Laboratory Measurements of Zeta Potential in Fractured Lewisian Gneiss: Implications for the Characterization of Flow in Fractured Crystalline Bedrock

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Abstract: Despite the broad range of interest and possible applications, the controls on the electric surface charge and the zeta potential of gneiss at conditions relevant to naturally fractured systems remain unreported. There are no published zeta potential measurements conducted in such systems at equilibrium, hence, the effects of composition, concentration and pressure remain unknown. This study reports zeta potential values for the first time measured in a fractured Lewisian gneiss sample saturated with NaCl solutions of various concentrations, artificial seawater and artificial groundwater solutions under equilibrium conditions at confining pressures of 4 MPa and 7 MPa. The constituent minerals of the sample were identified using X-ray diffraction and linked to the concentration and composition dependence of the zeta potential. The results reported in this study demonstrate that the zeta potential remained negative for all tested solutions and concentrations. However, the values of the zeta potential of our Lewisian gneiss sample were found to be unique and dissimilar to pure minerals such as quartz, calcite, mica or feldspar. Moreover, the measured zeta potentials were smaller in magnitude in the experiments with artificial complex solutions compared with those measured with NaCl, thus suggesting that divalent ions (Ca²⁺, Mg²⁺ and SO₄²⁻) acted as potential determining ions. The zeta potential was also found to be independent of salinity in the NaCl experiments, which is unusual for most reported data. We also investigated the impact of fracture aperture on the electrokinetic response and found that surface electrical conductivity remained negligibly small across the range of the tested confining pressures. Our novel results are an essential first step for interpreting field self-potential (SP) signals and facilitate a way forward for characterization of water flow through fractured basement aquifers.

Keywords: zeta potential; fractured gneiss; surface electrical conductivity; effect of concentration, composition and mineralogy; impact of fracture aperture and confining pressure

1. Introduction

Fractured crystalline basement aquifers are predominantly made of granites and gneisses are widespread [1] and are locally important for rural water supplies and as a geothermal resource (e.g., [2,3]). They are predominantly made of granites and gneisses and mostly occur in Precambrian shield regions covering large parts of Europe (especially Scandinavia), North and South America, Africa, Australia, and in Precambrian orogens such as the Lewisian basement of NW Scotland [3].

Unlike in tropical regions, where deep weathering of crystalline basement rocks produce a porous aquifer (saprolite) with relatively high water storage capacity [4], in temperate/cold regions where weathering has been limited or absent, or scraped by glacier
erosion, groundwater flow is mostly restricted to connected fracture networks and, while fracture porosity is generally low, permeability can be relatively high (e.g., [5,6]). Understanding the fluid flow and storage properties in fractured rocks is challenging due to the discrete nature, heterogeneity and scale effect of fracture systems (e.g., [7–9]). It is of major importance to understand fractured systems in terms of hydraulically active fractures (versus hydraulically inactive) controlled by fracture connectivity, and fracture geometry including aperture and direction, which overall results in strong anisotropy in groundwater flow.

Direct characterization of groundwater flow patterns in fractured rock aquifers usually involves borehole studies including hydraulic testing or tracer testing methods, or a combination of both (e.g., [10,11]). Due to low permeability and storativity of fractured aquifers, both methods usually provide information on aquifer and fracture network properties only in the close vicinity of the borehole [12]. It is challenging to extrapolate the information to site- or region-scale due to large, multiscale heterogeneities of fracture systems (e.g., [7,13,14]). Surface geophysical methods are increasingly used to support interpolation to larger scales by providing indirect quantitative information on the heterogeneities of fracture networks and associated equivalent hydrodynamic properties (e.g., [12,15]). Most common geophysical methods used for this purpose are electrical resistivity tomography (ERT, e.g., [16]), ground-penetrating radar (GPR, e.g., [17]), magnetic resonance [18], or a combination of these techniques.

However, these methods are usually applied to provide equivalent isotropic flow/storage properties of the bulk aquifer over a certain volume, or structural information, rather than the flow properties of the fracture networks themselves. They are also expensive to deploy in the field in terms of effort and equipment. More recently, alternative, low-cost surface geophysical methods, such as self-potential (SP), have shown promise to better quantify fracture flow, including anisotropy, in crystalline rock aquifers (e.g., [19–21]). SP arises in response to pressure concentration or temperature gradients associated with the motion of water or aqueous solutions [22]; therefore, any signal acquired from electrode arrays reflects the dynamic behavior of the effective aquifer pore space, i.e., hydraulically active fractures in fractured rock aquifers. The method is passive and can be applied using surface arrays installed into the overlying soil layer (e.g., [23]) or discrete observation boreholes equipped with electrodes (e.g., [24]) and, therefore, provide information at both high spatial and temporal resolution.

