHP), and ascribe two capacitance values to layers confined between the mineral
 98 surface and IHP, and between IHP and OHP. In addition, the QLM considers a stagnant diffuse
 99 layer, implying that the shear plane is located further away from the OHP. This makes these two
 100 models more computationally expensive compared with BSM, which uses only one capacitance and
 101 all adsorption reactions are considered to take place at OHP for calcite.



102

103 **Figure 1.** Electrical potential distribution at the calcite-water interface: a) – BSM (no distinction between IHP
 104 and OHP, the slip plane may or may not coincide with OHP); b) – TLM (IHP and OHP are considered separately
 105 with different surface complexation reactions taking place at each plane, the slip plane coincides with OHP);
 106 c) – QLM (IHP and OHP are considered separately with different surface complexation reactions taking place
 107 at each plane, the slip plane does not coincide with OHP and its distance from OHP can be constant or varying
 108 with salinity); d) – DLM (IHP and OHP are not considered at all as if there is no separation between the mineral
 109 surface and OHP, the slip plane may or may not coincide with mineral/OHP).

110 On the other hand, DLM assumes that both the protonation/deprotonation and salt ion adsorption
 111 reactions take place at the mineral surface thus ignoring the complexity of the Stern layer that
 112 comprises the IHP and OHP. Hence, compared to the BSM, TLM, and QLM, the DLM is not sensitive
 113 to the capacitance values and is only sensitive to the location of the so-called shear (or slip) plane
 114 at which ions can be mobilized by the flow and at which the electric potential is defined as the zeta

115 potential. According to the generally agreed theory of the electrical double layer, concentrations of
116 counter-ions that populate the region beyond the OHP (in the diffuse layer) obey the Boltzmann
117 distribution law where they decrease exponentially towards concentrations of ions in electroneutral
118 bulk electrolyte (outside the diffuse layer). The location of the slip plane is uncertain and is usually
119 used as a fitting parameter to match the model predictions to experimental data [10]. The assumed
120 location of the slip plane can also be used in BSM, TLM and QLM as additional fitting parameter, but
121 in contrast with DLM the modelled zeta potential using these models depends on adjustable
122 capacitance values and therefore, the location of the slip plane can be kept constant for matching
123 different experimental datasets. The QLM published by Alizadeh and Wang [24] assumed that the
124 slip plane was dynamic and moved further away from the mineral surface as the ionic strength of the
125 bulk electrolyte decreased. Although the dynamic slip plane model successfully reproduced
126 experimental results, the hypothesis was not clearly justified by the authors or confirmed
127 experimentally.

128 However, despite the plethora of published SCM studies, all of the simulated results appear to be
129 specific to an experimental dataset. Heberling and co-authors [6,10] used fractional charges of
130 individual surface site, the approach that was consistent with that taken by Wolthers et al. [23], who
131 used the Charge Distribution MultiSite Ion Complexation or CD-MUSIC modeling approach.
132 However, the magnitude of the surface site charges was different. On the other hand, Van Cappellen
133 et al. [26] used integer surface site charges in their BSM. Furthermore, Heberling et al. [6,10] used
134 equilibrium constants for adsorption reactions at OHP limited by ion-binding ones that corresponded
135 to pairing of ions in the bulk electrolyte. In contrast, Pokrovsky and Schott [27] estimated their
136 equilibrium constants for adsorption reactions from the correlation between stability of surface sites
137 and stability of the same molecules in the solution. As a result, the equilibrium constants used by
138 Pokrovsky and Schott [27] in their BSM exceeded by several orders of magnitude those used by
139 Heberling et al. [6,10] in their BSM. Considering the equilibrium constants of
140 protonation/deprotonation reactions that take place at the mineral surface, a significant variation in
141 their values was also evident across different studies. For instance, Heberling et al. [10] used five-
142 fold larger equilibrium constant for deprotonation of hydrated Ca site compared with their later study
143 [6]. At the same time, Wolthers et al. [23] used twelve orders of magnitude larger equilibrium constant
144 for the same reaction compared with Heberling et al. [10].

145 Previously published studies that used either BSM or TLM relied on constant Stern layer
146 capacitance, but the reported values used in these studies varied between 0.45 F/m², 1.24 F/m² [11]
147 , 17 F/m² [27], 52 F/m² [28] and 100 F/m² [23]. Therefore, the inconsistency in these values suggests
148 a high degree of freedom and unconstrained variation of simulated zeta potential. Moreover, most
149 of the published SCM simulated the zeta potential of calcite in contact with low salinity (typically less
150 than 0.1 M) single salt solutions (e.g., NaCl) therefore ignoring the effect of divalent ions such as
151 Mg²⁺ and SO₄²⁻ and high ionic strength. Studies that included such reactions reported inconsistent
152 equilibrium constants, with Heberling et al. [6] predicting adsorption of Ca²⁺ on negative surface sites
153 to be controlled by an equilibrium constant ten orders of magnitude smaller compared with the
154 equilibrium constant calculated from two consecutive reactions described by Pokrovsky et al. [28]
155 and Pokrovsky and Schott [29]. All in all, a unique combination of adsorption reactions, their
156 equilibrium constants, capacitance values and charges of the surface sites used in published studies
157 appear to be very specific to the modelled experimental datasets, thus limiting the use of each model
158 for a very narrow range of ionic strength, pH and composition of electrolytes in contact with the
159 calcite mineral.

160 Therefore, the aim of this study is to develop and validate a new SCM that accurately computes the
161 zeta potential of any calcite-water system as long as the experimental conditions are reproduced in
162 the model. The developed model has proven to be accurate in simulating and predicting zeta

163 potentials at both equilibrium and non-equilibrium conditions, high to low salinity, for any carbonate
 164 rock type as long as the dominating mineral is calcite, and for any solution composition as long as
 165 the dominating salt is NaCl and concentration of SO_4^{2-} does not exceed that of Ca^{2+} and Mg^{2+} .

166 2. Methodology

167 To develop a robust predictive model requires detailed description of experimental parameters and
 168 conditions. Therefore, we have chosen the data reported by Al Mahrouqi et al. [12] as the most
 169 comprehensive experimental work and will briefly describe the experimental conditions and main
 170 conclusions.

171 2.1. Description of the experimental data on zeta potential in intact limestone samples

172 Streaming potential measurements were performed by Al Mahrouqi et al. [12] and Alroudhan et al.
 173 [19] on three different carbonate rock samples: Estailades [12], Ketton [12] and Portland [19]. The
 174 petrophysical and mineralogical properties of the rock samples are shown in **Table 1**.

Property/rock	Ketton	Estailades	Portland
Porosity (%)	23.0 ± 0.5	28.0 ± 0.5	20.0 ± 0.5
Permeability (Darcy)	1.4 ± 0.4	0.13 ± 0.2	0.005 ± 0.001
Formation factor (F)	13.87 ± 0.5	12.92 ± 0.5	22.04 ± 0.5
Composition (%)	97 ^a – calcite	97 ^a (95 ^b) – calcite	96.6 ^a – calcite
	3 ^a magnesium	3 ^a (4 ^b) – magnesium	3.4 ^a – quartz
		(1 ^b) – anhydrite	

175 **Table 1.** Petrophysical and mineralogical properties of three different carbonate rock samples. Ketton and
 176 Estailades were used in the experiments of by Al Mahrouqi et al. [12], Portland was used by Alroudhan et al.
 177 [19].

178 **a** – mineralogy reported by Al Mahrouqi et al. [12] using X-ray diffraction (XRD). Magnesium was reported to
 179 be likely incorporated into the limestone as dolomite.

180 **b** – recent study by Udoh and Vinogradov [30], identified and confirmed that there was 4% dolomite
 181 ($\text{CaMg}(\text{CO}_3)_2$) and 1% anhydrite (CaSO_4) in the Estailades sample using XRD.

182

183 The zeta potential measurements were conducted at room temperature (23°C) with NaCl solutions
 184 of varying concentration between 0.05 M (mol/L) and 5.0 M NaCl at equilibrium conditions. Effluent
 185 fluid samples were regularly collected for chemical analyses to determine the concentration of Na^+ ,
 186 Ca^{2+} , Mg^{2+} , Cl^- , and SO_4^{2-} . The experimentally confirmed presence of Ca^{2+} , Mg^{2+} , and SO_4^{2-} was
 187 explained by partial dissolution of rock samples during the equilibration with NaCl solutions and
 188 leaching of these ions.

189 These studies concluded that the zeta potential of carbonate rock samples was controlled by the
 190 concentration of the potential determining ions (PDIs), which were identified to be Ca^{2+} , Mg^{2+} , and
 191 SO_4^{2-} .

192 2.2. Calcite surface complexation model development

193 2.2.1. Basic definitions and assumptions

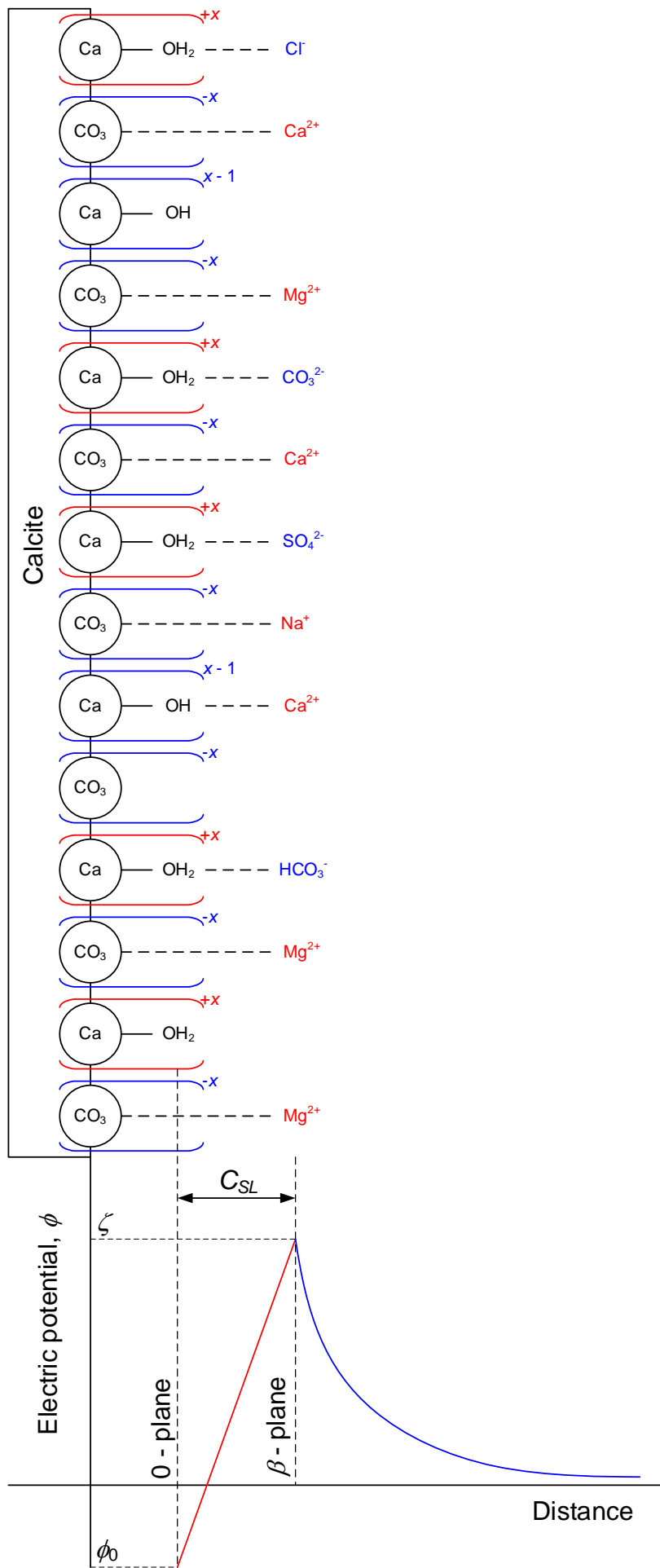
194 A BSM was used in this study to model the zeta potential of pure calcite in contact with aqueous
 195 solutions. The modelling approach was consistent with the most comprehensive and empirically
 196 justified model developed by Heberling et al. [6,10]. Since the content of calcite in all three tested
 197 carbonate rock samples (Estailades, Ketton, Portland) was found to be greater than 95%, it was

198 assumed that calcite was the most dominant mineral in the samples so that only calcite surface sites
199 were considered in the model.

