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On Permeability Prediction From Complex Conductivity Measurements Using Polarization Magnitude and Relaxation Time

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Abstract Geophysical length scales determined from complex conductivity (CC) measurements can be used to estimate permeability k when the electrical formation factor F is known. Two geophysical length scales have been proposed: (1) the specific polarizability ω normalized by the imaginary conductivity σ′′ and (2) the time constant τ multiplied by a diffusion coefficient Ds. The parameters ω and Ds account for the control of fluid chemistry and/or varying mineralogy on the geophysical length scale. We evaluated the predictability of two CC permeability models: (1) an empirical formulation based on ω normalized chargeability m and (2) a mechanistic formulation based on τ. The performance of the CC models was evaluated against measured k and further compared against that of well-established k estimation equations that use geometric length scales. Both CC models predict permeability within one order of magnitude for a database of 58 sandstone samples, with the exception of samples characterized by high pore volume normalized surface area Spor. Variations in ω and Ds likely contribute to the poor model performance for the high Spor samples, which contain significant dolomite. Two observations favor the implementation of the ω-based model over the τ-based model for field-scale k estimation: (1) a limited range of variation in ω relative to Ds and (2) ω field measurements are less time consuming to acquire relative to τ. The need for a reliable field-estimate of F limits application of either model, in particular the ω model due to a high power law exponent associated with F.

1. Introduction

The reliable in situ estimation of permeability remains one of the most challenging problems in hydrogeological characterization. Permeability can vary by orders of magnitude over short distances due to heterogeneity across multiple scales. Permeability is essential in the planning and implementation of water management projects including determining well yields, understanding recharge rates, and fluxes to surface water and designing storm water infrastructure. At environmental sites, permeability exerts a fundamental control on contaminant transport and determines the effectiveness of remediation strategies. Borehole tests to estimate permeability often do not fully capture site heterogeneity and may require upscaling to representative scales that determine flow and transport processes. Consequently, novel methods for measuring spatial variations of permeability at the field-scale are needed.

Indirect estimation of permeability is built upon equations that utilize a representative length scale associated with a dominant pore-scale dimension controlling fluid flow (Carman, 1939) coupled with an electrical formation factor F describing the ratio between tortuosity and porosity (Banavar & Johnson, 1987; Katz & Thompson, 1987, 1986). The models utilizing geometric length scales are summarized in Figure 1. Pape et al. (1987) proposed one such equation that is referred to as the PaRiS model. This model uses the inverse of the pore volume normalized surface area Spor as the representative length scale, with fitting parameters calibrated against a database composed of a wide range of sandstone samples. Another equation proposed by Banavar and Johnson (1987) is based on the Katz and Thompson (KT) model (Katz & Thompson, 1987). A dynamically interconnected pore radius Λ is derived from the mercury injection capillary pressure (MICP) method and used as the representative length scale in permeability estimation. In both cases, such
representative geometric length scales cannot be directly measured in situ, thereby limiting the application of these permeability prediction models at the field scale.

Geophysical measurements offer the possibility of indirectly determining representative length scales needed for in situ permeability prediction (Revil et al., 2012; Revil & Florsch, 2010; Slater, 2007). Using sensors placed on the surface and/or in boreholes, the variation of hydrogeological properties across multiple scales can be inferred from geophysical measurements (see Binley et al., 2015, for review). Complex conductivity (CC), often referred to as spectral-induced polarization (SIP), is an electrical geophysical method that provides measures that have been shown to be well correlated with representative geometric length scales controlling permeability (Binley et al., 2005; Scott & Barker, 2003; Slater, 2007). The low frequency (<1 kHz) CC response is determined by diffusive polarization mechanisms associated with ions in the electrical double layer (EDL) forming at the mineral-fluid interface. Both the magnitude and the relaxation time of this double layer polarization have been correlated with geometric length scales extracted from the grain or pore size distribution of the rock (Binley et al., 2005; Borner et al., 1996; Revil, 2013; Slater & Lesmes, 2002; Weller et al., 2010). Mechanistic and empirical models have been developed to explain the dependence of the CC parameters on geometric length scales derived from the pore or grain size distribution (Leroy et al., 2017, 2008; Revil et al., 2015; Weller et al., 2015a).

The link between CC parameters and representative geometric length scales has encouraged efforts to develop permeability prediction equations based on CC measurements as summarized in Figure 1. These models rely on a measure of the polarization strength (an imaginary conductivity $\sigma''$ or normalized charge-ability term $m_n$) or the dominant relaxation time (typically a characteristic time constant $\tau$) (Revil et al., 2015; Weller et al., 2015a). Unlike the geometric length scales $1/S_{por}$ and $\Lambda$, $\sigma''$ and $\tau$ are dependent on the pore filling fluid chemistry (Niu et al., 2016; Revil & Skold, 2011; Weller & Slater, 2012), surface mineralogy (Abuseda et al., 2016; Kruschwitz et al., 2016; Revil, 2012) and are a function of measurement frequency. Weller et al. (2011) introduced the concept of specific polarizability $\epsilon_p$ to represent the control of the fluid chemistry and/or mineralogy on polarization magnitude. Similarly, Revil (2013) defined values of a diffusion coefficient $D_1$ to accommodate such factors in a mechanistic model describing the CC of soils and rocks (Revil et al., 2015).

Accounting for the fluid conductivity and/or mineralogy controls on CC measurements allows equivalent geophysical length scales to be defined to replace the classical length scale appearing in the geometric models. These representative geometric length scales are defined as $\epsilon_p/\sigma''$ and $\tau D_1$ (Figure 1). They are coupled in permeability prediction equations with $F$, which is used to describe the porosity and tortuosity in the same way as the geometric models described above.

One such approach based on an electrical analog of the PaRiS model involves using a form of $\sigma''$ for $S_{por}$, as justified by the strong correlation between $\sigma''$ and $S_{por}$ observed in numerous studies (e.g., Kruschwitz et al.,

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**Figure 1.** Graphical overview of geometric and geophysical length scales in relation to mechanistic (Banavar & Johnson, 1987; Katz & Thompson, 1987; Revil et al., 2015) and empirical permeability models (Pape et al., 1987; Weller et al., 2015a). Mechanistic length scales have units of L or L² (i.e., m or m²). Empirical length scales have units of μm. The polarization magnitude model $k_T$ implicitly incorporates a single value of the specific polarizability, $\epsilon_p$. 

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Similarly, an electrical analog of the Katz and Thompson (KT) model has been proposed where \( D_1 \) is used as a proxy for \( \Lambda \), a substitution that is justified by the strong empirical correlation between \( \tau \) and pore radius observed in numerous studies (e.g., Revil et al., 2014). One challenge in using the equivalent geophysical length scales is uncertainty in the variation in \( c_p \) and \( D_1 \) associated with fluid chemistry and mineralogy. Using single values of \( c_p \) and \( D_1 \) to represent variations in the interfacial chemistry and mineralogy will limit the predictive capability of these permeability equations in natural settings where fluid chemistry and mineralogy vary (Kemna et al., 2012; Kruschwitz et al., 2010; Revil et al., 2012).