A numerical study [25] investigated the potential application of SP to characterize fractured systems, and the authors demonstrated that the simulated SP signal not only accurately predicted preferential orientation of hydraulically active fractures but also identified fracture-matrix fluid exchange. The reported results indicated that the SP signal was primarily sensitive to hydraulically active fractures having matrix-fracture fluid exchange, thus implying that a non-zero porosity and near-zero permeability should be assigned to the matrix, which accommodated a significant volume of water but did not contribute significantly to the flow, and would be required to characterize the flow using SP. Assuming a strictly impervious matrix (zero porosity and zero permeability), resulted in a negligibly small electrokinetic signal. However, the authors assumed their fracture apertures to range between 0.2 mm and 2 mm, thus implying near-zero pressure gradients along the fractures and, therefore, a negligible SP response. In real fractured crystalline rocks, the aperture is expected to be a few orders of magnitude smaller (as discussed below); hence, the SP signal in such systems would be measurable even without matrix-fracture fluid exchange, thus, making the SP method applicable for such systems. In support of our hypothesis, a field SP survey on a crystalline bedrock [20] clearly demonstrated that azimuthal SP gradients tightly correlated with the investigated fracture strike orientation even though the matrix of the studied field site was impermeable and non-porous, consistent with the nature of crystalline bedrocks. Moreover, azimuthal resistivity surveys conducted on the same site showed the expected orientation of high water content, but were completely misaligned with the measured fracture network, thus making the SP method superior.
for identifying hydraulically active fractures, essential for accurate characterization of groundwater storage and flow.

Other published studies used the SP method to characterize and predict temporal variation of head gradients in a fractured aquifer as the results of diurnal tide cycles [26], and also to characterize saltwater intrusion into freshwater chalk aquifers (e.g., [24,27]). These studies demonstrated that the flow of freshwater and/or seawater through fractures was responsible for observed temporal and spatial variation of the SP measured in an observation borehole, even though the matrix was nearly impermeable. Moreover, the observed SP precursor that preceded seawater breakthrough was demonstrated to result from spatial variability of the so-called exclusion efficiency that was primarily associated with existing anisotropy in mineralogy and permeability of the aquifer. Therefore, not only was the SP method proven to be applicable to fractured systems, but it was also shown to rely on the system’s anisotropies to characterize dynamic behavior of flows.

Another modelling study on heterogeneous hydrocarbon reservoirs (Jackson et al., 2012a) during oil production showed that the SP measured in the production well would successfully predict the shape of the waterfront encroaching on the well, even when it was hundreds of meters away from the well. The study simulated the expected SP response from a mineralogically homogenous reservoir with permeability anisotropy, which did not intercept the well or could not be detected by seismic surveys. Hence, the study reported results from the so-called forward modelling so that a thorough inversion of the SP signal along with information on expected spatial variation of mineralogy would be required to quantify the exact distance of the permeability anisotropy from the well and the expected flow velocities across the reservoir from the measured SP signal. Nonetheless, even without such information, SP was shown to be useful in qualitatively predicting the waterfront shape (e.g., [22,28]) and quantitatively locating it relative to the instrumented well.

However, the inversion of the SP field measurements (as explained above) and gain of both qualitative and quantitative interpretation of the groundwater flow (i.e., preferential flow paths, anisotropy of flow and hydrodynamic properties, pressure gradients, local flow velocities) requires values of a key petrophysical property, termed the zeta potential, which characterizes electrochemical interactions that take place at the mineral-water interface. The zeta potential depends on the aquifer lithology, mineralogy, pore space topology (e.g., [29,30]) as well as on groundwater properties, specifically ionic composition, pH and concentration of aqueous solutions [31].

Numerous studies have reported zeta potentials measured in sandstones (e.g., [31–33]), carbonates (e.g., [34]) and granites [35] in contact with aqueous solutions of varying ionic strength and composition. However, no such measurements have been reported for gneiss-bose rocks, which form large parts of crystalline basement regions, and are the focus of this study. Therefore, the aim of this study is to thoroughly investigate the zeta potential of fractured gneiss samples saturated with simple and complex aqueous solutions under a range of confining pressures.

2. Materials and Methods

The rock sample used in this study was cored from a weathered gneiss boulder collected from the Precambrian Lewisian Gneiss Complex in Northwest Scotland, near Clashnessie. The cylindrical rock sample (38 mm diameter, 54 mm length) was subjected to uniaxial loading under controlled stress and artificially fractured along pre-existing cracks. To preserve the cylindrical shape of the sample, it was wrapped in duct tape during the loading. The initial and final state of the sample is shown in Figure 1.
Figure 1. Rock sample used in this study was cut from the weathered outcrop boulder (a) collected from the Clashnessie area denoted by the red star on the inset map (refer to the geological map of the area, S1, in the supplementary material); the core sample was naturally fractured as demonstrated by the top (b), bottom (d) and side (f) views; the sample was loaded along the longitudinal axis resulting in at least one fracture that extended from the top face (c) to the bottom one (e); to preserve its cylindrical shape during uniaxial loading, the sample was wrapped in duct tape (g) and experienced fracturing that corresponded to bulk displacements of 1 mm, 1.4 mm and 1.7 mm, as demonstrated by the load [kN] vs. displacement [mm] graph (h).

To mimic the fluids expected to saturate the fractured aquifer in similar conditions as encountered at the site where the sample was taken, i.e., in a coastal setting of NW Scotland, we used four aqueous solutions to measure the streaming potential coupling coefficient: (1) the artificial groundwater (AGW) comprising the main salts, consistent with the groundwater chemical composition as reported by BGS [36] for groundwater samples taken from similar Lewisian gneiss on the Isle of Harris (about 50 km West of the site, blue circle in Figure 1a; (2) the artificial seawater (ASW) formulated from the main ionic species; (3) low-salinity NaCl (LS); (4) high-salinity NaCl (HS) solutions. The ionic strength of LS and AGW was $7.35 \times 10^{-3}$ M (M = moL·L$^{-1}$), while the ionic strength of HS and ASW was 0.7 M, hence the ionic strengths of NaCl and the corresponding complex solution were identical between all low- and all high-salinity experiments. Therefore, the impact of ionic strength on the zeta potential measured with NaCl in comparison with complex solutions was eliminated to isolate the effect of the chemical composition. The ionic composition and equilibrium pH of all solutions used in this study are presented in Table 1.
Table 1. Chemical composition, equilibrium pH and electrical conductivity of tested solutions. The solution pH was measured using Mettler Toledo FiveGo pH meter, the electric conductivity of all solutions was measured using Jenway 4520 conductivity meter with the corresponding errors reflecting the instrument inaccuracy and measurement repeatability. AGW and LS solutions were prepared from high-salinity stock solutions of 0.735 M ionic strength, hence, errors in molar concentrations of all four tested solutions resulting from inaccuracies in weighing of the constituent salts are estimated at ±3% of the reported values.