200 The measured zeta potential with all tested carbonate rocks reflected presence of minerals other
201 than calcite (e.g., dolomite and anhydrite), which led to a non-zero concentration of Mg^{2+} and SO_4^{2-}
202 found in the bulk solution as a result of mineral dissolution during the equilibration process.
203 Therefore, we artificially added the equilibrium concentration of Mg^{2+} and SO_4^{2-} measured in the
204 experiment to our geochemical model of pure calcite. The amount of Ca^{2+} leached into NaCl
205 solutions during the equilibration was either added artificially to the model and equilibration was
206 switched off, or it was not added to the initial solution while the equilibration was activated (see
207 Section 2.3 for details).

208 Consistent with Heberling et al. [6,10], calcite (1 0 4) face was assumed to dominate the interface
209 due to its high abundance and the total surface site density of calcium and carbonate surface sites
210 on the calcite (1 0 4) face was set to 4.95 sites per nm^2 for each type of site. However, unlike
211 Heberling et al. [6], this study assumed that the zeta potential could be calculated by considering the
212 shear plane coinciding with the OHP (see **Figure 2**), i.e., $\phi_\beta = \phi_d = \zeta$ – the assumption that was
213 also used by Li et al. [11] (no stagnant diffuse layer). Li et al. [11] considered that the assumption of
214 the presence of a stagnant diffuse layer in Heberling et al. [10] was due to the nature of electrokinetic
215 measurements that were not corrected for surface conductivity effects thus reducing the
216 experimental zeta potentials and then necessitating smaller modelled zeta potentials to be further
217 away from the mineral surface, to match the data.

218 The 0-plane was defined as the hydrolysis layer located 2.3 Å from the calcite surface defined by
219 the surface of calcium ions, at which only protonation/deprotonation of surface sites took place [6].
220 The surface charge at 0-plane was found to be negative regardless of the concentration and
221 composition of the adjacent solution and pH range of 5-11 tested in our study, similar to the results
222 reported by Heberling et al. [6]. On the other hand, the β -plane was set at a distance of 4.6 Å from
223 the surface [6], and we assumed that all the specific adsorption reactions between the surface sites
224 and salt ions from the bulk electrolyte only occurred at the OHP where ions are adsorbed as outer-
225 sphere complexes.



227 **Figure 2.** Illustration of the Basic Stern Model used in this study. The 0-plane corresponds to the mineral
 228 surface where only protonation/deprotonation reactions take place. We used fractional surface charge $x = 0.5$
 229 for all surface sites. The Stern Layer is confined between the 0-plane and β -plane, with the latter corresponding
 230 to the OHP where all adsorption reactions take place and where the zeta potential (ζ) is defined. C_{SL} is the
 231 Stern layer capacitance.

232

233 2.2.2. Adsorption reactions and equilibrium constants

234 All modelled adsorption reactions between calcite surface sites and ions are listed in **Table 2**. The
 235 original set of adsorption reactions proposed by Heberling et al. [6,10] and Li et al. [11] was extended
 236 to include the reactions between the surface sites and Mg^{2+} and SO_4^{2-} (R9 – R11 in **Table 2**). The
 237 equilibrium constant of R1 that corresponds to deprotonation of hydrated calcium surface sites was
 238 fixed to the value of $\text{Log}K_1 = -0.5$ to yield the relative abundance of positive and negative surface
 239 complexes at 0-plane according to Heberling et al. [6].

240 Among published studies on SCM of calcite, there is no consensus on whether protonation of
 241 $>CO_3^{-0.5}$ surface sites on 0-plane should be considered. Some of these studies included the reaction
 242 $>CO_3^{-0.5} + H^+ \rightleftharpoons CO_3H^{+0.5}$ (e.g., [10,23,27]), while other studies (e.g., [6,11]) disregarded it. We
 243 could not find any published justification for inclusion of $>CO_3^{-0.5}$ protonation based on the
 244 experimental data or theoretical analysis. However, there have been several publications in support
 245 of excluding the protonation of the $>CO_3^{-0.5}$ surface sites. One theoretical study on hydration of (1 0
 246 4) calcite surface [31] demonstrated that neutral H_2O molecules preferentially ‘chemisorbed’ into $>$
 247 $Ca^{+0.5}$ to form $>CaOH_2^{+0.5}$ site, and existence of $>CO_3H^{+0.5}$ surface sites was attributed to hydration
 248 of carbonate surface groups ($>CO_3OH_2^{-0.5}$) but not direct protonation of the $>CO_3^{-0.5}$ sites.
 249 Therefore, in the absence of $>CO_3OH_2^{-0.5}$ there could be no $>CO_3H^{+0.5}$ sites resulting from
 250 subsequent dissociation of water molecules $>CO_3OH_2^{-0.5} \rightleftharpoons CO_3H^{+0.5} + OH^-$. These conclusions
 251 were supported by a variety of experimental measurements reported in the paper. Another numerical
 252 study [32] also predicted low fraction of protonated $>CO_3H^{+0.5}$ sites (<20%) on a variety of calcite
 253 mineral geometries for pH between 5 and 11, which is consistent with pH values tested with our
 254 model. For pH greater than 7, the fraction of protonated carbonate sites decreased to below 5%
 255 according to this study. Moreover, inclusion of the $>CO_3^{-0.5}$ surface site protonation would result in
 256 increased pH under simulated equilibrium conditions, thus mismatching the experimentally
 257 measured values. Effect of $>CO_3^{-0.5}$ protonation on equilibrium pH will be discussed in more detail
 258 in Section 3. For these reasons and following Heberling et al., [6] we did not consider hydration of
 259 carbonate surface sites in our model.

No	Reactions	$\text{Log}K^{\min}$	$\text{Log}K^{\max}$	$\text{Log}K^{\text{opt}}$
0-plane reactions				
R1	$>CaOH_2^{+0.5} \rightleftharpoons CaOH^{-0.5} + H^+$	-0.5	-0.5	-0.5
β -plane reactions				
R2	$>CaOH^{-0.5} + Na^+ \rightleftharpoons CaOH^{-0.5} - Na^+$	0.09 ^a	0.56 ^b	0.56
R3	$>CaOH_2^{+0.5} + Cl^- \rightleftharpoons CaOH_2^{+0.5} - Cl^-$	-2.10 ^c	0.45 ^d	-0.64
R4	$>CaOH^{-0.5} + Ca^{2+} \rightleftharpoons CaOH^{-0.5} - Ca^{2+}$	1.68 ^b	3.40 ^e	3.40

R5	$> \text{CaOH}_2^{+0.5} + \text{HCO}_3^- \Leftrightarrow \text{CaOH}_2^{+0.5} - \text{HCO}_3^-$	0.04 ^b	12.50 ^f	10.65
R6	$> \text{CaOH}_2^{+0.5} + \text{CO}_3^{2-} \Leftrightarrow \text{CaOH}_2^{+0.5} - \text{CO}_3^{2-}$	-7.07 ^b	6.00 ^g	-4.59
R7	$> \text{CO}_3^{-0.5} + \text{Ca}^{2+} \Leftrightarrow \text{CO}_3^{-0.5} - \text{Ca}^{2+}$	1.68 ^b	3.40 ^h	3.40
R8	$> \text{CO}_3^{-0.5} + \text{Na}^+ \Leftrightarrow \text{CO}_3^{-0.5} - \text{Na}^+$	0.56 ^b	3.40 ⁱ	0.56
R9	$> \text{CaOH}^{-0.5} + \text{Mg}^{2+} \Leftrightarrow \text{CaOH}^{-0.5} - \text{Mg}^{2+}$	1.66 ^a	3.40 ^j	2.81
R10	$> \text{CaOH}_2^{+0.5} + \text{SO}_4^{2-} \Leftrightarrow \text{CaOH}_2^{+0.5} - \text{SO}_4^{2-}$	-2.10 ^g	3.30 ^k	3.30 → 11.9
R11	$> \text{CO}_3^{-0.5} + \text{Mg}^{2+} \Leftrightarrow \text{CO}_3^{-0.5} - \text{Mg}^{2+}$	1.68 ^l	3.40 ^l	2.81

260 **Table 2.** Adsorption reactions and their equilibrium constants. Equilibrium constant for R1 was not optimised.
261 **a** – Song et al. [33].
262 **b** – Heberling et al. [6].
263 **c** – only one value for the equilibrium constant for R3 was found in the literature (0.45), hence we considered
264 $\text{Log}K^{\text{min}}$ to be equal to that of R10 due to similarity.
265 **d** – Li et al. [11].
266 **e** – only one value for the equilibrium constant for R8 was found in the literature (1.68), hence we considered
267 $\text{Log}K^{\text{max}}$ to be equal to that of R7 due to similarity.
268 **f** – combined from two reactions reported in Pokrovsky and Schott [29].
269 **g** – Qiao et al. [34].
270 **h** – combined from two reactions reported in Pokrovsky et al. [28].
271 **i** – only one value for the equilibrium constant for R8 was found in the literature (0.56), hence we considered
272 $\text{Log}K^{\text{max}}$ to be equal to that of R7 due to similarity.
273 **j** – only one value for the equilibrium constant for R9 was found in the literature (1.66), hence we considered
274 $\text{Log}K^{\text{max}}$ to be equal to that of R4 due to similarity.
275 **k** – Qiao et al. [35].
276 **l** – this is a new additional reaction introduced in our model with no values for the equilibrium constant found
277 in the literature, hence we assumed both $\text{Log}K^{\text{min}}$ and $\text{Log}K^{\text{max}}$ to be equal to those of R7 due to similarity.

278
279 For reactions R2 – R11 we identified ranges of possible variation of equilibrium constants. The
280 ranges were based on previously published values thus defining the minimum and maximum
281 possible equilibrium constant for each reaction, $\text{Log}K^{\text{min}}$ and $\text{Log}K^{\text{max}}$, respectively. Note, that due to
282 lack of published information on some equilibrium constants, their maximum and minimum values
283 were assumed to be similar to those of other adsorption reactions or calculated from published
284 values of consecutive reactions (see caption of **Table 2** for details).

285 We used the Pitzer theory (pitzer.dat database of PREEQC [36,37]) to calculate the ion activity
286 coefficients for all complexation reactions that take place in the bulk solution at high ionic strength
287 systems (> 0.5 M). For ionic strengths below 0.5 M it was found that there was no noticeable
288 difference in the modelled results obtained using either the Debye-Huckel (phreeqc.dat database of
289 PHREEQC) or Pitzer theory to calculate the ion activity coefficients. Therefore, the Pitzer theory was
290 used throughout the entire range of tested ionic strengths.