An inherent limitation of both the geometric (PaRiS, KT) and CC-based permeability predictions is that \( F \) must be known. Researchers have used MICP to estimate \( F \) (Amirtharaj et al., 2011; Katz & Thompson, 1987, 1986) with limited results due in part to unknown sample inhomogeneity (Amirtharaj et al., 2011). Reliable field-scale estimates of \( F \) are difficult (or impossible) to acquire unless pore fluid conductivity is very high (e.g., brine fluids) such that the surface conductivity is assumed negligible. Borner et al. (1996) first suggested that CC measurements could be used to estimate surface conductivity from \( \sigma^\infty \) measurements, allowing the measured real part of the conductivity to be corrected for surface conduction permitting a field-scale estimation of \( F \). Several researchers have subsequently examined the relationship between \( \sigma^\infty \) and the surface conductivity measured using laborious multisalinity measurements both for sandstone (Revil et al., 2015; Weller et al., 2013) and limestone (Halisch et al., 2014) rocks in an attempt to generalize models for \( F \) estimation.

In this paper, we present a study using 58 sandstone samples where extensive physical and geophysical laboratory data were acquired. We examine the predictive capability of two recently proposed models of permeability estimation and use these models “as-is” with the predetermined constants and coefficients defined in the original models. Within the context of these two models, we: (1) explore the relative importance of the length scales (geometric and geophysical) and \( F \) on permeability prediction; (2) consider the limitations of CC permeability prediction due to the uncertainty in the parameters \( c_p \) and \( D_1 \) required to define geophysical length scales; and (3) investigate the effectiveness of proposed methods to estimate \( F \) by correcting for surface conductivity at low salinities using \( \sigma^\infty \) such that these permeability models could be applied to field scale CC data.

### 2. Permeability Models

#### 2.1. PaRiS and Katz-Thompson (KT) Geometric Models

The PaRiS model is based on flow of fluid through a network of capillary tubes whereas the KT model is based on the application of percolation theory applied to a broad distribution of pore sizes. Central to both models is the definition of an effective hydraulic radius \( r_{\text{eff}} \) (\( \mu \text{m} \)) representative of the pore size controlling fluid flow. In the PaRiS equation, \( r_{\text{eff}} \) is related to \( S_{\text{por}} \):

\[
r_{\text{eff}} \approx \frac{2}{S_{\text{por}}},
\]

\[
S_{\text{por}} = S_t \rho_s \frac{1 - \phi}{\phi},
\]

where \( S_t \) (\( m^2/g \)) is the specific surface area from Brunauer-Emmett-Teller (BET) analysis, \( \phi \) is the porosity, and \( \rho_s \) is the density of the solids. In the KT equation, \( r_{\text{eff}} \) is equivalent to \( \Lambda \) (Banavar & Johnson 1987; Katz & Thompson 1986) as determined from MICP,

\[
r_{\text{eff}} \approx \Lambda.
\]

The PaRiS model is based on a modified Kozeny-Carman equation (Carman, 1939), where \( F \) replaces the ratio of the pore capillary tortuosity to the porosity (e.g., Guéguen & Palciauskas, 1994, p. 196). The PaRiS model also considers the fractal nature of internal surface roughness of sedimentary rocks. Pape et al. (1987) proposed the following model:

\[
k_{\text{PaRiS}} = \frac{475}{S_{\text{por}}^{3.13} F}
\]

where \( k_{\text{PaRiS}} \) is permeability predicted by the PaRiS model with originally presented units of Darcy (D) \( (1 \text{ D} = 9.869 \times 10^{-13} \text{ m}^2) \) and where \( S_{\text{por}} \) is in \( \mu \text{m}^{-1} \).
The KT permeability equation (Katz & Thompson 1986) is expressed as

\[ k_{KT} = \frac{I_c^2}{cF}, \]  

(5)

where \( k_{KT} \) (m²) is permeability predicted by the KT model and \( I_c \) (µm) is the equivalent pore diameter corresponding to the pressure at which the maximum incremental mercury intrusion occurs using MICP. The constant \( c \) is a scaling constant equal to 226. Katz and Thompson (1986) equated \( I_c \) to the inflection point on the MICP cumulative pore size distribution curve, considered to represent the threshold at which the pore space of a rock becomes hydraulically interconnected (i.e., sufficient fluid saturation for fluid flow). Johnson et al. (1986) reformulated this original model in terms of \( K \) (µm) and Banavar and Johnson (1987) equating a pore radius \( (I_c/2) \) to \( K \) by a scaling constant \( a \). This reformulation results in

\[ k_{KT} = \frac{\Lambda^2}{8F}. \]  

(6)

Modeling the pore network as a distribution of cylindrical pores of differing radii, Banavar and Johnson (1987) derived two scaling constants \( a \) (equal to 0.51 and 0.34) to represent different sizes of pores and multiplied \( I_c/2 \) by \( a \) to calculate \( \Lambda \) (i.e., \( \Lambda = alc/2 \)). Revil et al. (2014) equated equations (5) and (6) with \( c = 226 \) (equation (5)) to derive \( a \) equal to 0.19.

2.2. Equivalent Geophysical Length Scale Models

Weller et al. (2015a) present an empirical model for permeability estimation that uses the inverse of the imaginary conductivity \( \sigma'' \) as the representative length scale. This is justified by the linear proportionality between \( \sigma'' \) and \( S_{por} \) demonstrated by Borner et al. (1996) and Weller et al. (2015b, 2010) where the linear coefficient represents the specific polarizability \( c_p \) (i.e., \( \sigma'' = c_pS_{por} \)) (in Siemens, S). Based on 58 clean and shaly sandstones samples from 17 formations, Weller et al. (2015a) presented a new empirical equation linking \( \sigma'' \) and \( F \) to permeability

\[ k_{\sigma''} = \frac{2.66 \times 10^{-7}}{\sigma''^{0.66}} f_{5.35}. \]  

(7)

where \( k_{\sigma''} \) (m²) is the permeability predicted using \( \sigma'' \) (mS/m) at a frequency of 1 Hz. Weller et al. (2015a) implicitly incorporated a single value of \( c_p \) in equation (7), represented in the numerator. The value of \( c_p \) was previously found to be equal to \( 10 \times 10^{-12} \) S for 114 samples (Weller et al., 2010), and \( 30 \times 10^{-12} \) and \( 3 \times 10^{-12} \) S for sandstones from the Bahariya Formation with a high and low magnetic susceptibility (greater than and less than \( 100 \times 10^{-6} \) (dimensionless, SI units)), respectively (Abuseda et al., 2016). Using the methylene blue (MB) method to determine \( S_{por} \), Weller et al. (2015b) found \( c_p \) equal to \( 7.5 \times 10^{-12} \) S for 60 unconsolidated samples saturated with a conductive fluid with a conductivity close to 100 mS/m. Regardless of the sample or method used to determine \( S_{por} \), values of \( c_p \) vary by at most one order of magnitude.