|        | AGW          | LS            | ASW          | HS           |
|--------|--------------|---------------|--------------|--------------|
| [Na⁺] M | 1.522 × 10⁻³ | 7.350 × 10⁻³  | 0.512        | 0.700        |
| [Ca²⁺] M | 1.612 × 10⁻³ | -             | 0.013        | -            |
| [Mg²⁺] M | 0.288 × 10⁻³ | -             | 0.042        | -            |
| [Cl⁻] M | 5.072 × 10⁻³ | 7.350 × 10⁻³  | 0.577        | 0.700        |
| [SO₄²⁻] M | 0.125 × 10⁻³ | -             | 0.022        | -            |
| pH     | 6.0 ± 0.1    | 6.5 ± 0.1     | 6.2 ± 0.1    | 6.9 ± 0.1    |
| σₗw, S·m⁻¹ | 0.069 ± 0.001 | 0.097 ± 0.001 | 5.83 ± 0.01  | 6.13 ± 0.01  |
| Ionic strength, M | 7.350 × 10⁻³ | 7.350 × 10⁻³ | 0.700        | 0.700        |

To characterize the chemistry and mineralogy of the gneiss sample used in our experiments, XRD analysis was carried out on powdered offcuts. Moreover, a chip from the sample was polished to produce a thin section, which was analyzed using scanning electron microscopy (SEM). The main constituent minerals from the SEM analysis were identified to be: quartz (c. 35 wt.%), Na-feldspars (plagioclase including albite, c. 60 wt.% in total) and micas (muscovite and biotite, c. 5 wt.% in total). These mineral proportions are characteristic of felsic Lewisian gneiss, as previously reported (e.g., [37,38]).

Prior to the streaming potential measurements, the rock sample was cleaned using deionized water and methanol following the procedure described by Alroudhan et al. [39]. Note that the cleaning procedure did not have any noticeable effect on the sample, as measurements of the zeta potential and permeability were repeated several times after cleaning and returned the same values within the experimental uncertainty. Subsequently, the sample was saturated with the solution of interest and placed in a vacuum oven for 24 h to establish 100% of water saturation. The saturated rock sample was mounted in a core holder made of PEEK, similar to that described in detail by Vinogradov et al. [31]. The experimental setup used in this study, including a pair of Ag/AgCl external electrodes used to measure the streaming potential, was the same as the one reported in previous studies by Vinogradov and co-workers (e.g., [33]).

We used the paired-stabilized (PS) method [40] to measure the streaming potential coupling coefficient, from which the zeta potential was interpreted using the Helmholtz-Smoluchowski equation (e.g., [41]) and assuming negligible contribution of the surface electrical conductivity to the effective electrical conductivity of the aqueous solution occupying a mm-scale fracture (as discussed in subsequent sections):

\[
C_{SP} = \left. \frac{\Delta V}{\Delta P} \right|_{j=0} = \frac{\varepsilon \zeta}{\mu \sigma_w}
\]

where \(C_{SP} [V/Pa]\) is the streaming potential coupling coefficient calculated from the measured voltage \(\Delta V [V]\) and the pressure difference \(\Delta P [Pa]\) across the sample when the total current density \(j [A/m^2]\) is zero; \(\varepsilon [F/m]\) is the permittivity of water; \(\zeta [V]\) is the zeta potential; \(\mu [Pa.s]\) is the dynamic viscosity of water; and \(\sigma_w [S/m]\) is the electrical conductivity of the aqueous solution. Each measurement of the streaming potential coupling coefficient was carefully analyzed for potential error sources including voltage noise level, stability of voltage during the experiment and repeatability. The resulting cumulative
errors are presented by the individual error bars in Figures 2–4, as discussed below (all data accompanied by examples of SP experiment and error analysis are provided in S2 of the Supplementary Material).

Figure 2. Zeta potential of gneiss samples compared with published data on other types of rocks. Circles denote NaCl solutions, triangles correspond to AGW and squares are ASW. Filled symbols correspond to confining pressure of 7 MPa. Low ionic strength is denoted by blue symbols while the high ionic strength data corresponds to red symbols. The solid lines that outline domains of reported data for carbonates and sandstones are based on the published data summarized in Table 2.