291 2.2.3. BSM – capacitance of the Stern layer and electric potential distribution within EDL

292 The electric potential distribution within the Stern layer, between 0-plane and β -plane, is described
293 by a linear variation, similar to the concept of two parallel plates capacitor which it can be expressed
294 as follows:

$$\sigma_0 = C_{SL} (\phi_0 - \phi_\beta), \quad (1)$$

295 where σ_0 is the surface charge density at 0-plane ($\text{C}\cdot\text{m}^{-2}$), C_{SL} is the specific Stern layer capacitance
 296 (or Stern layer capacitance for simplicity) between 0-plane and β -plane ($\text{F}\cdot\text{m}^{-2}$), ϕ_0 is the electric
 297 potential at 0-plane (V) and ϕ_β is the electric potential at β -plane (V). The Stern layer capacitance
 298 can also be expressed as:

$$C_{SL} = \frac{\varepsilon_0 \varepsilon_r}{x}, \quad (2)$$

299 where ε_0 is vacuum permittivity ($\text{F}\cdot\text{m}^{-1}$), ε_r is relative permittivity of the Stern layer and x is the
 300 distance between 0-plane and β -plane or the thickness of the Stern layer (m). Beyond the β -plane,
 301 in the diffuse layer, the electrical potential distribution is described using the Gouy-Chapman theory
 302 based on the Poisson-Boltzmann equation. The electrical potential magnitude decreases
 303 exponentially with the distance from the mineral surface and ion concentrations follow a Boltzmann
 304 distribution.

305 2.2.4. Two-step optimization

306 Our model was developed through a two-step optimization process applied to experimental zeta
 307 potential results obtained with Estailades and Ketton rock samples saturated with equilibrated NaCl
 308 solutions of ionic strengths between 0.05 M and 5 M.

309 The first-step optimization aimed at determining the equilibrium constants of R2-R11, while the
 310 $\log K_{R1}$ and the Stern layer capacitance (C_{SL}) were kept constant and equal to -0.5 and 1.24 F/m^2 ,
 311 respectively, in accordance with the study of Li et al. [11]. The optimization of equilibrium constants
 312 of the surface complexation reactions and Stern layer capacitance was a necessary step, as
 313 previous studies considered either low salinity (e.g., [11]) or moderate salinity solutions (e.g.,
 314 seawater model by Song et al. [33]) using different sets of the parameters for each ionic strength,
 315 thus making their models suitable to very specific experimental conditions. To the best of our
 316 knowledge there has been no study that tried SCM for a range of salinities, and especially for ionic
 317 strengths above 0.5 M which are typical for many subsurface settings. The first-step optimization
 318 was conducted in the following manner:

- 319 • Concentration of all ions (Na^+ , Ca^{2+} , Mg^{2+} , Cl^- and SO_4^{2-}) and pH reported by Al Mahrouqi et al.
 320 [12] were used as input parameters. The concentration of carbon related ions (HCO_3^- , CO_3^{2-}),
 321 termed C(4) ions in PHREEQC, was not reported in the corresponding paper, therefore it was
 322 calculated from the charge balance equation. It was also assumed that the calculated C(4)
 323 concentration could exceed that of Ca^{2+} and C(4) ions that originate from the dissolved
 324 atmospheric CO_2 in aqueous solution at equilibrium with air. This assumption is consistent with
 325 the experimentally confirmed dissolution of CaCO_3 during the equilibration process that
 326 resulted in 10^{-3} M of dissolved Ca^{2+} , and hence the same concentration of C(4) [12,19], which
 327 is higher than the equilibrium concentration of $1.5 \times 10^{-5} \text{ M}$ of C(4) in solution at equilibrium with
 328 air [11].
- 329 • The dissolution and precipitation of calcite was switched off at this optimization step by
 330 implementing PHREEQC code 'Calcite 0 0' in 'equilibrium_phases' section [6,22].
- 331 • The R2 – R11 equilibrium constants were optimized by minimizing the objective function, f ,
 332 that defines the normalized difference between the observed and the simulated zeta potential
 333 and pH:

$$f = \sum \left(\frac{\zeta_{obs} - \zeta_{sim}}{\delta_{\zeta}} \right)^2 + \sum \left(\frac{pH_{obs} - pH_{sim}}{\delta_{pH}} \right)^2, \quad (3)$$

334 where ζ_{obs} is the observed zeta potential (mV), ζ_{sim} is the simulated zeta potential (mV), δ_{ζ} is
 335 the experimental uncertainty of the zeta potential (mV), pH_{obs} is the observed pH, pH_{sim} is the
 336 simulated pH and δ_{pH} is the experimental uncertainty of pH. Note that pH was included in the
 337 objective function since the computed value for a given set of equilibrium constants could be
 338 different from the input (experimental) value. Therefore, both directly measured properties
 339 (zeta potential and pH) were included in **Eq. 3**. The optimization was conducted by combining
 340 the PHREEQC geochemical code with an optimization software, PEST, in which the Gauss
 341 Marquardt Levenberg method was implemented.

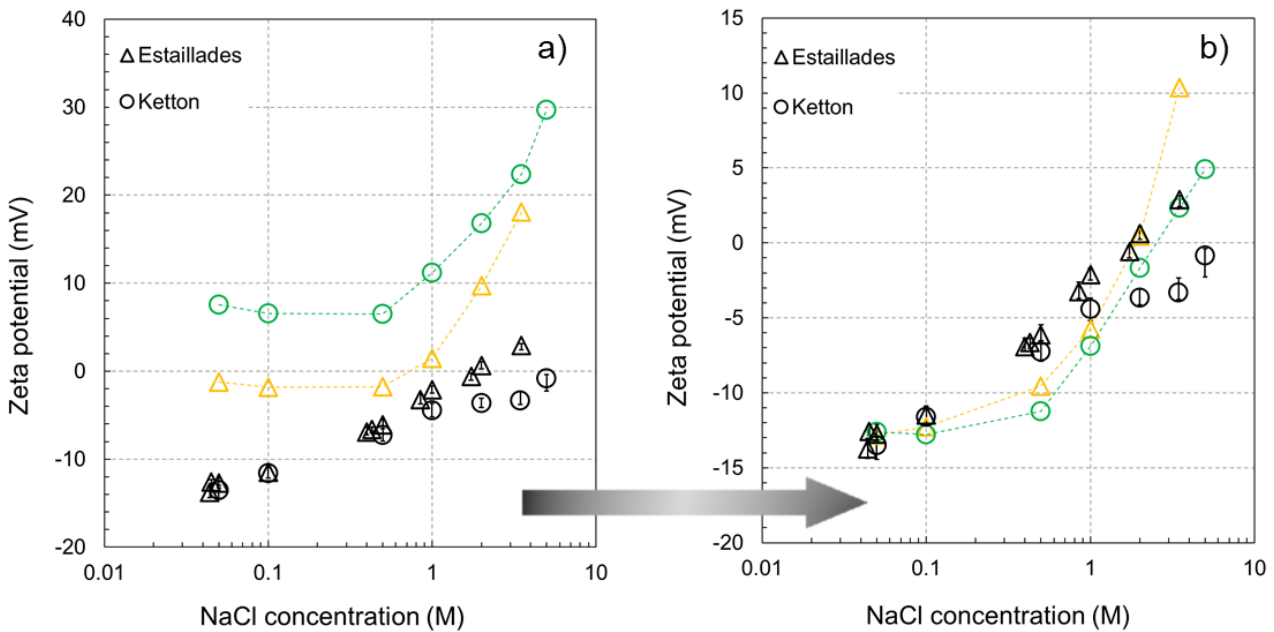
342 • The optimization was repeated three times, with the initial values of all equilibrium constants
 343 being either i) the minimum found in literature ($\text{Log}K^{\text{min}}$), ii) the maximum reported in the
 344 literature ($\text{Log}K^{\text{max}}$), or iii) the median within the tested range (see more details in **Table 2**
 345 caption). This repetitive optimization procedure was required to find the global minimum of f ,
 346 which corresponded to the best match between the experimentally measured and computed
 347 zeta potential and pH, so that the equilibrium constants that corresponded to the smallest f
 348 were taken forward.

349 • The results of the first step optimization procedure are shown in **Table 2** ($\text{Log}K^{\text{opt}}$ column).

350 It was found that the calculated zeta potentials of Estailades and Ketton were positive except for
 351 Estailades at ionic strength below 1 M. In contrast, the experimental results showed that the zeta
 352 potential of both rock samples remained negative at low ionic strength to become less
 353 negative/positive at high ionic strength (**Figure 3a**). Moreover, the calculated Ketton zeta potential
 354 was more positive compared with that of Estailades, while the experimental data showed an
 355 opposite trend. The more positive zeta potential measured in Estailades sample could not be
 356 explained by different rates of calcite dissolution as the equilibrium Ca^{2+} concentration was found to
 357 be higher in Ketton sample (Figure 9a in Al Mahrouqi et al. [12]), which would imply a more positive
 358 zeta potential, consistent with our model.

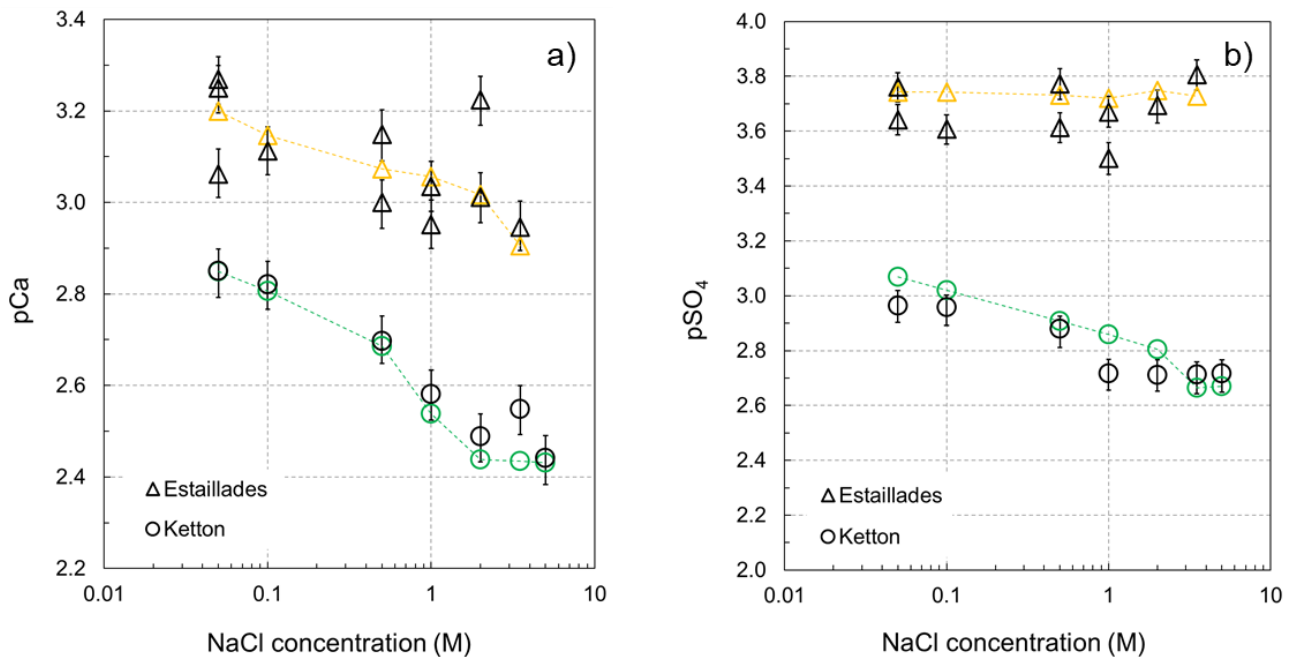
359 However, a considerably higher concentration of SO_4^{2-} was also reported in experiments with Ketton
 360 (Figure 9b in Al Mahrouqi et al. [12]), and more negative zeta potential compared with that of
 361 Estailades sample was attributed to it. The authors demonstrated that the zeta potential was not
 362 very sensitive to SO_4^{2-} concentration (Figure 8b in Al-Mahrouqi et al. [12]), as long as concentration
 363 of all other ions remained unchanged thus implying that high sensitivity to SO_4^{2-} only took place when
 364 sulfate content was less or equal to that of the divalent cations. However, the paper did not consider
 365 a possibility of Ca^{2+} ions acting as bridges to SO_4^{2-} ions [38–40] thus enhancing sulfate adsorption
 366 when elevated concentration of Ca^{2+} is found in bulk solution. Such bridging ability of Ca^{2+} implies
 367 that higher concentration of Ca^{2+} (and/or Mg^{2+}) combined with higher concentration of SO_4^{2-} (as was
 368 observed by Al Mahrouqi et al. [12]) should result in overall higher adsorption of sulfate and more
 369 negative zeta potential. Keeping in mind that dissolved SO_4^{2-} could only originate from undetected
 370 minerals such as anhydrite (CaSO_4) or epsomite ($\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$), a higher concentration of sulfate
 371 ions would always be accompanied by a higher equilibrium concentration of Ca^{2+} and/or Mg^{2+} as it
 372 was indeed reported by Al-Mahrouqi et al. [12]. Therefore, we assumed that the reactivity of calcite
 373 to SO_4^{2-} was considerably higher for rock samples that leached more sulfate and the constraint on
 374 $\text{Log}K^{\text{max}}$ for R10 was removed, which resulted in a substantially better match to the experimental
 375 data (**Figure 3b**) and the corresponding value of the optimized equilibrium constant, $\text{Log}K^{\text{opt}}$, of 11.9
 376 (Table 2). We expect that including Ca^{2+} – SO_4^{2-} ion bridging in the surface complexation reactions
 377 would lower $\text{Log}K^{\text{opt}}$. Such inclusion implemented in SCM, would require currently unavailable