Rather than relying on a single frequency measure of \( \sigma'' \), a Debye decomposition approach can be used to integrate over a range of frequencies to give a measure of the polarization strength that encompasses the frequency dependence of the CC response (Nordsiek & Weller, 2008). Weller et al. (2015a) also considered the normalized chargeability \( m_n \), obtained from a Debye decomposition where the equivalent predictive equation is

\[ k_{m_n} = \frac{8.69 \times 10^{-7}}{m_n^{0.79f_{5.38}}} \]  

(8)

where \( k_{m_n} \) (m²) is the permeability predicted using \( m_n \) (mS/m). Again, the numerator value of \( 8.69 \times 10^{-7} \) implicitly incorporates a single value of \( c_p \). Equations (7) and (8) will be referred to herein as geophysical polarization magnitude permeability models.

Revil et al. (2012) present a mechanistic model based on the KT model (equation (6)) for permeability estimation using the CC relaxation time \( \tau \) (s) (the exact definition of this parameter is discussed below) in conjunction with a diffusion coefficient \( D_1 \) (m²/s) as a representative length scale. This development is built on
work originally based on spherical particles in suspension (Schwarz, 1962), where the relaxation time $\tau$ is related to the grain size radius, $R$ by,

$$\tau = \frac{R^2}{2D_+},$$

with $D_+$ related to the effective ionic mobility $\mu_+$, temperature $T$, the Boltzmann’s constant $k_b$, and the charge of counterions in the Stern layer $q_+$ by the Nernst-Einstein relationship $D_+ = k_b T \mu_+ / |q_+|$. Based on different $\mu_+$ values for a shaly-sand (Vinegar & Waxman, 1984) and unconsolidated sand sediments (Koch et al., 2011), Revil (2013) and Revil et al. (2012) proposed two values of $D_+$: $1.3 \times 10^{-9}$ m$^2$/s for clean sands and $3.8 \times 10^{-12}$ m$^2$/s for clayey material. Revil et al. (2012) proposed that $\tau$ is related to $\Lambda$ by

$$\tau = \frac{\Lambda^2}{2D_+}.$$  

Substitution of equation (10) into equation (6) yields the permeability prediction model proposed by Revil et al. (2012)

$$k_s = \frac{\tau D_+}{4F}.$$  

The application of equation (11) for permeability prediction is reliant on an accurate estimate of $D_+$. The $k_s$ model will be referred to herein as the geophysical time constant permeability model.

3. Sample Descriptions and Methods

Our sample database consists of 58 sandstone samples where hydraulic and physical property measurements (permeability, BET specific surface area, MICP pore size distribution, porosity) as well as CC spectral data and/or parameters are recorded.

3.1. Santa Susana and Hydrite Samples

The majority of our database is composed of 43 previously unpublished samples from two sites: the Santa Susana Field Laboratory (California, USA) with 22 samples and the Hydrite Chemical Company (Wisconsin, USA) with 21 samples. The Santa Susana site is a Cretaceous age deep-sea turbidite deposit composed primarily of sandstone with minor interbeds of shale and siltstone and occasional conglomerate of similar mineralogy. The sedimentary rocks at the Hydrite site are Cambrian to Ordovician in age and are associated with a variety of marine, shoreface, and terrestrial depositional environments. Consequently, the lithology types encountered at the Hydrite site are more heterogeneous including sandstones with variable grain size, mineralogy, and amounts of subordinate clay and silt, shales, siltstones, and dolostones.

A triple tube coring system was used with a coring diameter of 2.4 in (6.1 cm) (i.e., HQ3) at both sites. Two boreholes were cored at Santa Susana site, designated RD109 and C3. At Hydrite site, two boreholes were again cored, designated MP24S and MP25S. Cores were preserved to retain moisture content and reduced exposure to oxygen on site and sent to the laboratory for subcoring to a diameter of 3.8 cm. Smaller rock samples were subsampled at the same depths as the subcores, such that physical measurements could be collected and compared on collocated samples.

At Santa Susana site, 11 subcores were prepared from borehole RD109 over approximately a 105 m interval (30–135 m below ground surface (bgs)) and 11 subcores were prepared from borehole C3 over approximately a 35 m interval (7–42 m bgs). At Hydrite site, five horizontal subcores (designated 2H in Table 1) and seven vertical subcores (designated 1V in Table 1) were prepared from borehole MP24S over a 20 m interval between 34 and 54 m bgs. Four horizontal subcores and five vertical subcores were prepared from borehole MP25S over an approximately 14 m interval from 40 to 54 m bgs. Within this paper, horizontal and vertical subcores at Hydrite site were considered independent samples such that this analysis does not consider any effects of anisotropy.

Klinkenberg corrected (Klinkenberg, 1941) gas permeability was measured on the subcores. The subcores were saturated by applying a vacuum to at least $5 \times 10^{-2}$ mbar, adding the saturating fluid, and then applying 34.5 bar (500 psi) for 20 min. The fluid was boiled and infused with nitrogen gas prior to saturation.
<table>
<thead>
<tr>
<th>Sample name</th>
<th>Rock type and/or formation location</th>
<th>Physical properties</th>
<th>CC parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$S_{\text{ve}}$ ($\mu m^{-1}$)</td>
<td>$l_c$ ($\mu m$)</td>
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**Table 1**

Physical Properties and CC Parameters for Samples

**Physical Properties**

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<tr>
<th>Sample name</th>
<th>Rock type and/or formation location</th>
<th>$S_{\text{ve}}$ ($\mu m^{-1}$)</th>
<th>$l_c$ ($\mu m$)</th>
<th>Permeability (m²)</th>
<th>Porosity (%)</th>
<th>$\phi$ (mS/m)</th>
<th>$F$ (-)</th>
<th>$\phi^2$ (mS/m)</th>
<th>$\rho_0$ (mS/m)</th>
<th>$\tau_p$ (s)</th>
<th>$\tau_{\text{min}}$ (s)</th>
</tr>
</thead>
</table>

**CC parameters**

(Optional)

**Notes**

1. For the calculation of $\phi^2$, we used $\phi^2 = \rho_0 \rho_1$.
2. The values with an asterisk (*) indicate the calculated values.
3. The values without an asterisk indicate the measured values.

