Dielectric Constant and Electrical Resistivity of Natural-State Cores

GEOLOGICAL SURVEY BULLETIN 1052-H

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By G. V. Keller and P. H. Licastro

Experimental and Theoretical Geophysics

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CONTENTS

Abstract .......................................................................................................................... 257
Introduction ...................................................................................................................... 257
Acknowledgments ............................................................................................................ 258
Experimental work .......................................................................................................... 258
  Preparation of core material .......................................................................................... 258
  Presentation of data for natural-state cores ................................................................. 258
  Measurements on dry cores ......................................................................................... 263
Evaluation of the data in terms of dispersion theories ..................................................... 265
Summary and conclusions ............................................................................................... 283
Literature cited .................................................................................................................. 284

ILLUSTRATIONS

Figure 68. Dielectric constants of natural-state Morrison cores ..................................... 259
69. Resistivity of natural-state Morrison cores ............................................................... 260
70. Loss angles measured on natural-state Morrison cores ............................................. 260
71. Phase angle versus water content at a frequency of 10 kc ........................................ 263
72. Dielectric constants of dried Morrison cores measured in vacuum ......................... 264
73. Equivalent circuit for electrode polarization ............................................................. 266
74. Dispersion curves showing effect of electrode-polarization capacity decaying as an inverse power of the frequency ............................................................... 267
75. Static dielectric constant versus water content ......................................................... 267
76. Distribution times as a function of the distribution parameter b ................................. 270
77. Wagner's equation for dispersion of the real dielectric constant in a substance with a normal distribution of relaxation times ......................................................... 271
78. Equation for dispersion of the imaginary dielectric constant in a substance with a normal distribution of relaxation times ......................................................... 271
79. Dielectric ratio as a function of distribution parameter and frequency ................. 272
80. Example of the graphical determination of Wagner's parameters b and z 0 ................. 273
81. Theoretical dielectric constant as a function of distribution parameter and frequency .......................................................... 274
82. Graphical determination of the maximum dielectric constant and the parameter k of Wagner's equation .......................................................... 275
83. Dielectric constants in core Mo-18 ....................................................................... 275
Figure 84. Equivalent circuit for a dielectric material having more than one relaxation time .................................................. 277
85. Dielectric constant for a laminated mixture of quartz and water .............................................. 278
86. Idealized pore structure .................................................. 278
87. Tracing made from a photomicrograph of a stained thin section, showing the pore structure of a typical sandstone in the Salt Wash member ............................................. 279
88. Number of immobile cations in a typical sandstone of the Salt Wash member .............................................. 281

Tables

Table 1. Characteristics of the dielectric dispersion of natural-state cores .............................................. 276
2. Evaluation of dielectric constant under certain hypothetical conditions .............................................. 282
EXPERIMENTAL AND THEORETICAL GEOPHYSICS

DIELECTRIC CONSTANT AND ELECTRICAL RESISTIVITY OF NATURAL-STATE CORPS

By G. V. Keller and P. H. Licastro

ABSTRACT

As part of a program to obtain basic data on the physical properties of the rock in and near uranium-mineralized zones, measurements of electrical resistivity and dielectric constant were made on parts of 27 cores from the Morrison formation in the Colorado Plateau uranium province. For frequencies between 50 cycles and 30 megacycles per second, resistivity ranged from $10^2$ to $10^{12}$ ohm-centimeters, and the dielectric constant ranged from 4.0 to more than $10^4$. The water content of the cores seemed to be the controlling factor: the high resistivities were associated with low water content and the high dielectric constants with high water content.

INTRODUCTION

The electrical properties of the Morrison formation in the Uravan mineral belt of the Colorado Plateau have been measured in place by electric logging and in the laboratory on cores with restored water saturation (Keller, 1959). The resistivity of different members of the Morrison formation in drill holes ranged from 8 to 2,000 ohm-meters. The resistivity of the sandstone members of the Morrison was found to be highest in areas that were considered most favorable for the occurrence of ore, probably because of the lower water saturation or lower salinity of the water in the favorable areas.