378 experimental data to validate the model and therefore, data acquisition and the corresponding SCM
 379 adjustments will be conducted in a follow-up study.



380
 381 **Figure 3.** Calculated zeta potential after the first step optimization a) with the constraint $-2.10 \leq \log K_{R10} \leq$
 382 3.30 , b) without the constraint $-2.10 \leq \log K_{R10}$. Black symbols denote the experimental results from
 383 AlMahrouqi et al. [12].

384 The calculated concentration of Ca^{2+} and SO_4^{2-} (note, that input pH and concentration of Mg^{2+} were
 385 constant at 8.3 and $\text{pMg} = 4.19$, respectively, as reported by Al-Mahrouqi et al. [12], $\text{pMg} = -\log C_{\text{Mg}}$
 386 with C denoting the ion concentration in M) for both rock samples is shown in **Figure 4**. The
 387 computed solution pH and concentration of all other ions (those that are not shown in **Figure 4**)
 388 remained unchanged and equal to the input values regardless of $\text{Log}K^{\text{opt}}$ for R10, hence the
 389 corresponding plots for these ions are not included in the figure.



390
 391 **Figure 4.** Computed pCa (a) and pSO₄ (b) from the first optimization step. Black symbols represent the
 392 experimental data [12]. The results in color correspond to the modelled zeta potential that appears in **Figure**
 393 **3b**.

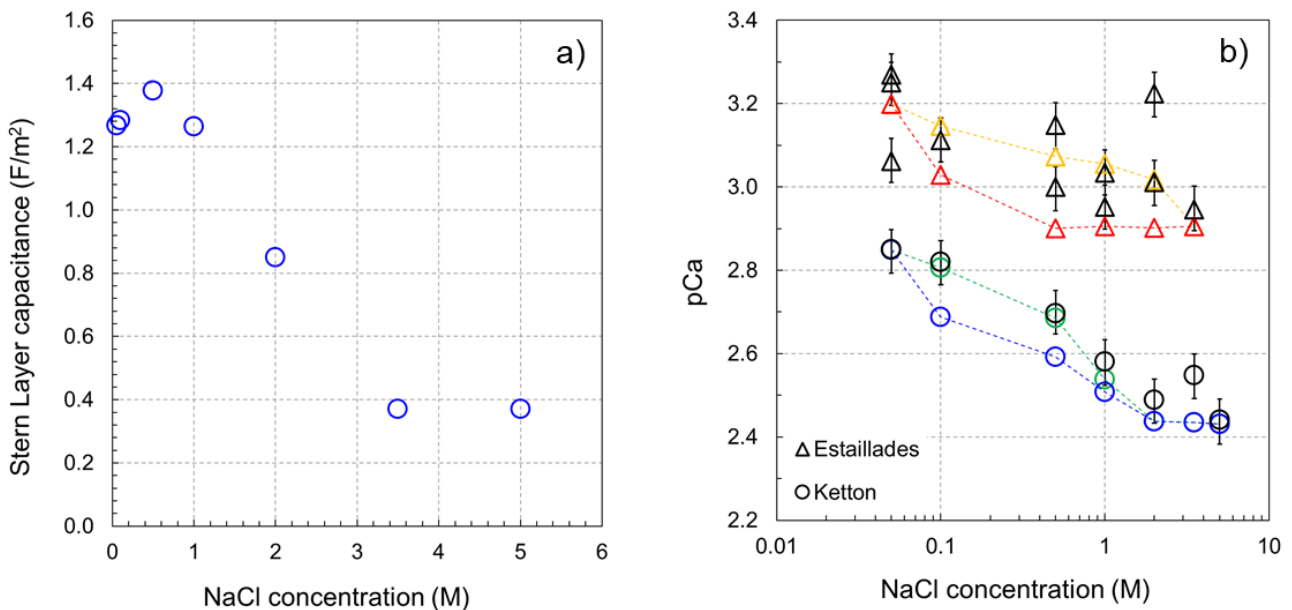
394

395 After increasing the $\text{Log}K^{\text{opt}}$ for R10 to the value of 11.9, the match to the experimental results
396 significantly improved as shown in **Figure 3b** for ionic strength ≤ 2 M. However, the simulated zeta
397 potential at ionic strength greater than 2 M remained significantly more positive compared with the
398 observed values, and for the Ketton sample it became positive as opposed to the negative zeta
399 potential obtained from the experiments. Therefore, additional modification to the model was made.

400 The second step optimization was initiated by fixing the optimal values of all equilibrium constants
401 obtained from the first step optimization. Then, using the same objective function as in the 1st
402 optimization step (**Eq. 3**), the 2nd step optimization was implemented for each rock sample and each
403 concentration of NaCl solution using the following variables:

- 404 • The concentration of Ca^{2+} and SO_4^{2-} was allowed to vary within the reported experimental
405 uncertainty.
- 406 • A variable capacitance of the Stern layer was assumed and allowed to vary between $0.2 \text{ F}\cdot\text{m}^{-2}$
407 2 and $1.4 \text{ F}\cdot\text{m}^{-2}$ (see more detailed discussion in section 2.2.5).

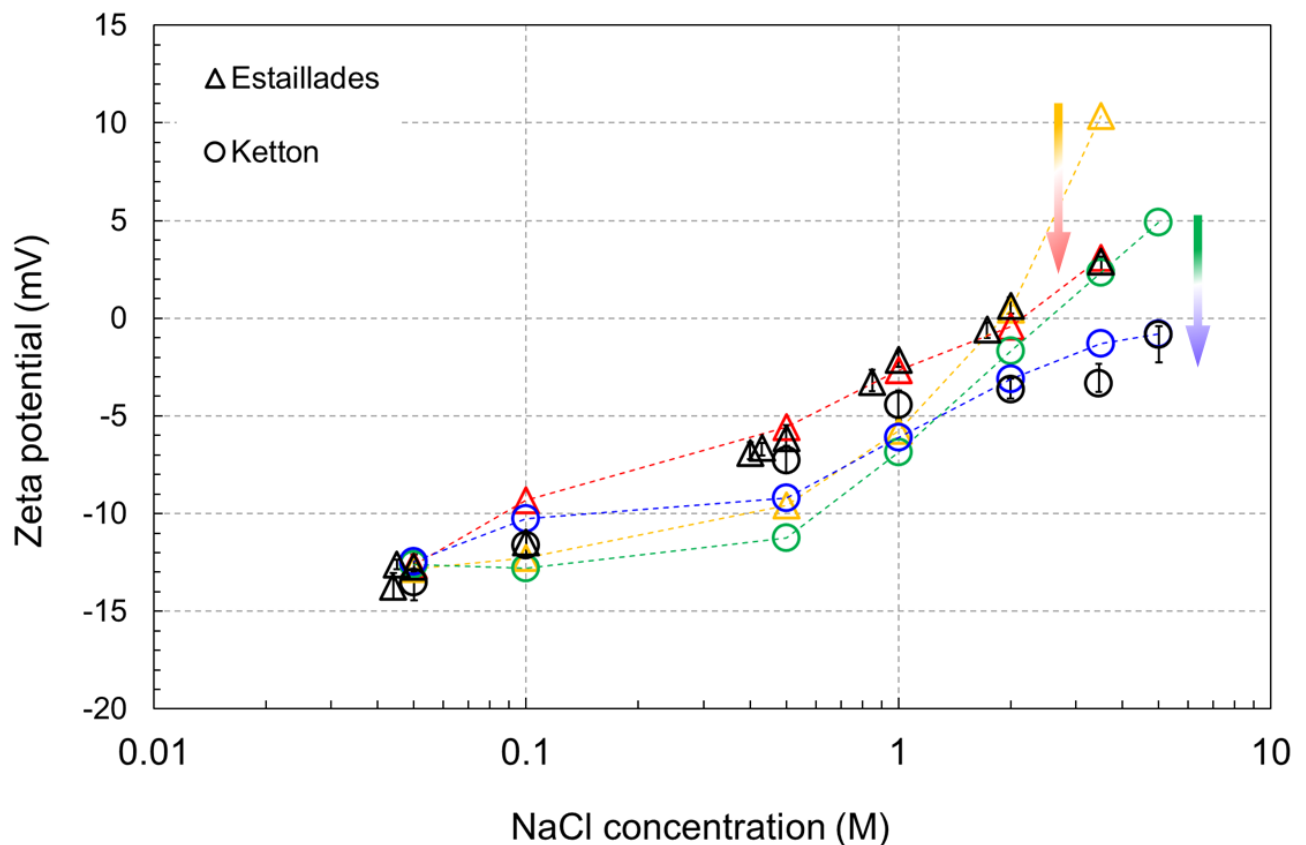
408 The resulting from the 2nd optimization step pH and concentration of SO_4^{2-} and Mg^{2+} remained the
409 same compared with the results of the 1st optimization step. However, to obtain a better match to the
410 experimentally measured zeta potential required a non-monotonic change of the optimized Stern
411 layer capacitance (**Figure 5a**) and increased concentration of Ca^{2+} (see the yellow \rightarrow red shift for
412 Estailades and green \rightarrow blue shift for Ketton in **Figure 5b**). The computed zeta potential was found
413 to be in a good agreement with the experimental data after the variable capacitance was
414 implemented (see the yellow \rightarrow red shift for Estailades and green \rightarrow blue shift for Ketton in **Figure 6**).