*References*

2. Abuseda et al. (2016)
to minimize air bubbles within the pores. Gravimetric porosity was calculated from the mass difference between saturated and dry subcores divided by submersible volume. MICP (up to 2,068 bar/30,000 psi), nitrogen-gas BET and X-ray diffraction (XRD) data were acquired on the colocated smaller samples and used to determine specific surface area and mineralogical composition, respectively.

CC measurements were collected on subcores saturated first with a low salinity solution and subsequently with a high salinity solution. The low salinity solution contained calcium chloride and magnesium sulfate in concentrations similar to the site groundwater and had a fluid conductivity of 66.5 mS/m. The high salinity solution was a sodium chloride solution with a fluid conductivity of 8 S/m. The CC measurements of subcores saturated with the high salinity solution were used to determine the electrical formation factor \( F \).

CC measurements were acquired using sample holders designed, tested, and calibrated following the procedure in Kemna et al. (2012) to remove measurement artifacts that can occur due to the electrode positions. A Portable Spectral-Induced Polarization (PSIP) instrument (Ontash & Ermac Instruments, NJ, USA) recorded CC over a frequency range of \( 10^{2}–10^{3} \) Hz, with five measurements per decade of frequency, for a total of 36 measurements. To avoid errors due to temperature fluctuations, the measurements were collected on sample holders placed in an environmental chamber set to 25°C. To ensure equilibration of temperature and ion exchange processes between the saturating pore fluid and mineral surface, measurements were collected twice daily (for approximately 1 week) until the difference between a previously collected measurement of both the resistance and phase was less than 2%; data collected at 1 Hz were used to quantitatively compare successive measurements, but the entire spectrum was visually inspected for consistency. Resistance and phase data were converted to complex conductivity using the geometric factor for the sample holder, which was experimentally determined from measurements on water samples of precisely known complex conductivity.

### 3.2. Previously Published Samples

We supplement this extensive new data set with 15 samples from multiple sources including: five Eocene sandstone samples of the Shahejie Formation (CS samples, China) (Zhang & Weller, 2014), seven samples of the Cretaceous Bahariya Formation (Egypt) (Abuseda et al., 2016), and three unpublished samples from various locations in Germany (Bentheimer, Elbe-sandstone, Green sand). Physical and geophysical measurements were available for our analysis including BET derived \( S_{m} \), permeability, \( F \), and CC spectra.

### 3.3. Analysis of the CC Spectra

A characteristic relaxation time is typically used to represent CC data. Here we used two characteristic \( \tau \) values: \( \tau_{\text{mean}} \) which is calculated from Debye decomposition and takes into account the entire CC spectrum, and \( \tau_{\text{pc}} \) which is associated with a characteristic frequency \( f_{\text{char}} \). Since different behaviors can be observed in the CC spectrum, the procedure to choose \( f_{\text{char}} \) can vary (Revil et al., 2015; Schwarz, 1962). Within our sample database, the \( \sigma'' \) spectra exhibited either a well-defined peak frequency \( f_{p} \) or a corner frequency \( f_{c} \), where \( \sigma'' \) rapidly decreased toward lower frequencies (Revil et al., 2015). If the \( \sigma'' \) spectrum contained a well-defined peak frequency, a parabola was fit to the five data points centered on this peak and used to

### Table 1. (continued)

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Rock type and/or formation</th>
<th>location</th>
<th>Physical properties</th>
<th>CC parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>( S_{m} ) (( \mu m^{-1} ))</td>
<td>( l_{c} ) (( \mu m ))</td>
</tr>
<tr>
<td>22H Bahariya</td>
<td></td>
<td></td>
<td>37.81</td>
<td>5.17</td>
</tr>
<tr>
<td>28H1 Bahariya</td>
<td></td>
<td></td>
<td>1.34</td>
<td>8.41</td>
</tr>
<tr>
<td>43H1 Bahariya</td>
<td></td>
<td></td>
<td>2.49</td>
<td>16.27</td>
</tr>
<tr>
<td>53H2 Bahariya</td>
<td></td>
<td></td>
<td>3.89</td>
<td>14.65</td>
</tr>
<tr>
<td>55 Bahariya</td>
<td></td>
<td></td>
<td>19.66</td>
<td>10.68</td>
</tr>
<tr>
<td>Other BH6 Bentheimer</td>
<td></td>
<td></td>
<td>4.12</td>
<td>35.72</td>
</tr>
<tr>
<td>ES-14 Elb-sandstone</td>
<td></td>
<td></td>
<td>1.06</td>
<td>31.75</td>
</tr>
<tr>
<td>GR1 Green sand</td>
<td></td>
<td></td>
<td>31.33</td>
<td>36.26</td>
</tr>
</tbody>
</table>

* A corner value of \( \tau \) was used for this sample.
estimate the apex, equal to \( f_p \). If there was no clear peak, a spline was fit to the data points and \( f_p \) was chosen as the intersection of two linear fits to the data following the method of Revil et al. (2015). The time constant was then calculated from \( \tau_{pc} = 1/2nf_{pc} \), where \( p \) or \( c \) indicates either peak or corner. Examples of peak and corner selection for two samples are shown in supporting information Figure S1.

To compare the predictive capabilities of the different models, the average absolute logarithmic deviation between predicted parameters, \( p^* \) and measured parameters \( p \) was determined by:

\[
d = \frac{1}{n} \sum_{j=1}^{n} |\log_{10} p_j - \log_{10} p_j^*|,
\]

where \( d \) represents the average, or mean absolute deviation on a logarithmic scale. A \( d \) value equal to 1 indicates that the average misfit between the predicted data point and the measured parameter is one order of magnitude. In regards to permeability model estimates, one order of magnitude above or below measured values (i.e., \( d = 1.0 \)) is commonly considered to yield an acceptable estimation.

### 4. Results

Table 1 summarizes all physical properties and CC parameters for the 58 samples used in this study. Permeability ranges from \( 9.21 \times 10^{-17} \) to \( 2.84 \times 10^{-12} \) m\(^2\) (0.0933–2,875 mD), spanning more than four orders of magnitude. The other physical properties also demonstrate the variation in the pore geometry among these sandstones: porosity ranges from 10.1% to 24.0%; \( F \) ranges from 11.7 to 59.6; \( S_{por} \) ranges from 1.06 to 86.03 \( \mu \)m\(^2\). We report \( \sigma^2 \) at a frequency of 1 Hz to facilitate direct comparison with previous studies (Weller et al., 2010, 2015a, 2015b).

#### 4.1. PaRiS and Katz-Thompson (KT) Geometric Models

We first test the predictions of the PaRiS model (equation (4)) and KT model using 0.19 \( l_2 = \Lambda \) in (equation (6)) based on the geometric length scales, 1/\( S_{por} \), and \( \Lambda \), respectively (Figure 2). As we are only interested in examining the predictive capabilities of these models in a field setting, no effort has been made to optimize the coefficient or scaling constants in the equations. The 1:1 line is shown as a solid line in Figure 2 and the dashed lines represent one order of magnitude of variation above and below the 1:1 line.