During 1952 a program of natural-state coring was begun by the U.S. Geological Survey in the Long Park area (Manger, Gates, and Cadigan, 1954) to obtain basic data on physical properties of rocks in and near uranium-mineralized zones. Drilling was done using air and oil-base drilling mud so that the water content of the cores could be preserved as nearly as possible in its original condition. These cores offered an opportunity to obtain truly representative values of the resistivity.

Parts of 27 cores of sandstones and mudstones of the Morrison formation, taken from 2 drill holes in the Long Park area, were used in
this study. Both dielectric constant and electrical resistivity were measured from frequencies of 50 cycles to 30 megacycles per second.

ACKNOWLEDGMENTS

The work herein reported was done in part under the program carried on by the Geological Survey on behalf of the Division of Raw Materials, U.S. Atomic Energy Commission. Some of the laboratory work was carried out at the Pennsylvania State University, where equipment and laboratory facilities were made available by the University.

EXPERIMENTAL WORK

PREPARATION OF CORE MATERIAL

For measurements of dielectric constant and electrical resistivity, a thin wafer was cut from each of the cores. The plane faces of these wafers were coated with conductive silver paint to form a pair of electrodes. The capacity and resistance of each piece of core were determined by using the core as one arm of a bridge circuit. A second arm contained a capacitor and resistor which were varied to match the capacity and resistance of the core. The bridge was operated at discrete frequencies from 50 cycles per second to 30 megacycles per second.

The core holder consisted of a brass plate placed in contact with the bottom of a core and a point electrode on a micrometer adjustment in contact with the top. Use of a conical-point electrode, which has practically no capacitance of its own, and wafer-shaped cores eliminated the necessity for Kelvin shielding of the cores and reduced considerably the complexity of the bridge adjustment.

PRESENTATION OF DATA FOR NATURAL-STATE CORES

The dielectric constants measured on the 27 natural-state cores are presented graphically in figure 68. The data are presented as dielectric ratios: that is, the capacity of a core divided by the capacity that an air condenser of the same dimensions would have. In order to obtain dielectric constants in cgs units from figure 68, it is necessary to multiply the ratios by the factor $8.85 \times 10^{-4}$ farads per centimeter, the dielectric constant of air.

The values of dielectric constant are strongly dispersed, ranging from 4 to more than $10^6$ at the low frequency end of the spectrum. At frequencies above 10 megacycles per second, the dielectric constants are all less than 80, as might be expected. In cores with very low water saturations, such as Nos. 10 and 12, there is almost no variation of dielectric constant with frequency.
Dielectric constant of natural-state cores

Figure 68.—Dielectric constants of natural-state Morrison cores.
Figure 69. Resistivity of natural-state Morrison cores.
The results of the resistivity measurements on these same cores are presented in figure 69 in cgs units, ohm-centimeters. In the dry cores (see, for example, Nos. 10 and 12) there is a linear inverse relation between resistivity and frequency over practically the entire range of frequencies studied. On the other hand, in the wet cores such as Nos. 21 and 23 there is no variation of resistivity with frequency.

The dispersion of resistivity and the anomalous behavior of dielectric constant raise a question as to which is the more important in controlling current flow. When a voltage is impressed on a material, the current that flows can be considered as consisting of two parts: one in phase with the applied voltage, resulting in ohmic conductivity; and another in quadrature with the applied voltage, termed the displacement current. The ratio of these two currents is the tangent of the phase angle between the applied voltage and the resultant current and is a measure of the relative importance of resistivity and dielectric constant in determining the response of a material in an applied electric field. The ratio of ohmic current to displacement current was determined for each of the 27 cores by

\[ \tan \delta = \frac{1}{\omega \varepsilon_0 \rho} \]

where
- \( \delta \) = the phase angle
- \( \varepsilon \) = the relative dielectric constant of a core
- \( \varepsilon_0 \) = the dielectric constant of air
- \( \omega \) = the frequency in radians per second
- \( \rho \) = the electrical resistivity

The phase angles are shown graphically in figure 70.

Conduction in the cores studied ranges from predominantly ohmic (\( \tan \delta > 1 \)) to predominantly displacement conduction (\( \tan \delta < 1 \)). The cores with the highest dielectric constant are those in which conduction is almost entirely ohmic; and in those with very low dielectric constant, conduction is almost entirely by displacement currents.