415

416 **Figure 5.** Optimized Stern layer capacitance (a) and pCa as a function of NaCl concentration. Black symbols
417 in b) correspond to the experiments of Al Mahrouqi et al., [12]. The yellow \rightarrow red and green \rightarrow blue shift represent
418 manual adjustment of pCa to obtain a better match to the computed zeta potential in **Fig. 6**.

419



420

421 **Figure 6.** Calculated zeta potential after the first and the second step optimization. Black symbols correspond
 422 to the experimental data of Al Mahrouqi et al. [12]. The yellow→red (Estailades) and green→blue (Ketton)
 423 shifts demonstrate a significant improvement of the match after applying variable Stern layer capacitance.

424

425 2.2.5. Variable Stern layer capacitance

426 As mentioned in the previous section, in order to obtain a good match with the experimentally
 427 measured zeta potential it was required to allow the Stern layer capacitance (C_{SL}) to vary and to
 428 decrease globally. Following the definition of C_{SL} (**Eq. 2**), its optimized concentration dependence
 429 (**Figure 5a**) can be explained by either: i) constant distance between the 0-plane and β -plane, x , and
 430 decreasing relative permittivity of the Stern layer, ϵ_r , with increasing NaCl concentration; ii) constant
 431 relative permittivity and increasing with salinity x ; iii) increasing with salinity x and decreasing ϵ_r ; or
 432 iv) decreasing x and ϵ_r with increasing salinity so that the rate of decrease of ϵ_r exceeds that of x .
 433 Previously published studies have suggested that hydration diameter of cations decreases
 434 considerably with increasing salinity, hence x might become smaller as smaller in diameter ions may
 435 have a closer distance of approach to the mineral surface (see for example, results obtained from
 436 an analytical model for Mg^{2+} in Afanas'ev and Ustinov [41] and for Na^+ in Afanasiev et al. [42] using
 437 the same approach, both verified by various experimental data) thus restricting the possible
 438 explanation for the decreasing C_{SL} to option iv). This assumption is also supported by Brown et al.
 439 [43] who suggested a substantial compression of the Stern layer with increasing electrolyte salinity,
 440 which was interpreted from experimental observations including EPM. In contrast, some molecular
 441 dynamics studies (e.g., [44]) suggest that there is no change in x with increasing salt concentration
 442 due to the presence of the hydrolysis layer. At the same time, an electric field within the Stern layer
 443 that is exerted on polar water molecules results in its polarization and may lead to a substantial
 444 decrease in ϵ_r with increasing salinity (e.g., [45]), consistent with options i), iii) and iv). In the light of
 445 the above arguments, we assumed option i) in our model, implying constant x and decreasing with
 446 salinity ϵ_r . Note, that we tested hypothesis iv) and found that C_{SL} is significantly less sensitive to the

447 variation of x (even assuming the maximum possible range between fully hydrated Na^+ radius of 4.5
 448 Å and crystallographic Na^+ radius of 0.5 Å at 5 M [46]) compared with allowed variation of ε_r (between
 449 the value that corresponds to diluted electrolyte, ~80 at a temperature of 20°C and that of structured
 450 water in the Stern layer, ~6 [10]), but this sensitivity analysis is not presented here.

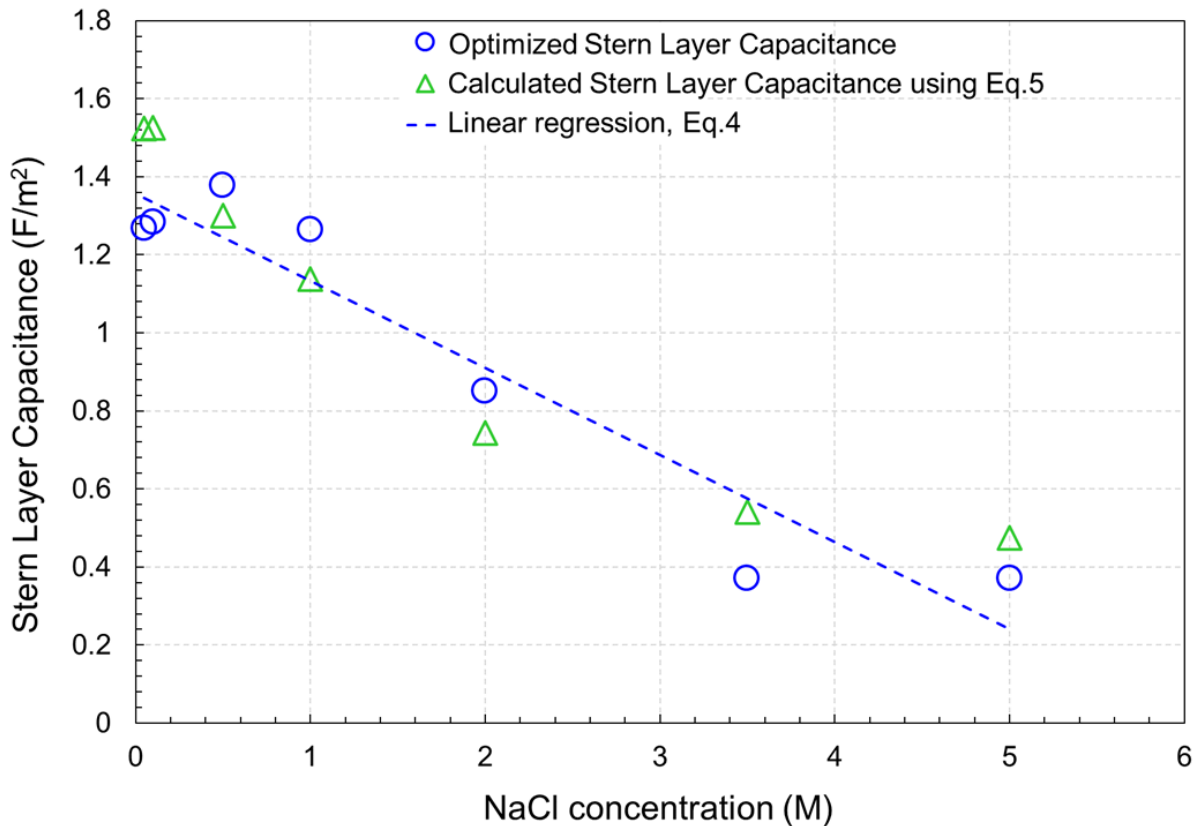
451 For simplicity we used a linear regression through optimized values of C_{SL} as shown in **Figure 7**:

$$C_{SL} = -0.2232 \times IS + 1.357 [F \cdot m^{-2}] \quad (4)$$

452 Where IS is the ionic strength of the solution of interest (M). To verify the regression is physically
 453 meaningful we assumed a constant distance between 0-plane and β -plane of $x = 2.3$ Å [6] and used
 454 a published equation for the Stern layer relative permittivity [47,48]:

$$\varepsilon_r = \frac{\varepsilon_z - \varepsilon_s}{1 + b \left(-\frac{d\phi}{x}\right)^2} + \varepsilon_s \quad (5)$$

455 where ε_z is the relative permittivity of the bulk electrolyte (at a given ionic strength and temperature),
 456 ε_s is the relative permittivity near the mineral surface (~6 [48]), $d\phi$ is the electrical potential difference
 457 (V) between 0-plane with β -plane calculated by our model, and $b = 1.2 \times 10^{-17} \text{ m} \cdot \text{V}^{-1}$ is a constant
 458 [48]. The values of C_{SL} calculated using ε_r computed from Eq. 5, and $x = 2.3$ Å are plotted in **Figure**
 459 **7** and are in good agreement with the optimized values thus confirming the validity of our approach.
 460 Note, that $d\phi$ for Eq.5 were computed by PHREEQC using the linear regression of C_{SL} (blue line in
 461 **Figure 7**) and provided an excellent match to the zeta potentials, so that non-linear regressions for
 462 C_{SL} were not tried as this was not the focus of this study.



463
 464 **Figure 7.** Optimized Stern layer capacitance compared with values obtained using **Eq. 5** and approximated
 465 by the linear regression fitted to the optimized values in blue (**Eq. 4**; quality of match to the optimized values
 466 of capacitance in blue is $R^2 = 0.91$) as a function of ionic strength.

467

468 2.3. Calcite surface complexation model implementation

469 To use our model for predicting the zeta potential of calcite in contact with aqueous solutions requires
470 identification of input parameters and model options that accurately replicated the reported
471 experimental conditions. Therefore, the following steps should be taken to obtain an accurate
472 prediction of the zeta potential:

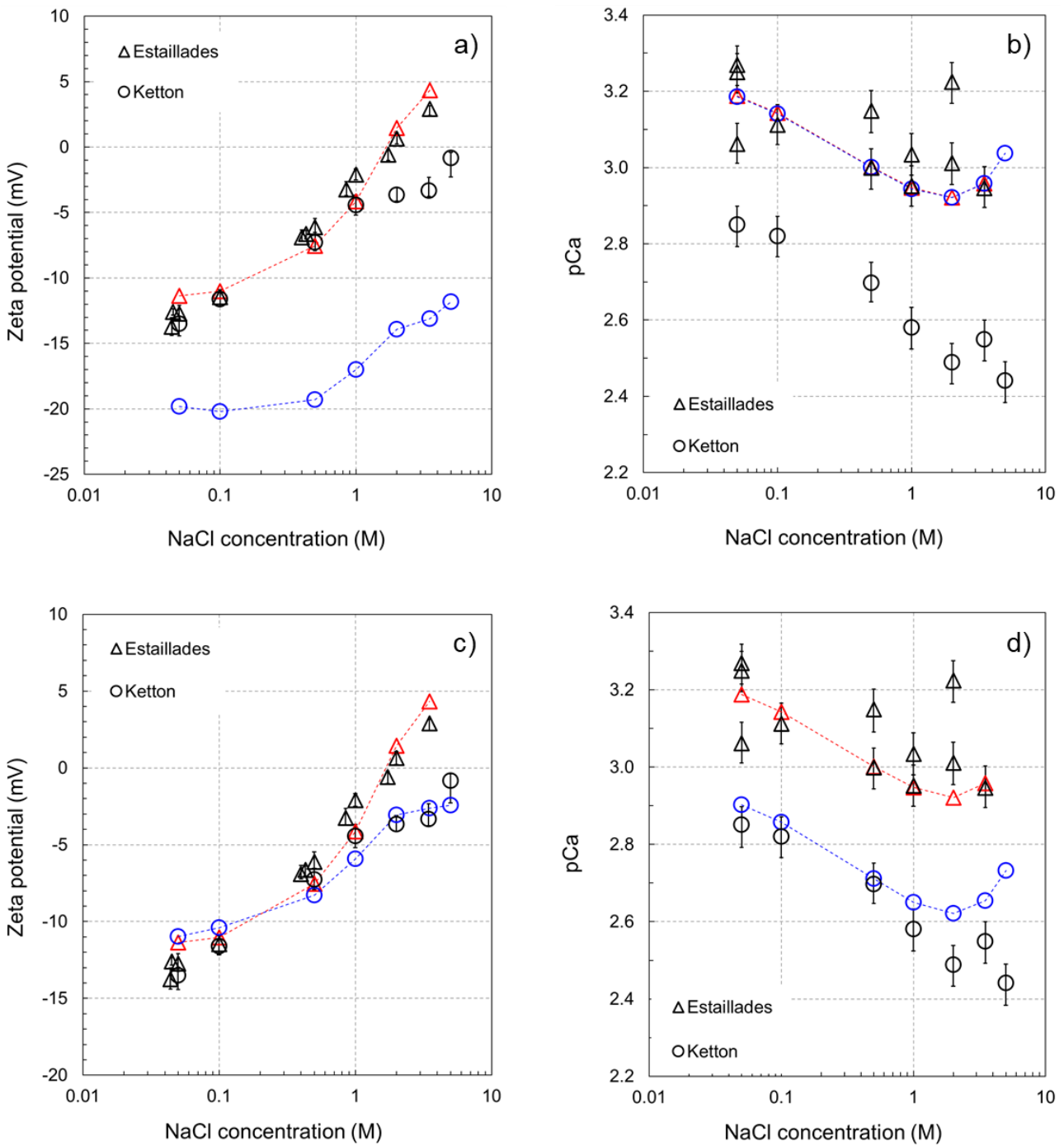
- 473 • Define the *saturation index* and the *amount* (typically, 10 mol to 20 mol to allow sufficient
474 amount of calcite to equilibrate with the solution of interest) in 'EQUILIBRIUM_PHASES'
475 section (see **Appendix A**). A non-zero *amount* should be chosen to replicate chemical
476 equilibration between calcite and water of the experiment of interest, while the *saturation index*
477 should be adjusted so that the computed Ca^{2+} concentration matches the experimental value.
- 478 • Define input concentrations of all ions in the modelled solution consistent with the reported
479 values. Note, that the input concentration of Ca^{2+} should be kept at zero and
480 dissolution/precipitation of calcite switched on, when simulating the zeta potential experiments
481 under equilibrium conditions, in which case content of Ca^{2+} in the solution is computed by
482 PHREEQC and cross-compared with the experimental data. Otherwise, the measured
483 concentration of Ca^{2+} should be used while switching off the calcite dissolution option (see
484 examples in the **Appendix A**).
- 485 • Use the optimized equilibrium constants for R1-R11 (**Table 2**).
- 486 • For a given ionic strength, calculate the Stern layer capacitance using the linear regression
487 equation (**Eq. 4**) and allow PHREEQC to compute the zeta potential. To validate the result
488 cross-compare the simulated pH and concentrations of all ions against the measured values.