The colorscale of the data points represents \( S_{por} \) and highlights the general trend of a decrease in measured permeability with an increase in \( S_{por} \). Using the PaRiS model (Figure 2a), permeability of the high \( S_{por} \) samples is underpredicted, with several data points deviating by more than one order of magnitude. The permeability of several low \( S_{por} \) samples is overpredicted.

Using the KT model (Figure 2b), most permeabilities are underpredicted but fall within one order of magnitude of the actual measured values. Based on the \( d \) values, the KT model (equation (6) using \( \Lambda \) \( d = 0.729 \)) yields a better prediction of permeability than the PaRiS model \( d = 1.023 \).

![Figure 2. Permeability predictions using geometric length scales (a) PaRiS model per equation (4) \( d = 1.023 \) and (b) KT model with \( \Lambda \) per equation (6) \( d = 0.729 \).](image-url)
Table 2. These plots show the expected trends: permeability decreases with increasing \( \sigma_{1\text{Hz}}^e \) and \( m_n \); and permeability increases with increasing \( \tau_{PC} \) and \( \tau_{mean} \) (based on regression exponents presented in Table 2). The relationship between \( F \) and permeability is shown in Figure 4. Similar to Figure 3, a linear regression on the logarithmic values was used to determine the coefficients for a power law relationship between \( F \) and measured permeability. The relationship between \( F \) and measured permeability shows, as expected, that measured permeability decreases with \( F \) (Figure 4).

4.2.2. Polarization Magnitude Geophysical Models

Predictions of permeability for the polarization magnitude geophysical models are shown in Figure 5. Permeability is well-predicted (within one order of magnitude) from equations (7) and (8) (Figure 5) with the exception of samples with a high \( S_{PC} > 60 \mu \text{m}^{-1} \) (see Table 1). The \( d \) values for the polarization magnitude geophysical models demonstrate less than within one order of magnitude misfit for single frequency \( (\sigma_{1\text{Hz}}^e) \) (0.770) versus multifrequency \( (m_n) \) (0.807) CC parameters (Table 3). The polarization magnitude geophysical models are a better predictor of permeability than the PaRiS model (equation (4)) based on misfit alone (1.023 versus values stated above) (Table 3).

A single value of \( c_p \) (as in equations (7) and (8)) may not represent all samples, contributing to the poor prediction of permeability for the high \( S_{PC} \) samples. Applying a linear least squares regression to \( \sigma_{1\text{Hz}}^e \) versus \( S_{PC} \) yields an optimal \( c_p \) value of 4.9 \( \times \) 10\(^{-10} \) S, although the coefficient of determination is low \((R^2 = 0.40)\). In contrast, individually calculating \( c_p \) for each sample shows a large range of potential \( c_p \) values based on the mean of 16.7 \( \times \) 10\(^{-12} \) S and standard deviation of 14.5 \( \times \) 10\(^{-12} \) S (maximum = 88.9 \( \times \) 10\(^{-12} \) S, minimum = 1.6 \( \times \) 10\(^{-12} \) S).

4.2.3. Time Constant Geophysical Model

Permeability predictions from equation (11) using the time constants \( \tau_{PC} \) and \( \tau_{mean} \) with \( F \) directly determined from the high salinity data are shown in Figures 6a and 6b, respectively. A single value of \( D_+ \) equal to 3.8 \( \times \) 10\(^{-12} \) m\(^2\)/s (clayey-sand) was assumed since our samples are best represented as clayey sandstones rather than clean sandstones. Permeability is well-predicted (within one order of magnitude) again with the exception of the samples with high \( S_{POD} > 60 \mu \text{m}^{-1} \). In addition, a few low \( S_{POD} \) are also outliers. Figure 6 demonstrates that the overall predictions of \( k \) are similar when using either \( \tau_{PC} \) or \( \tau_{mean} \) (Table 3).

A complicating factor in the use of equation (11) to calculate \( k \) is determining an appropriate value of \( D_- \). If the two proposed values of 1.3 \( \times \) 10\(^{-7} \) m\(^2\)/s for clean sands and 3.8 \( \times \) 10\(^{-12} \) m\(^2\)/s for clayey material (Koch et al., 2011; Revil, 2013; Revil et al., 2012) best represent \( D_- \) values for all samples, then by rearranging equation (11), the values of \( \log 4Fk \) plotted versus \( \log \tau_{PC} \) should fall on two distinct lines; the slopes of which corresponds to \( D_- \). However, as seen in Figure 7a, this is not the case; there is a large scatter in this plot and the samples do not consistently fall along either line similar to the observations of...
Weller et al. (2016). The large scatter indicates a wide range of $D_1$ which does not appear to be associated with individual formations. By calculating individual values of $D_1$ for each sample, using equation (11) and the known values for $F$, $k$, and $\tau_{pc}$, we can group the samples and determine a more appropriate value of $D_1$. From Figure 7b, higher $S_{por}$ samples are better represented by a lower $D_1$ and lower $S_{por}$ samples are better represented by a higher $D_1$. We thus define three linearly spaced groups of samples based on the values of $S_{por}$ (0–29 $\mu$m$^{-1}$; 30–57 $\mu$m$^{-1}$; 58–86 $\mu$m$^{-1}$) and calculate $D_1$ for each group. The values for $D_1$ determined from this approach are: $4.96 \times 10^{-12}$ m$^2$/s for the low $S_{por}$ samples, $2.13 \times 10^{-12}$ m$^2$/s for the midrange $S_{por}$ samples and $6.98 \times 10^{-14}$ m$^2$/s for the high $S_{por}$ samples. The new lines representing these $D_1$ values are plotted alongside the data in Figure 7b. Using these new $S_{por}$-specific $D_1$ values, the permeability was predicted from equation (11); the results from the updated permeability model are shown versus the measured permeability in Figure 8. Unsurprisingly, using these $S_{por}$-specific $D_1$ values improves the permeability prediction relative to using a single assumed value of $D_1$ alone. However, some outliers are not reconcilable using such variations in $D_1$.

5. Discussion

We have performed a first-of-a-kind comparative analysis of two recently proposed approaches to permeability prediction on 58 sandstone samples using geophysical length scales that can be defined from complex conductivity data and analyzed the uncertainty in these predictions that can be partly attributed to the variation in the electrochemical parameters $D_1$ and $c_p$. We have also evaluated permeability predictions from “as-is” geophysical length scales against predictions from geometric length scales. Using the geometric length scales $1/S_{por}$, and $\Lambda$ in these formulations, permeability is reasonably predicted (Figure 2), although the coefficients in the PaRiS model (Figure 2a) are not optimal values for our database. The geophysical length scales $c_p/\sigma_{1Hz}$ and $c_p/m_n$ (equations (7) and (8); Figure 5) and $\tau_{pc}D_+$ and $\tau_{mean}D_+$ (equation (11); Figure 6) were shown to provide estimates of permeability to within one order of magnitude for most samples. In general, the geophysical length scale models perform as well as the geometric length scale permeability models based on analysis of residuals (Table 3). Amongst the geophysical permeability models, the $\tau_{mean}$-based model performs best relative to $\tau_{pc}$, $\sigma_{1Hz}$, and $m_n$ models. Interestingly, all geophysical models outperform the PaRiS model based on the geometric length scale.