Table 2. Summary of experimentally measured zeta potential in relevant samples saturated with NaCl and natural solutions. In the table, LS and HS correspond to low-salinity and high-salinity data, SS stands for sandstones, SP corresponds to sand packs, LS refers to limestones, SW is natural seawater from southern UK, TW is tap water and ALSW is artificial low-salinity water. Exact values of ionic strength in the table vary between 0.001 M and 0.01 M for low salinity, and between 0.5 M and 0.75 M for high-salinity data. For details, refer to the cited papers. The data sources are: a—Vinogradov et al. [31]; b—Walker and Glover [32]; c—Jaafar et al. [42]; d—Vinogradov et al. [43]; e—Vinogradov and Jackson [44]; f—Li et al. [45]; g—Tosha et al. [35]; h—Reppert and Morgan [46]; i—Morgan et al. [47]; j—Ishido and Mizutani [48]; k—Al Mahrouqi et al. [34]; l—Jackson et al. [49]; m—Collini et al. [50]; n—Heberling et al. [51]; o—Heberling et al. [52].

| Sample, Mineralogy | Solution   | Zeta Potential, mV | Source |
|--------------------|------------|--------------------|--------|
|                    | LS         | HS                 |        |
| Grey shaded area   | NaCl       | −52                | −20    | a, b   |
| Fontainebleau SS; >99% quartz | NaCl     | −77                | −25    | b      |
| Lochaline SS; >99% quartz     | NaCl     | −26                | −16    | a      |
| Stainton SS; 90% quartz, 5% clays and feldspar | NaCl | −52                | −20    | a, b   |
Table 2. Cont.

| Sample; Mineralogy                              | Solution | Zeta Potential, mV | Source |
|------------------------------------------------|----------|--------------------|--------|
| St Bess SS; 90% quartz, <5% clays              | NaCl     | −31                | −16    | a, c   |
| St Bees SS; 90% quartz, <5% clays              | SW       | N/A                | −13    | a      |
| Doddington SS; 69% quartz, 5% clays            | NaCl     | −22                | −10    | e      |
| SP; >99% quartz                                | TW       | −100               | N/A    | a      |
| SP; >99% quartz                                | NaCl     | −20                | −12    | d      |
| Berea SS; 90% quartz, up to 4% clays           | NaCl     | −20                | −17    | f      |
| Boise SS; 47% quartz, 26% plagioclase, 3% clays | NaCl     | −20                | −20    | b      |
| Inada granite; N/A                             | KCl      | −35                | N/A    | g      |
| Westerly granite, 38% plagioclase, 29% quartz  | NaCl     | −20                | N/A    | h      |
| Crushed Westerly granite; N/A                 | KCl/NaCl | −65                | N/A    | i      |
| Inada granite; N/A                             | KNO₃     | −70                | N/A    | j      |

White shaded area

| Ketton LS; 97% calcite, 3% dolomite             | NaCl     | N/A                | −6     | k      |
| Estailades LS; 95% calcite, 4% dolomite, 4% anhydrite | NaCl     | N/A                | −6     | k      |
| Portland LS; 96.6% calcite, 3.4% quartz        | NaCl     | N/A                | −9     | k      |
| Estailades LS; 95% calcite, 4% dolomite, 4% anhydrite | SW       | N/A                | −1.25  | l      |
| Reservoir BA LS; N/A                           | ALSW     | −4                 | N/A    | m      |
| Reservoir BD LS; N/A                           | ALSW     | −10                | N/A    | m      |
| Outcrop TE LS; >99% calcite                    | Artificial | −15              | −2     | m      |
| Crushed Iceland spar; 100% calcite             | NaCl     | −5 [n]             | +4 [o] | n, o   |

To replicate different depth of fractured gneiss of the aquifer of interest and, hence, different fracture aperture, two different confining pressures were tested with all four solutions: 4 MPa and 7 MPa.
3. Results and Discussion

3.1. Effect of Rock Type and Mineralogy

We begin with comparing the zeta potentials measured in the fractured gneiss sample against published data on pure sandstone (>99% quartz), clayey sandstone (2–5% clay content), sand packs (>99% quartz), carbonate (>95% calcite) and granite samples saturated with aqueous solutions of similar ionic strength and composition, and under equilibrium conditions (i.e., rock samples were thoroughly equilibrated with aqueous solutions). The results presented in Figure 2 indicate that the zeta potentials measured with gneiss saturated with high-salinity electrolytes were generally more negative compared with values obtained...
in carbonates saturated with aqueous solutions of ionic strengths within the range of those tested in this study, and were more positive across the salinity range compared with the data from sand packs, as well as from clean and clayey sandstones. All of our results in Figure 2 are color-coded with red symbols denoting the ionic strength of 0.7 M and the blue symbols corresponding the ionic strength of $7.35 \times 10^{-3}$ M.

The individual data points included in the grey and white shaded areas are summarized in Table 2, for which zeta potentials measured in intact and crushed sandstone, carbonate and granite samples have been reported. To the best of our knowledge, there have been no published data on zeta potentials measured with basement rocks (e.g., granite or gneiss) saturated with high ionic strength electrolytes, hence, the corresponding column is populated with N/A (not available).

The majority of the data detailed in Table 2 indicate that the zeta potentials of gneiss saturated with high-salinity solutions (0.7 M) are negative and larger in magnitude compared with carbonates, but smaller in magnitude compared with sandstones. However, there are two exceptions, with the zeta potential of $-10$ mV measured in Doddington sandstone saturated with 0.5 M NaCl solution, and the zeta potential of $-9$ mV measured in Portland limestone saturated with equilibrated 0.5 M NaCl solution, both of which are comparable with the values obtained for the gneiss sample saturated with ASW. We attribute these anomalous reported zeta potentials to a higher clay content in the Doddington sample and quartz inclusion in the Portland limestone, which made these samples’ response to 0.5 M NaCl solutions to be dissimilar to that of pure silica or calcite rocks, but comparable with the signal measured with gneiss.