489 3. Model validation, results and discussion

490 Our model described in previous sections was tested against the experimentally measured zeta
491 potential in intact carbonate core samples saturated with aqueous solutions reported by Al Mahrouqi
492 et al. [12], Jackson et al. [49] and Li et al. [11]. The developed model simulated the zeta potential
493 from SPM conducted on three different intact carbonate samples saturated with equilibrated NaCl
494 solutions [12], one intact carbonate sample saturated with equilibrated artificial solutions [49], and
495 crushed Iceland spar sample saturated with unequilibrated NaCl solutions [11].

496 3.1. Estailades and Ketton samples saturated with equilibrated NaCl solution [12]

497 We began modelling the zeta potential with defining the input equilibrium concentration of Na^+ , Cl^- ,
498 Mg^{2+} , SO_4^{2-} as reported in the paper. The initial input concentration of Ca^{2+} was set to zero M, partial
499 CO_2 pressure is set to $10^{-3.44}$ atm (consistent with atmospheric CO_2), and dissolution of calcite is
500 switched on (zero default value of the *saturation index* and 20 mol *amount*) to replicate the
501 equilibrium experimental conditions. Applying the specific capacitance values calculated using **Eq.**
502 **4** for each value of tested ionic strength our model computed the zeta potential and the resulting
503 equilibrium concentration of Ca^{2+} , and the results are shown in **Figures 8a** and **8b** (note, that
504 simulated pH and concentration of all ions except Ca^{2+} were found to be identical to the experimental
505 values used as input parameters). It can be seen from **Figures 8a** and **8b** that simulated results for
506 the Ketton sample underestimate the concentration of Ca^{2+} and consequently overestimate the
507 negative zeta potential. To be consistent with the experimental data, the calcite dissolution rate for
508 Ketton was enhanced by setting the *saturation index* to 0.8, so that the modelled equilibrium
509 concentration of Ca^{2+} became comparable with the measured one, which resulted in an excellent
510 correlation between the modelled and measured zeta potential (**Figures 8c** and **8d**).



511

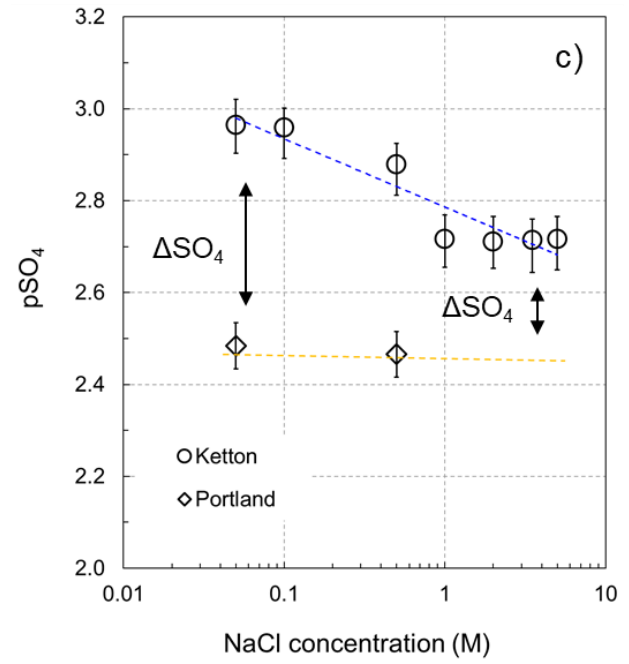
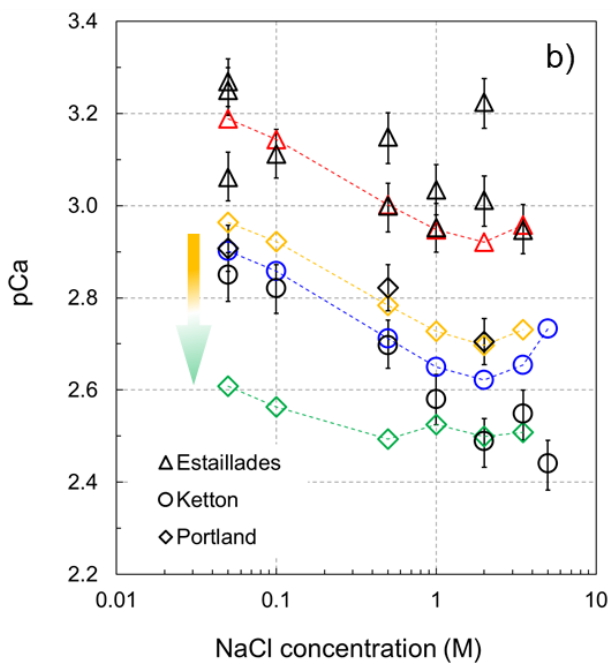
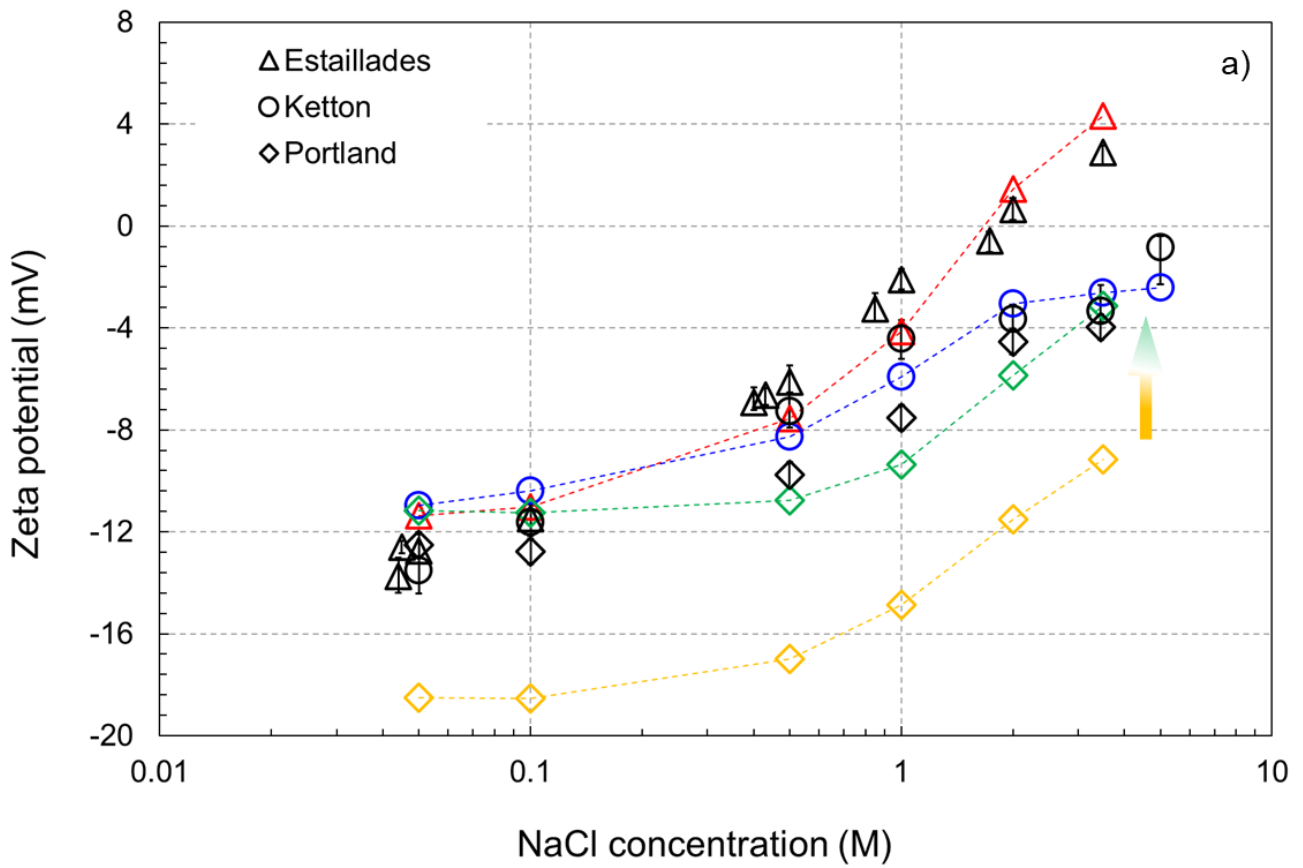
512 **Figure 8.** Experimentally measured (black symbols) and simulated zeta potential (a, c) and pCa (b, d) at
 513 equilibrium conditions for Estailades and Ketton samples saturated with NaCl solutions. The modelled results
 514 were obtained assuming identical *saturation index* for both rock samples toward NaCl solutions (a, b), and
 515 higher *saturation index* for Ketton (0.80) sample relative to the default value (0) of Estailades (c, d).