Many samples with high $S_{por}$ appear as outliers in the geophysical length scale models while $S_{por}$ does not have as large of an influence on outliers in the geometric length scale models. In the geophysical models

![Figure 5](file://path/to/image.png)  
Figure 5. Permeability predicted for the polarization magnitude geophysical model using (a) $\sigma_{1Hz}$ (equation (7)) ($d = 0.770$) and (b) $m_n$ (equation (8)) ($d = 0.807$).
permeability is overestimated for high Spor samples. The most likely cause of the permeability overestimation for high Spor samples is mineralogical variation and its associated impact on $c_p$ and $D_1$. X-ray diffraction results show that high Spor samples are characterized by the presence of dolomite (Table 1), which might cause variations in $c_p$ relative to quartz (Halisch et al., 2014) given that $c_p$ accounts for the control of the interfacial chemistry and sample mineralogy on the polarizability. The high Spor samples containing dolomite (supporting information Figure S2) are interpreted to have a lower $c_p$ than other samples which is similar to finding reported by Halisch et al. (2014) on carbonates. Similarly, these samples have a lower $D_1$ than the assumed single value for a clayey-sand ($3.8 \times 10^{-12} \text{m}^2/\text{s}$).

Mineralogical variations may not be responsible for the overprediction of permeability in high Spor samples in the polarization magnitude geophysical model. One alternative possibility is that the calibration parameters in this model cause this effect, e.g., if the power law exponent on $r_0$ is too low. To assess this possibility, a multilinear regression using the form of equation (7) was performed to solve for the numerator and the $r_0$ exponential coefficients (supporting information Figure S3b). For the 58 samples in this database, it was found that the $r_0$ exponential coefficient increased only slightly from 0.66 to 0.96. Additionally, high Spor samples still appear as outliers, suggesting a structural and/or mechanistic behavior in these sandy dolostones that differs from other samples in our database. Interestingly, calibrating the coefficients in the polarization magnitude geophysical model (equation (7)) results in a slightly better $d$ value ($d = 0.515$) than the calibration of equation (11) using the Spor-specific $D_1$ coefficients ($d = 0.539$).

To further assess the impact of mineralogical variability on permeability prediction, $d$ values were calculated for all reference equations by omitting samples with high Spor ($\geq 60 \mu \text{m}^{-1}$) (Table 3). Residuals for permeability predictions using geometric parameters Spor and $K$ remain fairly constant, with a slight improvement

### Table 3

Summary $d$ Misfits for Permeability Predictions Using Reference Equations

<table>
<thead>
<tr>
<th>Key parameter type</th>
<th>Description</th>
<th>Equation</th>
<th>Eqn #</th>
<th>Figure #</th>
<th>$d$ misfits</th>
</tr>
</thead>
<tbody>
<tr>
<td>Geometric</td>
<td>PaRiS model</td>
<td>$k_{\text{PaRiS}} = \frac{475}{\text{Spor}^{0.77}}$</td>
<td>4</td>
<td>2a</td>
<td>1.023</td>
</tr>
<tr>
<td></td>
<td>Katz and Thompson (KT)-type</td>
<td>$k_{\text{KT}} = \frac{A}{F}$</td>
<td>6</td>
<td>2b</td>
<td>0.729</td>
</tr>
<tr>
<td></td>
<td>Polarization magnitude $\sigma''$</td>
<td>$k_{\sigma''} = \frac{2.66 \times 10^{-7}}{F}$</td>
<td>7</td>
<td>5a</td>
<td>0.770</td>
</tr>
<tr>
<td></td>
<td>Polarization magnitude $m_{\text{ini}}$</td>
<td>$k_{m_{\text{ini}}} = \frac{8.4 \times 10^{-7}}{F}$</td>
<td>8</td>
<td>5b</td>
<td>0.807</td>
</tr>
<tr>
<td></td>
<td>Time constant $t_{\text{pc}}$</td>
<td>$k_{t_{\text{pc}}} = \frac{t_{\text{pc}}D_1}{4F}$</td>
<td>11</td>
<td>6a</td>
<td>0.779</td>
</tr>
<tr>
<td></td>
<td>Time constant $t_{\text{mean}}$</td>
<td>$k_{t_{\text{mean}}} = \frac{t_{\text{mean}}D_1}{4F}$</td>
<td>11</td>
<td>6b</td>
<td>0.707</td>
</tr>
<tr>
<td></td>
<td>Time constant $t_{\text{pc}}$ and $D_1$ (Spor)</td>
<td>$k_{t_{\text{pc}}} = \frac{t_{\text{pc}}D_1}{4F}$</td>
<td>11</td>
<td>8</td>
<td>0.539</td>
</tr>
</tbody>
</table>

(Figures 5 and 6), permeability is overestimated for high Spor samples. The most likely cause of the permeability overestimation for high Spor samples is mineralogical variation and its associated impact on $c_p$ and $D_1$. X-ray diffraction results show that high Spor samples are characterized by the presence of dolomite (Table 1), which might cause variations in $c_p$ relative to quartz (Halisch et al., 2014) given that $c_p$ accounts for the control of the interfacial chemistry and sample mineralogy on the polarizability. The high Spor samples containing dolomite (supporting information Figure S2) are interpreted to have a lower $c_p$ than other samples which is similar to finding reported by Halisch et al. (2014) on carbonates. Similarly, these samples have a lower $D_1$ than the assumed single value for a clayey-sand ($3.8 \times 10^{-12} \text{m}^2/\text{s}$).

Mineralogical variations may not be responsible for the overprediction of permeability in high Spor samples in the polarization magnitude geophysical model. One alternative possibility is that the calibration parameters in this model cause this effect, e.g., if the power law exponent on $r_0$ is too low. To assess this possibility, a multilinear regression using the form of equation (7) was performed to solve for the numerator and the $r_0$ exponential coefficients (supporting information Figure S3b). For the 58 samples in this database, it was found that the $r_0$ exponential coefficient increased only slightly from 0.66 to 0.96. Additionally, high Spor samples still appear as outliers, suggesting a structural and/or mechanistic behavior in these sandy dolostones that differs from other samples in our database. Interestingly, calibrating the coefficients in the polarization magnitude geophysical model (equation (7)) results in a slightly better $d$ value ($d = 0.515$) than the calibration of equation (11) using the Spor-specific $D_1$ coefficients ($d = 0.539$).