On the other hand, the zeta potentials in the gneiss sample saturated with low-salinity electrolytes ($7.35 \times 10^{-3}$ M) were found to be more positive compared with both sandstone and carbonate samples, with the exception of $-5$ mV recorded with crushed Iceland spar saturated with 0.01 M NaCl mixed with 1 mM CaCl$_2$ solution [51], and $-4$ mV measured in reservoir carbonate sample BA of unknown mineralogy saturated with artificial 0.01 M electrolyte containing Ca$^{2+}$ and Mg$^{2+}$ [50]. The results with Iceland Spar were obtained at an unconfirmed chemical equilibrium with solutions and showed strong dependence on the concentration of added CaCl$_2$, so that the measured zeta potential varied between $-3$ mV with 5 mM of CaCl$_2$ and $-14$ mV with 0.1 mM of CaCl$_2$, thus, limiting the applicability of direct comparison between our results and those reported by Heberling et al. [51]. In contrast, the results reported by Collini et al. [50] were obtained under equilibrium conditions, but the solution used by the authors contained a considerable amount of Mg$^{2+}$ (1 mM), Ca$^{2+}$ (0.3 mM) and no SO$_4^{2-}$ in their 0.01 M solution. It is generally agreed that divalent ions act as potential determining ions (PDIs) for clays (e.g., [53]), so that divalent cations make the zeta potential more positive while divalent anions tend to make the zeta potential more negative. Hence, in absence of SO$_4^{2-}$ in the tested low-salinity solution, the presence of Ca$^{2+}$ and Mg$^{2+}$ made the zeta potential more positive compared with pure NaCl and seawater in sandstones, other carbonates or gneiss.

There have been numerous reported measurements of the zeta potential in carbonate samples (e.g., refer to Figure 3 in Jackson et al. [23]) and the values in these studies ranged between $-20$ mV and $+20$ mV for low-salinity (ionic strength below 0.01 M) solutions with pH of 7. However, as discussed by Al Mahrouqi et al. [34], none of these measurements were conducted under conditions of thermo-chemical equilibrium between the tested solutions and rock minerals, hence, these data cannot be directly compared with our results.

Moreover, considering the mineralogy of our gneiss sample, we have found a limited number of published studies reporting measurements of the zeta potential of plagioclase [54] including albite [55], muscovite [56] and biotite [57]. None of these studies conducted their electrophoretic mobility measurements under equilibrium conditions, hence, the tested minerals could dissolve during the experiments with single salt aqueous solutions. This situation would be similar to that reported by Alroudhan et al. [39] and Al Mahrouqi et al. [34] for carbonate samples, or the one reported by Reppert and Morgan [46] for granite, all of which observed varying zeta potential during their experiments
until it stabilized at the final equilibrium value after hundreds of hours of equilibration. Therefore, the measured zeta potentials of plagioclase, albite, biotite and muscovite systems could not be compared with our data and between each other in any other way, but qualitatively. Furthermore, it is worth noting that the scarce data available for these minerals cannot be utilized for interpreting the SP signal measured in fractured gneiss aquifers, which are expected to be fully equilibrated with aqueous solutions, thus emphasizing the importance of our laboratory results. The reported laboratory data on mica and feldspar minerals generally suggest that the zeta potentials remain negative across the pH range of interest (6–8) but become progressively smaller in magnitude following the order: $|\zeta_{\text{plagioclase}}| < |\zeta_{\text{albite}}| \cong |\zeta_{\text{biotite}}| < |\zeta_{\text{muscovite}}|$ with the values reported for plagioclase ranging between $-10$ mV and $+15$ mV obtained with $10^{-2}$ M di-sodium tetraborate solution [54].

All in all, our experimental results demonstrated that all zeta potentials of Lewisian gneiss, saturated with low- to high-salinity aqueous solutions consisting of a single salt or a mixture of different salts, were unique. The main constituent minerals of this rock type are quartz, feldspar and mica, so the electrochemical response to ionic species in tested solutions is expected to reflect the individual contribution of each mineral. Since there are no reported zeta potential measurements of feldspars and micas in contact with various aqueous solutions at equilibrium conditions, we hypothesize that the electric surface charge on these complex minerals is controlled by multi-valent PDIs, similar to the mechanism of the surface charge development on calcite surface (e.g., [58]). Therefore, the resulting surface charge and the corresponding zeta potential of gneiss-hosted feldspars in our experiments caused the zeta potential to become less negative compared with pure quartz systems. Micas in contact with aqueous solutions ([55,56]) were reported to produce more negative zeta potential compared with feldspars ([54,57]). At the same time, the negative zeta potentials measured with micas were smaller in magnitude compared with negative values on silica surface, consistent with a published study of silica and mica minerals in contact with $10^{-3}$ M KCl solution [56]. Moreover, previously published studies also reported a shift in the zeta potentials towards more positive values with solutions containing divalent cations relative to NaCl/KCl measurements for all types of minerals, with a more pronounced shift reported with micas [59] and feldspars (comparing the results obtained with albite in contact with $10^{-3}$ M NaCl in Gülgünlü et al. [60] vs. $10^{-3}$ M CaCl$_2$ in Demir et al. [61]) compared with pure quartz (e.g., [62]). Hence, the zeta potential in our experiments responded to changes in chemical composition in a fashion similar to that of calcite/mica/feldspar and became more positive with divalent cations, while the electric charge on the gneiss-hosted quartz surface sites was mostly controlled by pH, thus, remaining insensitive to changes in the electrolyte composition. Moreover, the uniqueness of the gneiss zeta potentials was demonstrated by the impact of varying ionic strength, which appeared to be less pronounced than that obtained with either carbonates or pure sandstones (>95% quartz). Both the impact of composition and ionic strength on the zeta potential in gneiss samples will be discussed in more detail in the subsequent section.