516

517 3.2. Portland sample saturated with equilibrated NaCl solution [12]

518 Our model was then applied to simulate the zeta potential obtained from Portland limestone. To
 519 evaluate the equilibrium concentration of Ca^{2+} dissolved from the Portland sample we implemented
 520 a similar procedure by comparing the computed and measured equilibrium concentration of Ca^{2+} ,
 521 which was reported to be slightly lower than that for the Ketton sample (compare diamonds and
 522 squares in Figure 10a of Al Mahrouqi et al. [12]). A good agreement between the computed and
 523 measured Ca^{2+} concentration corresponded to the *saturation index* of 0.6. However, a significantly

524 higher, compared with either Estailades or Ketton, concentration of SO_4^{2-} was reported for
525 equilibrated solutions in contact with Portland (compare diamonds with squares in Figure 10b of Al
526 Mahrouqi et al. [12]), while the reported concentration of Mg^{2+} in all rock samples was the same (pMg
527 = 4.19). Since SO_4^{2-} in the equilibrated solution could only originate from dissolution of sulfate
528 containing minerals (anhydrite or epsomite), the equilibrium concentration of SO_4^{2-} would not be
529 expected to exceed that of the total concentration of Ca^{2+} and Mg^{2+} , which was not the case reported
530 for Portland sample. We could not explain this discrepancy, and thus assumed that the reported
531 concentration of Ca^{2+} was inaccurate and contained an experimental uncertainty, which resulted in
532 underestimated concentration of the cation. Therefore, we artificially added Ca^{2+} to our simulated
533 aqueous solution, so that the concentration of the added cation was equal to the difference in sulfate
534 concentration across the salinity range: $\Delta[\text{SO}_4^{2-}] = [\text{SO}_4^{2-}]_{\text{Portland}} - [\text{SO}_4^{2-}]_{\text{Ketton}}$. Note that the
535 difference in sulfate concentration depends on the salinity as shown in **Figure 9c**, so that a single
536 value of the *saturation index* could not be used thus justifying our approach of adding Ca^{2+} to the
537 solution artificially. Keeping the *saturation index* equal to 0.6, the total computed concentration of
538 Ca^{2+} increased by the amount equal to $\Delta[\text{SO}_4^{2-}]$ (as denoted by the yellow→green shift in **Figure**
539 **9b**), and we successfully modelled the zeta potential as presented in Figure 9a (the yellow→green
540 shift in **Figure 9a** shows the effect on the computed zeta potential made by adding Ca^{2+} to the
541 solution).



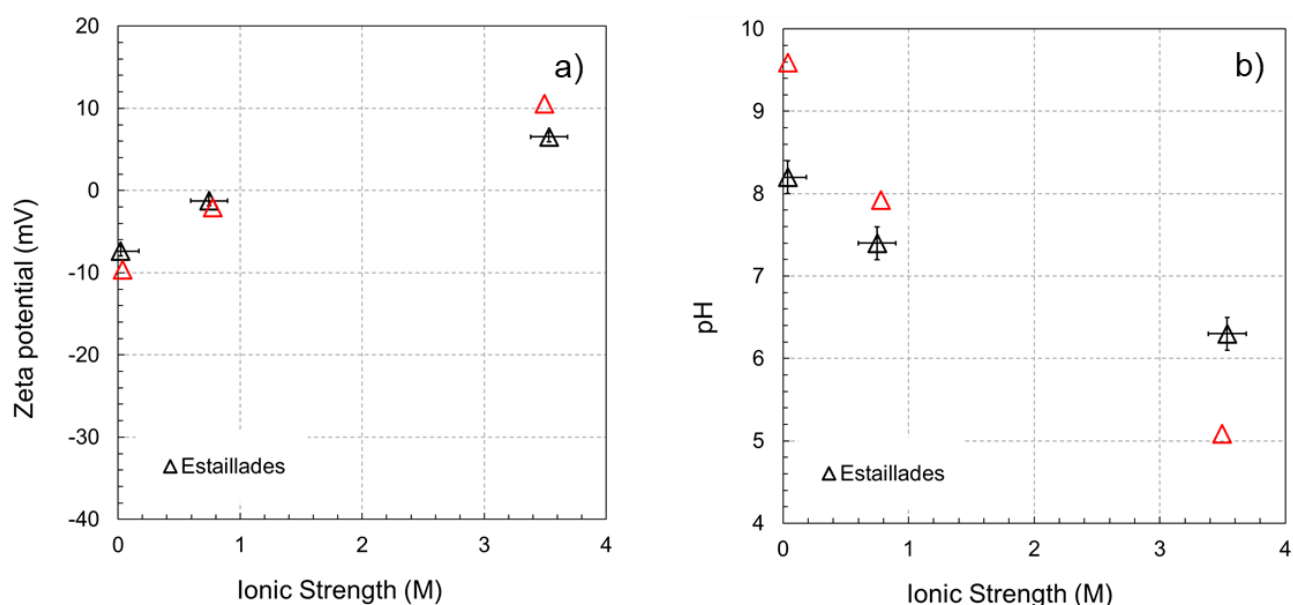
542

543 **Figure 9.** Measured and modelled zeta potential (a), pCa (b) and SO_4^{2-} (c) of three natural rock samples. The
 544 amount of artificially added Ca^{2+} to the model of Portland equals the reported difference in concentration of
 545 SO_4^{2-} as shown in (c) and explained in Section 3.2. Yellow diamonds represent the modelled zeta potential (a)
 546 and pCa (b) assuming *saturation index* of 0.6 and zero artificially added Ca^{2+} . The results of the model with
 547 artificially added Ca^{2+} are denoted by green diamonds.

548

549 **3.3. Estailades sample saturated with artificial natural solutions [49]**

550 Jackson et al. [49] measured zeta potential on Estailades rock sample saturated with three different
 551 equilibrated solutions: low salinity 20dSW (20 times diluted seawater), SW (seawater) representing
 552 moderate ionic strength of 0.749 M, and 3.537 M high salinity FMB (formation brine), with detailed
 553 composition provided in Table 1 in Jackson et al. [49]. The purpose of these experiments was to
 554 simulate the conventional and/or inverted low salinity waterflooding used to maximize oil recovery.
 555 To simulate the results, we modelled equilibrium conditions between calcite and the aqueous
 556 solutions by using zero initial concentration of Ca^{2+} and the same calcite dissolution rate as in Section
 557 3.1. for Estailades (default zero value of *saturation index*). The results of modelled zeta potential for
 558 all three solutions are presented in **Figure 10**. Our model could successfully reproduce the measured
 559 zeta potential (**Figure 10a**). However, the modelled equilibrium pH was significantly higher with
 560 20dSW and significantly lower with FMB experimental data (**Figure 10b**) despite the fact that
 561 modelled concentrations of all ionic species were found to be identical to the measured ones. We
 562 hypothesize that such discrepancy could result from the experimental protocol reported by Jackson
 563 et al. [49] who reported the initial (at time of preparation) pH and ionic concentration of solutions but
 564 not the final values established after equilibration with the rock sample.



565 **Figure 10.** Simulated zeta potential (a) and pH (b) of three different brine compositions for Estailades rock
 566 sample plotted as a function of ionic strength. The experimental data denoted by black symbols are extracted
 567 from Jackson et al. [49].
 568

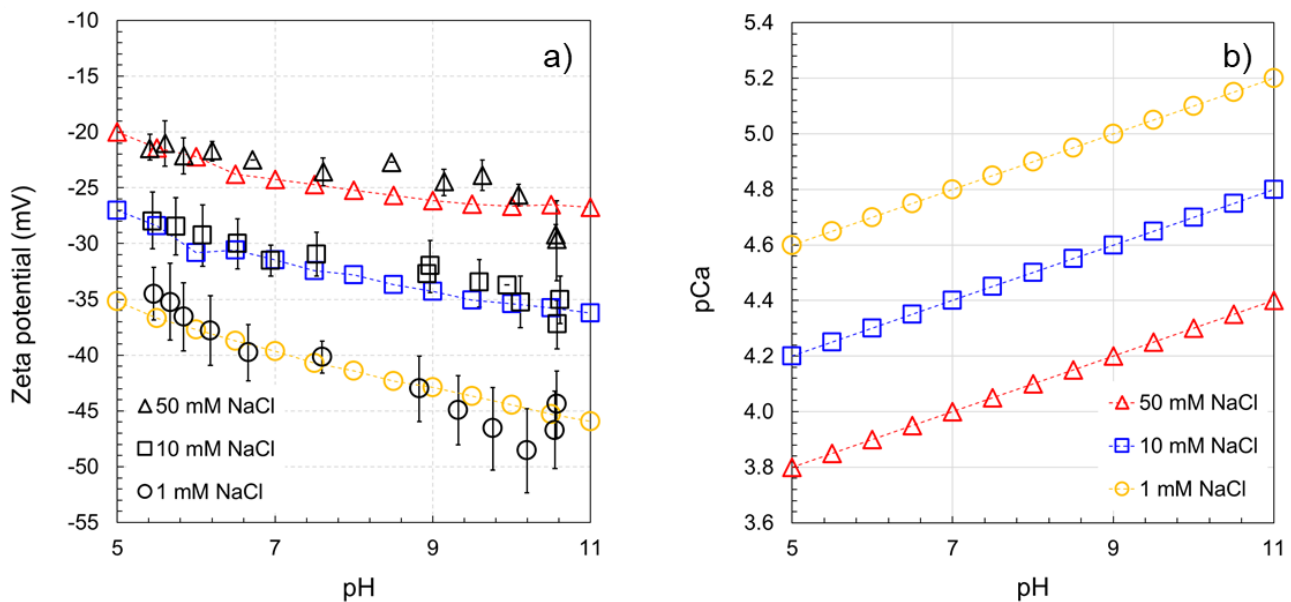
569
 570 It is worth mentioning, that equilibrium pH computed by our model in all tested aqueous solutions
 571 and rock samples matched the experimentally measured values apart from the results presented in
 572 **Figure 10**, where the simulated equilibrium pH was higher than experimental for 20dSW and SW and
 573 lower than experimental for FMB. Inclusion of protonation of $> \text{CO}_3^{-0.5}$ surface sites without changing
 574 the optimized equilibrium constants resulted in increased simulated equilibrium pH compared with
 575 the experimental values. Thus, the assumed exclusion of protonation of the $> \text{CO}_3^{-0.5}$ surface sites
 576 was confirmed for all tested parameters except for FMB, for which such reaction might need to be
 577 included, and this will be tested in a follow-up study.

578 3.4. Iceland spar saturated with NaCl at non-equilibrium conditions [11]

580 Finally, our model was tested against the data reported by Li et al. [11] on crushed Iceland spar
 581 saturated with 10^{-3} M, 10^{-2} M, and 5×10^{-2} M NaCl not equilibrated with the mineral. Description of the

582 experimental procedure reported by Li et al. [11] did not provide detailed information on the duration
583 of the streaming potential measurements. However, considering the expected high permeability of
584 the crushed samples (comparable to permeability of sandpacks reported by Vinogradov et al. [50])
585 and low salinity solutions used by Li et al. [11], it was assumed that the reported streaming potential
586 measurements on crushed Iceland spar did not last hundreds of hours required for complete
587 equilibration between the mineral and the tested solutions. Therefore, we assumed that only partial
588 equilibration was reached during the experiments and for that reason we modelled the data reported
589 by Li et al. [11] as obtained at non-equilibrium conditions so that any equilibration of calcite was
590 disabled in the model (zero *amount* was set, Section 2.3 and **Appendix A**). However, due to partial
591 equilibration expected in the experiments, we artificially added some non-zero initial concentration
592 of Ca^{2+} to the modelled solution. The concentration of dissolved Ca^{2+} in 0.05 M NaCl was adjusted
593 to $10^{-4.2}$ M (expressed as $\text{pCa} = 4.2$ in **Figure 11b**) at pH 9, consistent with observations reported by
594 Alroudhan et al. [19]. Firstly, we assumed that the amount of dissolved (and therefore artificially
595 added to the model) Ca^{2+} should depend on pH (lower pH would lead to higher concentration of
596 dissolved Ca^{2+} , consistent with higher dissolution rate reported by Anabaraonye et al. [51]).
597 Furthermore, the pH dependence of pCa in 0.05 M NaCl experiments was extended throughout the
598 entire range of pH (5-11) using a linear slope of $\Delta \text{pCa} / \Delta \text{pH} = 0.1$, consistent with calcite dissolution
599 rate reported by Chou et al. [52].