To further assess the impact of mineralogical variability on permeability prediction, $d$ values were calculated for all reference equations by omitting samples with high Spor ($\geq 60 \mu \text{m}^{-1}$) (Table 3). Residuals for permeability predictions using geometric parameters Spor and $K$ remain fairly constant, with a slight improvement

**Figure 6.** Permeability predicted from the time constant geophysical model (equation (11)) using (a) $t_{\text{pc}}$ ($d=0.779$) and (b) $t_{\text{mean}}$ ($d=0.707$) ($D_1 = 3.8 \times 10^{-12} \text{m}^2/\text{s}$).
in the PaRIS model and a slight decrease in for the KT-type model. All predictions using geophysical parameters $s_{\text{pc}}, s_{\text{mean}}, r_{00}$; and $m_n$ improve, with the largest improvement in the $k_1$ models not using an $S_{\text{por}}$-specific $D_{\ldots}$. Integration of lithological information in geophysical permeability models could provide an improvement over existing models.

What explains a smaller $c_p$ and $D_{\ldots}$ for our samples containing dolomite? Weller et al. (2016) suspect that, for samples with increasing clay content and increasing specific surface area, a decrease in the ion mobility results in a smaller $D_{\ldots}$. They suppose a stronger binding of ions at the surfaces dominates the diffusion in smaller pores associated with clay minerals, thereby reducing mobility. Similar to samples with high clay content, the high $S_{\text{por}}$ in these sandy dolostones may be reducing the effective ionic mobility $j_{\ldots}^n$, thereby reducing $D_{\ldots}$. In regards to smaller $c_p$ values in the sandy dolostone samples, the higher intracrystalline pore space (inferred from high $S_{\text{por}}$ measured in these samples) may be weakly polarized relative to silicates. Alternatively, smaller $c_p$ values may be attributed to the pH dependency of the CC response, specifically the difference in the point of zero charge (PZC) for dolomite versus silica. For Berea sandstones Lesmes and Frye (2001) found a reduced $r_{00}$ response at pH = 3, which corresponds to the PZC for silica; Skold et al. (2011) found a similar pH dependence for silica sands. In contrast, we compare the PZC of calcite, which has similar values to dolomite (Pokrovsky et al., 1999). The PZC of calcite has been reported within the pH range of 8–9.5 (Heberling et al., 2011; Somasundaran & Agar, 1967). Our CC data collection was performed at near neutral pH, i.e., much closer to the PZC for calcite relative to the PZC for silica. The $\sigma^n$ would then

Figure 7. (a) Individual diffusion coefficients ($D_{\ldots}$) for the sample database in Table 1 classified by lithological unit. Predicted values of $D_{\ldots} = 3.8 \times 10^{-12} \text{ m}^2/\text{s}$ for clayey-sand and $D_{\ldots} = 13 \times 10^{-12} \text{ m}^2/\text{s}$ for clean sands are shown by dashed and dotted lines respectively. (b) Same as Figure 7a with $S_{\text{por}}$ designated by the legend color scale. Binning samples by $S_{\text{por}}$ into three groupings and solving for the best fit $D_{\ldots}$ results in the three values shown (blue, cyan, and yellow lines).
presumably be reduced for calcite when compared to silicates. However, several researchers have suggested that pH has no direct influence of electrical surface properties in calcites, but rather can be used to determine speciation of Ca$^{2+}$ and CO$_3^{2-}$, which are the potentially determining ions (Amankonah et al., 1985; Foxall et al., 1979; Thompson & Pownall, 1989). While we suggest that the PZC plays a role in the reduced $c_p$ for the sandy dolostone samples, the mechanisms clearly need further study.

A major limiting factor in the application of both the geometric and geophysical length scale $k$ models is the use of the formation factor $F$. Since $F$ is not directly measurable from borehole electrical logging or electrical imaging surveys, the application of these models to fieldscale data is limited. We observe that $k$ has a strong dependence on $F$ (Figure 4), and so its inclusion in the permeability models is important. One solution to this limitation is to estimate $F$ from electrical parameters that can be measured in the field. For instance, the predicted electrical formation factor $F_p$ can be estimated from the real conductivity ($\sigma_0$) if $\sigma_w$ is known (i.e., $F_p=\sigma_w/\sigma_0$). This estimation assumes that the surface conduction is small, a questionable assumption even in relatively clean sands. In our database, $F$ is underestimated by $F_p$ for all but two samples (Figure 9a, $d=0.558$).

In order to overcome this critical limitation of electrical measurements for permeability estimation, Borner et al. (1996) first suggested that $F_p$ can be calculated as

$$F_p = \frac{\sigma_w}{\sigma_0 - \sigma_{surf}^\prime} = \frac{\sigma_w}{\sigma_0 - \sigma_{surf}^\prime} l$$

where $l$ is a linear coefficient that is assumed to relate $\sigma_0^\prime$ to the real part of the surface conductivity ($\sigma_{surf}^\prime$). Borner et al. (1996) suggested $l$ ranges between 0.01 and 0.15. More recent studies on larger databases have found $l = 0.042$ ($R^2 = 0.91$) (Weller et al., 2013) or $l = 0.037$ ($R^2 = 0.79$) (Revil et al., 2015) for sandstone samples. Based on the 58 samples in this study, the linear least squares regression gives $l = 0.044$ ($R^2 = 0.75$). We use $l = 0.042$ (Weller et al., 2013) to calculate $F_p$ for the 58 sandstone samples without any model calibration. This improves the prediction of $F$ relative to using $F_p=\sigma_w/\sigma_0$ although $F_p$ still mostly underpredicts true $F$ (Figure 9b, $d=0.403$). Using $l = 0.044$, the prediction improves slightly (Figure 9c, $d=0.336$). The impact of an incorrect estimate of $F$ on the permeability prediction depends in part on the relative weighting of $F$ in the models. For example, in the geophysical polarization magnitude equation given by equations (7) and (8), $F$ is raised to the power 5.35 and 5.38, respectively, resulting in large deviations from true permeability values when $F$ is in error. In contrast, $F$ is only raised to the power 1 in the geophysical time constant model (equation (11)). In these models, deviations from the true permeability do not increase so markedly when $F_p$ is used in place of $F$.

Figure 8. Using the time constant geophysical model (equation (11)) to solve for predicted permeability with three discrete values of $D_1$ defined based on $S_{por}$ (Figure 7b). $t_{pc}$ is used as the time constant.