3.2. Effect of the Ionic Strength and Chemical Composition

It is generally agreed that zeta potential decreases in magnitude with increasing ionic strength as a result of a collapse of the electrical double layer (e.g., [63]) and our results with ASW and AGW, as shown in Figure 3a, were consistent with this behavior. The measured zeta potentials of these systems remained negative but became smaller in magnitude with increasing ionic strength. In contrast, the zeta potentials measured with NaCl solutions remained constant within the experimental uncertainty with increasing salinity, as demonstrated by the horizontal dash-dotted line in Figure 3b. Our results are denoted by the red and blue symbols in Figure 3.

As shown in Figure 3a, the rate at which the zeta potential decreases in magnitude with increasing salinity in gneiss samples (blue and red symbols) is significantly lower than that reported for clean sandstones (green dashed line) and carbonates (black dashed...
saturated with NaCl solutions. However, a recent study [45] reported an anomalous salinity dependence of the zeta potential in clayey Berea samples. In this study the zeta potential obtained with NaCl solutions varied from $-18.3$ mV at 0.005 M to $-21.4$ mV at 0.5 M (green circles in Figure 3), thus, indicating much weaker and reversed concentration dependence compared with that of clean sandstones. At the same time, the magnitude of zeta potential in Berea NaCl experiments remained significantly larger compared with our gneiss results. In contrast, the zeta potentials of approximately $-7 \pm 0.3$ mV measured in Berea-CaCl$_2$ systems (black circles in Figure 3) remained unchanged between ionic strength of 0.008 M and 0.8 M. In this sense, the independant of salinity zeta potential in the clayey sandstones saturated with CaCl$_2$ was qualitatively similar to the observed trend in the gneiss sample saturated with NaCl, but the magnitude of the latter was consistently larger.

The zeta potentials in our gneiss sample saturated with aqueous solutions of complex composition became more negative with decreasing salinity and was consistently more positive compared with the results obtained with NaCl. Moreover, the zeta potential of $-7.2 \pm 0.2$ mV obtained with ASW (0.7 M) were comparable with those measured in Berea samples saturated with CaCl$_2$ ($-7.0 \pm 0.3$ mV) [45]) and Portland limestone saturated with equilibrated NaCl ($-8.0 \pm 0.5$ mV) [34]).

Figure 3 clearly indicates that zeta potentials of Lewisian gneiss saturated with aqueous solutions are unique in their response to varying ionic strength and brine composition. These observations suggest that gneiss mineralogy affects the zeta potential that strongly responds to presence of divalent ions, hence Ca$^{2+}$, Mg$^{2+}$ and SO$_4^{2-}$ are PDIs for this rock type, similar to the generally agreed feature of carbonates (Al-Mahrouqi et al. [34]), feldspars (Demir et al. [61]) and micas (Adamczyk et al. [59]). Therefore, accurate laboratory measurements of the zeta potential in gneiss should be carried out prior to attempting interpretation of field SP data.

### 3.3. Effect of Confining Pressure

In all of our experiments, we observed the zeta potentials to be independent of the confining pressure (Figure 4). The increasing confining pressure applied to the fractured rock sample resulted in a decrease of the apparent permeability from 51 mD (1 mD $\approx 10^{-15}$ m$^2$) at 4 MPa to 43 mD at 7 MPa. The observed decrease in the permeability corresponds to a closure of the fracture, thus potentially leading to an increased effect of the surface electrical conductivity.

To quantify the effect of the confining pressure on the measured streaming potential coupling coefficient, zeta potential and the fracture aperture, we considered the following geometry of the investigated rock sample (Figure 5).

All our streaming potential experiments were conducted under steady-state fluid flow conditions and the induced flow through the fracture was consistent with the direction of the rock sample. Therefore, for the geometry described in Figure 5 and the experimental conditions, the Navier–Stokes equations simplify to 1-D flow in the Cartesian coordinate system as follows:

$$\frac{\mu}{2\pi} \frac{d^2v_x}{dy^2} = \frac{\Delta P}{L} \quad (2)$$

where $v_x(y)$ is the fluid velocity in $x$-direction inside the fracture. Applying the no-slip boundary conditions at both fracture walls, the fluid velocity profile is given by:

$$V_x = \frac{1}{2\mu} \frac{\Delta P}{L} \left( y^2 - w^2 \right) \quad (3)$$

Multiplying Equation (3) by the cross-section area to the flow, $2wH$, and integrating across the fracture aperture (between $-w$ and $+w$) yields the well-known cubic law [64] for the total volumetric flow rate, $Q$:

$$Q = -\frac{2Hw^3}{3\mu} \frac{\Delta P}{L} \quad (4)$$
Figure 5. The tested fractured rock sample is assumed to have a single fracture of aperture $2w$, height $H$ and length $L$.