600 Secondly, we assumed that dissolved Ca^{2+} concentration should also depend on NaCl concentration
601 (higher NaCl concentration would lead to higher concentration of dissolved Ca^{2+} as reported in Al
602 Mahrouqi et al. [12]). Therefore, for NaCl solutions of 10^{-2} M and 10^{-3} M the end-points of pCa were
603 moved up the vertical scale by 0.4 units relative to one another to reflect on the salinity dependence
604 of calcite dissolution. The value of 0.4 was approximated from the reported calcite dissolution rate
605 as a function of NaCl concentration [53] and assuming duration of the steaming potential
606 measurement experiments conducted by Li et al. [11] to be approximately 10 hours. Ultimately,
607 ascribing the end-point values for Ca^{2+} concentration for each tested NaCl solution on the pH scale
608 (**Figure 11b**) and applying linear variation of pCa between these end-points, our model successfully
609 predicted the measured zeta potential (**Figure 11a**). Moreover, our model predicts that Ca^{2+} must
610 be added to the modelled solution in order to replicate the experimentally measured zeta potential.
611 Note, that $\text{C}(4)$ of the amount equal to the added Ca^{2+} must also be added to the solution in addition
612 to the initial concentration of $\text{C}(4)$ that reflects the equilibrium content of dissolved CO_2 . Despite the
613 fact that the computed from calcite dissolution or artificially added Ca^{2+} and $\text{C}(4)$ to the bulk
614 electrolyte was not reported by Li et al. [11], the adsorption reactions for these ions were included in
615 their SCM. Therefore, we are confident that our approach is consistent with Li et al. [11]. Our results
616 also explain how the experimental data from Li et al. [11] was successfully matched by means of
617 non-zero concentration of Ca^{2+} and $\text{C}(4)$ in the solution, both of which should increase with
618 decreasing pH. If such pH dependence of Ca^{2+} and $\text{C}(4)$ concentration was not applied to the model
619 of Li et al. [11], the simulated zeta potential would become more positive with increasing pH, which
620 is an inverted trend relative to that reported in the paper.



621

622 **Figure 11.** Modelled zeta potential (a) and pCa (b) of crashed Iceland spar samples saturated with three
 623 different NaCl solutions under non-equilibrium conditions. The modelled zeta potential is plotted in comparison
 624 with the experimental data [11] denoted by black symbols. The slopes and intercepts of the pCa as a function
 625 of pH are discussed and defined in Section 3.4.

626

627 3.5. Model capabilities, limitations and implication to carbonate-water subsurface settings

628 The developed robust SCM uses a set of justified and well-defined calcite surface sites and surface
 629 reactions' equilibrium constants combined with physically meaningful salinity dependence of the
 630 Stern layer capacitance. The model is capable to accurately predict zeta potential of a variety of
 631 carbonate-aqueous solution systems as long as calcite is the dominating mineral comprising natural
 632 rocks and the aqueous solution is dominated by NaCl. Moreover, an additional requirement for
 633 predictive capability of our model relates to the naturally occurring concentration of SO_4^{2-} in the
 634 solution, which has to reflect dissolution of sulfate containing minerals such as anhydrite or epsomite
 635 over geological deposition timescale. This requirement implies that the equilibrium concentration of
 636 SO_4^{2-} should not exceed the combined concentration of Ca^{2+} and Mg^{2+} .

637 The set of equilibrium constants proposed in this paper for ambient conditions (25°C and 1 atm) has
 638 not been tested to model experimental results obtained with aqueous solutions with artificially added
 639 SO_4^{2-} beyond natural occurrence, so that some amendments could be required to model such
 640 experiments. Moreover, we have not considered elevated temperature experimental conditions,
 641 which would require additional adjustments to the model: i) the equilibrium constants should be
 642 adjusted for elevated temperature by using for example the Van't Hoff equation; ii) the relative
 643 permittivity of the Stern layer should be recalculated; iii) the distance between the 0-plane and the
 644 β -plane should be reduced to account for reduced hydrated diameter of cations at elevated
 645 temperature (consistent with [50,54]); iv) the salinity dependence of the Stern layer capacitance
 646 should be amended taking into account ii) and iii) so that a new temperature specific regression is
 647 used.

648 Capabilities of our model have a broad range of applications since the approach reported here has
 649 demonstrated to provide an accurate prediction of the zeta potential of natural carbonate rock in
 650 contact with aqueous solutions, subject to the above conditions. Our model works best for equilibrium
 651 experimental conditions as long as concentration of all major ionic species is known. In this sense,
 652 our model can significantly improve our understanding of streaming potential measurements and

653 associated flows in shallow aquifers where temperature and pressure are low and therefore,
654 laboratory measurements of pH and concentration of all ions of the solution at aquifer conditions are
655 straight forward. Laboratory experiments should, in this case, assure establishment of full equilibrium
656 between rock and the solution of interest and fluid samples should be analyzed for molar
657 concentration of all ions to be used as input parameters of the model. The modelled zeta potential
658 can then be used to predict flow patterns in critical zones (e.g., [55]), permeability heterogeneities
659 (e.g., [56]) or even serve as an early warning of saline intrusion (e.g., [2,3]).

660 Our model is useful in situations where the exact concentration of all ions is unknown. For example,
661 the model can accurately assess the initial estimate of CO₂ geological storage efficiency. The model
662 would require input from the experimental data, given that laboratory experiments are conducted to
663 establish full chemical equilibrium between carbonate rock and a CO₂-saturated aqueous solution at
664 the target formation conditions of temperature and pressure. Such experiment should report at least
665 one measured concentration (of constituent ions such as Ca²⁺, Mg²⁺, SO₄²⁻ or protons, pH) to validate
666 the simulated by our model concentrations (similar to the experiments reported by Li et al. [57]). The
667 modelled zeta potential then can be used to interpret the wetting state using Derjaguin-Landau-
668 Verwey- Overbeek (DLVO) theory (e.g.,[58]) and the resulting residual trapping of the gas.

669 **4. Conclusions**

670 We report a robust surface complexation model of calcite-water interfaces. The model was
671 developed adopting a two-step optimization, in which the equilibrium constants of surface reactions
672 were initially optimized to match two experimental datasets, while the ionic strength dependence of
673 the Stern layer capacitance was obtained in the second optimization step. The model was applied
674 to successfully predict multiple experimental datasets with an excellent quality of match, and the
675 modelling results demonstrate that:

- 676 • The unique set of the optimized equilibrium constants can be used universally to simulate the
677 calcite-water zeta potentials obtained from the streaming potential measurements for all tested
678 systems and conditions.
- 679 • The Stern layer capacitance should decrease with increasing salinity to replicate high salinity zeta
680 potentials; the range of varying capacitances was found to be consistent with analytically
681 predicted values [48] and with values used in previously published numerical studies [45].
- 682 • Using experimental composition of investigated solutions as the only input, our model accurately
683 predicts zeta potentials of all tested systems and conditions including natural limestones
684 equilibrated with simple salt and complex solutions of ionic strengths between 0.05 M and 5 M,
685 and crushed calcite not equilibrated with NaCl solutions of ionic strength between 0.001 M and
686 0.01 M.
- 687 • To simulate zeta potentials of natural carbonate-water systems at equilibrium conditions, the
688 developed model requires knowledge of concentration of SO₄²⁻ and Mg²⁺ that leach from dolomite
689 and anhydrite inclusions, while the equilibrium concentration of Ca²⁺ is produced by the model.
- 690 • To simulate zeta potentials of calcite in contact with water under non-equilibrium conditions
691 requires knowledge of either measured in *real-time* Ca²⁺ concentration or duration of the
692 simulated experiment.
- 693 • Our model is fully predictive given the required input parameters (e.g., ion concentration) are
694 provided. However, additional surface reactions and/or model adjustments might be required to
695 simulate the zeta potentials for carbonate rock in contact with aqueous solutions in which SO₄²⁻

696 concentration exceeds that of Ca^{2+} and Mg^{2+} combined, this will be investigated in a follow-up
697 study.

698 Future work will also aim at acquiring additional experimental data obtained with varying
699 concentration of C(4) and SO_4^{2-} ions under equilibrium conditions to inform the surface complexation
700 model, which will be updated and/or modified to include the hypothesis of ion bridging outside OHP.

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- 883

884 **Appendix A – PHREEQC code examples**

885 In examples below, text that follows the # symbol is not part of the code, but comments aimed at
886 clarifying the meaning of program keywords and values

887

888 **EQUILIBRIUM_PHASES** #basic description

889 **Calcite 0 20**

890 # 0 is the default *saturation index*; 20 is the available *amount* of calcite in mol. The *saturation index*
891 can be either negative or positive to hinder or enhance calcite dissolution, respectively. To prevent
892 any dissolution or precipitation of calcite during the equilibration, the *amount* should be set to zero.

893

894 **CO2(g) -3.44**# defines partial CO₂ pressure at equilibrium with atmospheric air equal to 10^{-3.44} atm

895 -----

896

897 **EQUILIBRIUM_PHASES** #example 1 – simulation of established equilibrium between calcite and
898 water, *default* saturation index (Estailades). The computed equilibrium concentration of Ca²⁺
899 dissolved in 0.05 M NaCl is pCa=3.18

900 **Calcite 0 20**

901 **CO2(g) -3.44**

902

903 **SOLUTION 1**

904 **temp 25**

905 **units mol/L**

906 **Na 0.05**

907 **Cl 0.05 charge** #the **charge** keyword forces charge balance of the solution by adjusting
908 concentration of Cl

909 **Mg 0.0000645** #experimental

910 **S(6) 1.800E-4** #experimental

911 **SAVE SOLUTION 1**

912 -----

913

914 **EQUILIBRIUM_PHASES** #example 2 – simulation of established equilibrium between calcite and
915 water, *adjusted* saturation index (Ketton). The computed equilibrium concentration of Ca²⁺ dissolved
916 in 0.05 M NaCl is pCa=2.90

917 **Calcite 0.80 20**

918 **CO2(g) -3.44**

919

920 **SOLUTION 2**

921 **temp 25**

922 **units mol/L**

923 **Na 0.05**

924 **Cl 0.05 charge**

925 **Mg 0.0000645** #experimental

926 **S(6) 0.011** #experimental

927 **SAVE SOLUTION 2**

928 -----

929
930 **EQUILIBRIUM_PHASES** #example 3 – simulation of no established equilibrium between calcite and
931 water (Iceland Spar), default saturation index non-zero input concentration of Ca²⁺.
932 **Calcite 0 0**
933 **CO2(g) -3.44**
934
935 **SOLUTION 3**
936 **temp 25**
937 **units mol/L**
938 **Na 0.05**
939 **Cl 0.05 charge**
940 **Ca 0.0000631** #equivalent to pCa=4.2 interpreted from Figure 2a (Alroudhan, et al., 2016)
941 assuming 10 hours of partial equilibration during the experiment, which results in pH=9.
942 **SAVE SOLUTION 3**
943 -----