Figure 9. Predicted electrical formation factor $F_p$ versus measured formation factor $F$ where (a) $F_p=\sigma_w/\sigma_0$ ($d = 0.558$), (b) $F_p$ is calculated from equation (13) using $l=0.042$ ($d=0.403$) (Weller et al., 2013) and (c) $F_p$ is calculated from equation (13) using $l=0.044$ ($d=0.336$).
The selection of $\sigma''_{100Hz}$ in the polarization magnitude model (equation (7)) is arbitrary and primarily based on facilitating integration of a large number of independently determined data sets into model calibration (Weller et al., 2015a). To evaluate the effect of this choice on the permeability estimates, a sensitivity analysis was performed to quantify variations in $d$ as a function of measurement frequency when permeability prediction is based on equation (7) (supporting information Figure S4). Permeability is predicted to within one order of magnitude regardless of the frequency at which $\sigma''$ is selected and the variability of $d$ is small (0.757–0.788). In fact, the 1 Hz frequency has the median $d$ value, suggesting that permeability would be slightly better predicted at some other frequencies.

Field-based acquisition of CC parameters is increasing with the advent of new instrumentation and measurement techniques (Kemna et al., 2014; Slater et al., 2018). Considering the impracticality of applying the geometric permeability models in the field, the geophysical models might provide a solution to field-scale estimation of permeability. This study has highlighted that opportunity, as noted by the fact that the geophysical length scale in the polarization magnitude model provides a better prediction of permeability than the geometric length scale in the PaRiS model. However, significant challenges remain with the CC-based estimation of permeability. First and foremost is the limitation that $F$ is not readily measurable at the field scale. Beyond this, each geophysical model has limitations. In the polarization magnitude geophysical model (equations (7) and (8)), variations in fluid chemistry, are accounted for within $c_p$ whereas such factors control $D_1$ in the geophysical time constant model (equation (11)). Representative values of these parameters must be assumed for field-scale implementation of the models. Unsurprisingly, an improvement was seen in the time constant model with the application of three $S_{por}$-specific $D_1$ values (Figure 8 and Table 3). However, the diffusion coefficients will be unknown in the field, so this is not a practical approach to improving model performance. For our 58 samples, $S_{por}$-specific $D_1$ values vary by two orders of magnitude. In contrast, the majority of individual $c_p$ values vary by less than one order of magnitude. All other factors being equal, permeability prediction on high $S_{por}$ samples using the time constant geophysical model with a single value of $D_1$ would be more uncertain relative to polarization magnitude models utilizing a single $c_p$ value.

While CC data acquisition is increasing at the field scale, it is currently very challenging to acquire reliable broadband frequency spectra. In contrast, it is relatively straightforward to measure a single frequency $\sigma''$ and much less time consuming. All geophysical parameters used (i.e., $m_{\text{mrt}}$, $\tau_{pc}$, and $\tau_{\text{mean}}$) except $\sigma''$ necessitate collection of multifrequency data, with not much improvement in permeability prediction (Table 3). In addition, not all CC spectra indicate a peak frequency and the nonprescriptive selection of $\tau_{pc}$ in the time constant geophysical model adds further uncertainty to this approach, as there are several ways in which to estimate this parameter. These observations support the use of $\sigma''$ to estimate permeability at the field scale from electrical measurements.

This study explored the predictive capability of previously proposed PaRiS, KT-type, polarization magnitude, and time constant permeability models that utilize geometric and geophysical length scales. Limitations of the models, specifically with respect to field-scale applications, were considered. The primary limitation of all investigated models is the requirement of a reliable estimate of $F$, particularly where $F$ has a strong influence on the model. This is particularly evident in the polarization magnitude permeability models (equations (7) and (8)) where uncertainty in $F$ will result in large prediction errors due to the large power law exponents of 5.35 and 5.38. While this limitation has yet to be overcome in low salinity groundwater settings, the models are expected to perform well in (1) brine formations where $F$ can be reliably measured in the field assuming that the brine conductivity is known and low phase angles can be measured under field conditions and (2) where surface conduction effects are very low. Future work will explore permeability estimation from geophysical length scales in such high salinity (i.e., brine) and low surface conduction formations.

### 6. Conclusions

Complex conductivity offers the possibility of field-scale permeability prediction through the measurement of geophysical length scales analogous to geometric length scales. Permeability for most samples from an extensive database of sandstones can be predicted to within one order of magnitude from previously proposed CC models based on either a polarization magnitude or a time constant. Exceptions are samples characterized by high $S_{por}$ and mineralogical variability, particularly the presence of dolomite. The dependence of the geophysical length scales on mineralogical variability not captured in the use of single values

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of a specific polarizability ($c_p$) or diffusion coefficient ($D_s$) may explain the limited predictive capability of these $k$ estimation models. Several limitations to the geophysical permeability models have been identified including the time constraint imposed by the acquisition of multifrequency data in models using $m_m$, $T_{pc}$, and $\tau_{mean}$, the existence and selection of $T_{pc}$ in the covered frequency range and the large uncertainty in the appropriate value of a representative diffusion coefficient. However, the largest obstacle to the field-scale implementation of the permeability equations (both geometrical and geophysical) is a reliable estimation of electrical formation factor ($F$). A recently proposed method to estimate $F$ based on a surface conductivity correction using CC measurements improves $F$ predictions but still leaves significant errors in that will limit field-scale $k$ estimation using such models.

Notation

- $CC$: Complex conductivity (mS/m)
- $c_p$: Specific polarizability (S)
- $d$: Mean absolute deviation on a logarithmic scale
- $D_s$: Diffusion coefficient (m$^2$/s)
- $F$: Electrical formation factor ($\sim$)
- $k$: Permeability (m$^2$)
- $k_{KT}$: Modified Katz and Thompson permeability model (m$^2$)
- $k_{pc}$: Geophysical polarization magnitude permeability model from Weller et al. (2015a) using $\sigma_{1Hz}^2$ (m$^2$)
- $k_{mn}$: Geophysical polarization magnitude permeability model from Weller et al. (2015a) using $m_m$ (m$^2$)
- $k_r$: Permeability model from Revil et al. (2012) using time constant $T_{pc}$ or $\tau_{mean}$ (m$^2$)
- $l_c$: Equivalent pore diameter where maximum incremental intrusion occurs using MICP (µm)
- $\Lambda$: Dynamically interconnected pore radius (µm)
- $\mu$: Mercury injection capillary pressure
- $m_m$: Normalized chargeability (mS/m)
- $r_{eff}$: Effective hydraulic radius (µm)
- $S_{por}$: Pore volume normalized surface area (µm$^{-1}$)
- $\tau$: Time constant (s)
- $\tau_{mean}$: Time constant (s) obtained from a Debye decomposition
- $T_{pc}$: Peak and/or corner characteristic time constant (s)
- $\sigma^r$: Imaginary conductivity (mS/m)
- $\sigma_{1Hz}^0$: Imaginary conductivity (mS/m) at a frequency of 1 Hz

References