Now, equating the flow rate given by Equation (4) with that from the Darcy’s law, $Q = \frac{-2kHw}{\mu} \rho \tau$, the effective permeability is expressed as:

$$k = \frac{w^2}{3}$$

Equation (5) is consistent with a recently published study [65] in the limit of a single fracture, and it implies that for a decrease of approximately 16% in sample’s permeability (from 51 mD to 43 mD), the corresponding expected decrease in fracture aperture ($2w$) will be around 8% (from 0.782 mm at 4 MPa to 0.718 mm at 7 MPa). Decreasing fracture aperture implies a smaller distance of separation between the fracture walls, and as such, a higher confining pressure can potentially have an impact on the electrical properties of hydraulically active fractures via an increased impact of the surface electrical conductivity and the zeta potential.

To investigate the impact of fracture aperture on the electrokinetic properties of the sample, we consider the streaming current under the steady state experimental conditions and for geometry described in Figure 5:

$$I_s = 2 \int_0^w \Sigma(y)v_s(y)Hdy$$

where $\Sigma$ [C/m$^3$] is the volumetric charge density. The flow of mobile excess counter-ions that contribute to the streaming current is confined to the diffuse part of the electrical double layer (EDL), which has a thickness of approximately 3 nm for $7.35 \times 10^{-3}$ M and an even smaller thickness for higher salinity, so that the thickest expected EDL will still be two orders of magnitude smaller than the estimated fracture aperture. Therefore, we can use
this condition to linearize the flow velocity that contributes to the streaming current in the region adjacent to the fracture walls, where \( y \approx w \):

\[
V_x = \frac{1}{2\mu} \frac{\Delta P}{L} (y^2 - w^2) \approx \frac{1}{2\mu} \frac{\Delta P}{L} 2w(y - w)
\]  

(7)

Introducing a new variable \( \theta = y - w \), using Poisson equation for the charge density, \( \Sigma = -\varepsilon \frac{dV}{dy} \) (where \( V \) is the electric potential and \( \varepsilon \) is the permittivity of the aqueous solution), substituting Equation (7) into Equation (6), and integrating by parts yields an expression for the streaming current through the fracture:

\[
I_s = -\frac{2wH \varepsilon \zeta}{\mu} \frac{\Delta P}{L}
\]

(8)

where \( \zeta \) is the zeta potential at \( y = \pm w \). We now apply the boundary conditions of zero net current coming in or out of the fracture, and no charge separation along the fracture. In such case, the streaming current must be balanced by the conduction current of equal magnitude and opposite direction. The conduction current is given by Ohm’s law:

\[
I_c = 2wH \sigma_w \frac{\Delta V}{L} + 2H \sigma_s \frac{\Delta V}{L} = 2wH \frac{\Delta V}{L} [\sigma_b + \sigma_s \frac{w}{w}]
\]

(9)

where \( V \) is the electric potential difference (voltage) across the fracture, \( \sigma_w \) [S/m] is the electrical conductivity of the bulk aqueous solution and \( \sigma_s \) [S] is the surface electrical conductivity due to the presence of EDL. Using the condition of electro-neutrality, \( I_s = -I_c \), the streaming potential coupling coefficient can be expressed:

\[
C_{SP} = \frac{\Delta V}{\Delta P} = \frac{\varepsilon \zeta}{\mu [\sigma_w + \sigma_s \frac{w}{w}]}
\]

(10)

Note that Equation (10) explicitly expresses the effective electrical conductivity of brine with the contribution of the surface conductivity, but becomes equal to the Helmholtz-Smoluchowski equation (Equation (1)) when the surface conductivity is negligibly small. The zeta potential can now be expressed from the measured streaming potential coupling coefficient as:

\[
\zeta = C_{SP} \mu [\sigma_w + \sigma_s \frac{w}{w}] / \varepsilon
\]

(11)

For the two tested ionic strengths of \( 7.35 \times 10^{-3} \) M and 0.7 M, the permittivity and viscosity of the NaCl solutions can be calculated using the approach described in Saunders et al. [28], while \( C_{SP} \) and the electric conductivity of the solutions were directly measured during the experiments. Rock, fluid and electrokinetic properties for all our NaCl experiments are summarized in Table 3 along with estimates of fracture aperture made using Equation (5).

| ID | \( P \), MPa | \( \mu \), Pa·s | \( \varepsilon \) F·m\(^{-1} \) | \( C_{SP} \), mV·MPa\(^{-1} \) | \( \zeta \) | \( k \), mD | \( 2w \), nm |
|----|-------------|---------------|-----------------|-----------------|--------|--------|--------|
| LS | 4           | \( 9.33 \times 10^{-4} \) | 7.00 \times 10^{-10} | -86.4           | -11.12 | 51 ± 1 | 782    |
| LS | 7           | \( 9.33 \times 10^{-4} \) | 7.00 \times 10^{-10} | -89.5           | -11.51 | 43 ± 1 | 718    |
| HS | 4           | \( 9.89 \times 10^{-4} \) | 6.97 \times 10^{-10} | -1.21           | -10.52 | 51 ± 1 | 782    |
| HS | 7           | \( 9.89 \times 10^{-4} \) | 6.97 \times 10^{-10} | -1.23           | -10.69 | 43 ± 1 | 718    |
The fracture aperture estimated from the sample permeability is consistent with the experimentally measured values reported for artificially fractured granite sample [66]. In their study, Luo et al. [66] reported the hydraulic aperture of order of 200 mm, which is three orders of magnitude larger than the values estimated for our sample using Equation (5). However, the values of hydraulic conductivity of 13 cm/s measured by Lou et al. [66], which translates into the permeability of c. $13 \times 10^6$ mD, is identical to that predicted by Equation (5), thus validating our approach and estimates.

To assess the effect of the surface conductivity on the zeta potential, we used a model of the zeta potential of silica-NaCl systems [67], for which the expected surface conductivities for the two ionic strengths tested in our experiments were evaluated to be $3.0 \times 10^{-9}$ S for $7.35 \times 10^{-3}$ M, and $4.5 \times 10^{-9}$ S for 0.7 M solution. To characterize the impact of the surface conductivity on the zeta potential, we used the dimensionless Dukhin number ($\text{Du}$), defined by

$$
\text{Du} = \frac{\sigma_s}{\sigma_w w}
$$

(12)

From the estimates of the fracture aperture and surface conductivity, and measured bulk solution conductivities, $\text{Du}$ translates to $7.9 \times 10^{-2}$ for $7.35 \times 10^{-3}$ M solutions at 4 MPa of confining pressure; $8.6 \times 10^{-2}$ for $7.35 \times 10^{-3}$ M solutions at 7 MPa; $1.9 \times 10^{-3}$ for 0.7 M solutions at 4 MPa and $2.0 \times 10^{-3}$ for 0.7 M solutions at 7 MPa. All estimated $\text{Du}$ for our experimental conditions clearly satisfied the condition for a negligibly small contribution of the surface conductivity to the total effective conductivity of an aqueous solution, hence, a negligible effect on the zeta potential: $\text{Du} \ll 1$. Therefore, for all tested solutions, the surface electrical conductivity could be neglected, and Equation (1) was appropriate for interpreting the zeta potential from the measured streaming potential coupling coefficient.

To evaluate the aquifer depths at which surface conductivity becomes significant, we considered a study [66] that reported a near-linear correlation between the confining pressure and the hydraulic aperture of an artificially fractured granite sample. Thus, using estimates of the surface conductivity value for low-salinity solutions (Table 3), we were able to evaluate the confining pressure and the corresponding aquifer depth, at which the surface conductivity becomes significant, satisfying the condition of $\text{Du} \approx 0.1$. The aperture that satisfies $\text{Du} = 0.1$ was calculated to be 310 nm using Equation (12), and the corresponding confining pressure was found to be 3.8 MPa, corresponding to an aquifer depth of 380 m, which is significantly deeper than the typical useful (from a resource point of view) depth of fractured gneiss aquifers, which varies between a few meters and a few hundred meters. At these depths, the impact of the surface conductivity on the zeta potential becomes substantial, hence, appropriate care should be taken to interpret the SP and ERT field data.

Future experimental work will aim at acquiring additional zeta potential values in systems comprising other specimens of the Lewisian gneiss and confining pressures higher than 7 MPa. These experiments will complement the data reported here and expand the range of tested conditions not covered in this work, thus, further improving our understanding of the zeta potential at the gneiss-water interfaces under conditions relevant to naturally fractured systems. The results will also be used to inform future surface complexation and molecular dynamics simulation studies aimed at describing gneiss-water systems under realistic conditions.

4. Conclusions

We report measurements of the zeta potential carried out for the first time on a fractured Lewisian gneiss sample saturated with NaCl solutions, artificial groundwater (AGW), artificial seawater (ASW) and under 4 MPa and 7 MPa of confining pressure. The results show that:
• Zeta potentials of gneiss are unique and dissimilar to sandstones, carbonates and even to individual gneiss constitutive minerals, i.e., mica and feldspar.
• The negative zeta potential decreases with increasing salinity when the sample is saturated with AGW and ASW, but the rate of the decrease is smaller compared to any other mineral.
• The negative zeta potential is independent of salinity when using NaCl; this feature is similar to what was observed with clayey sandstone [45] but the zeta potential of gneiss was found to be more positive compared with clayey sandstones saturated with NaCl.
• Significant amounts of feldspar and mica present in our gneiss sample were found to be responsible for the high sensitivity of the sample to the presence of divalent ions (Ca$^{2+}$, Mg$^{2+}$ and SO$_4^{2-}$) compared with quartz. Therefore, these ions were identified as PDIs for gneiss, making the sample respond to compositional and concentration variations in a fashion similar to that of carbonate samples.
• The reported values of gneiss $C_{SP}$ remained independent of the confining pressure, thus, suggesting that the surface electrical conductivity in gneiss aquifers could be neglected. A simple fracture model was developed to evaluate depths, and resulting in situ confining pressures, until which fracture aperture would remain large enough to neglect the surface electrical conductivity; this depth was found to be 380 m, which is much deeper than most water resources applications.

The reported dependence of the fractured gneiss zeta potential on confining pressure, composition and salinity of aqueous solutions can be used to identify newly formed fractures along naturally weathered pre-existing clacks from the measured signal, thus paving a way to a novel passive non-intrusive method of quantitative characterization of fracture network in crystalline rocks.

The above results suggest that interpretation of SP field data requires knowledge of the zeta potential values, which should be used to calculate $C_{SP}$ using Equation (1). Subsequently, using the computed values of $C_{SP}$ and voltages measured in the field, the expected pressure gradients can be quantified without any requirement of accounting for surface electrical conductivity. Finally, the obtained values of the pressure gradients will provide essential information on hydraulically active fractures (zero SP response results in zero pressure gradient, hence, implies no flow in that direction) and average fracture permeability (higher pressure gradients correspond to lower average fracture permeability).

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/app12010180/s1: S1: Geological Map Lewisian gneiss; S2: Summary of All Experimental Data and Error Analyses.

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