The transition from displacement conduction at low water content to ohmic conduction at high water content is relatively sharp. This is shown in figure 71 by the phase-angle data at 10 kilocycles per second (kc). At this frequency the transition occurs at a water content of 3 percent. At lower frequencies the transition takes place at lower water contents; but even at frequencies as low as 50 cycles per second, the quadrature component of conduction was important in some of the sandstone cores studied.
Figure 70.—Loss angles measured on natural-state Morrison cores.
MEASUREMENTS ON DRY CORES

The behavior of resistivity and dielectric constant described in the preceding sections seems to be intimately associated with the presence of water in the cores. As a further check on this, measurements of dielectric constant and resistivity were made on the same group of cores after their fluid contents had been removed. The oil in the cores taken with oil-base mud was removed in a Soxhlet extractor with pentane. Most of the water was removed by baking the cores at 160°C for 72 hours. The cores were evacuated while still hot, resaturated with dry nitrogen, and then stored in a dry nitrogen atmosphere. Measurements were later made with the cores in a vacuum. This procedure ensured the removal of all water except that closely associated with minerals in the cores, and it prevented re-entrance of moisture from the atmosphere while measurements were being made.

Some dispersion is evident in the dielectric constants of the dry cores (fig. 72). The dielectric constants at low frequencies are as much as 20 percent more than those at the high frequencies, but the tremendous variations observed in the cores in their original state are absent. It is apparent that the observed dispersion was entirely dependent on the presence of water.

Determinations of resistivity and phase angle were made on these dried cores; but as might be expected, the ohmic conductivity is practically negligible. Because the in-phase current is such a small
Figure 72. Dielectric constants of dried Morgan core measured in vacuum.
part of the total current in the dried cores, only a very low accuracy was achieved in these determinations. It was found, however, that all dried cores behaved as did cores number 10 and 12 in their natural state; that is, the resistivity of the dried cores was inversely proportional to the frequency.

**EVALUATION OF THE DATA IN TERMS OF DISPERSION THEORIES**

Examples of dispersion of electrical resistivity and dielectric constant are abundant in the literature; though few measurements of dielectric constants as high as those observed on some of the cores in the present work have been reported. The most pertinent of these earlier measurements are those of H. M. Evjen (1948), who estimated a dielectric constant of $10^8$ to explain transients observed in a sedimentary section. Dielectric constants as large as $10^8$ or $10^9$ have been reported by Fricke (1925, 1926) for certain biological-cell suspensions, by Fricke and Curtis (1935) for kaolin-gelatin suspensions, by Koops (1951) for artificially prepared ferrites, by Von Hippel (1954) for bottom-round steak, by Dunlap and Makower (1945) for dehydrated carrots, and by Smith-Rose (1934) and Tarkhov (1948) for rocks and other earth materials.

Several explanations for such anomalous behavior of dielectric materials have been advanced by the various authors. Smith-Rose (1934) and Tarkhov (1948) have suggested that the high dielectric constants observed on moist rocks are caused by electrode polarization. It is a well-known fact that a pair of metallic electrodes immersed in an ionic solution will act as a condenser of large capacity if current is caused to flow between the electrodes. This capacity is confined to a narrow zone near the electrode surface and is caused in part by a change in concentration near the electrodes during current flow, resulting in a change of the normal electrode potential in opposition to the direction of current flow. In addition, the increased density of cations near the cathode and of anions near the anode results in a concentration potential, also opposed to the current flow. For steady-state direct currents, the capacity of a smooth electrode surface is of the order of 6 to 12 microfarads per square centimeter. The capacity of metal electrodes with a large specific surface area may be much larger, and polarization capacities as large as 0.02 farads per square centimeter have been reported (Jaffe and Rider, 1952). With an alternating current, these reactions are effective only if the ions have time to concentrate at the electrodes during each half cycle. It can be shown theoretically that if the diffusion potential is the only factor contributing to the polarization capacity, then that capacity should decrease as the square root of
In a study of polarization capacity at intermediate and high frequencies, Jolliffe (1923) found that the inverse square-root relation held good for platinum electrodes. Other investigators have found that the capacity dies off less rapidly than the square root of frequency, and still others have found an inverse proportionality.

Jolliffe (1923) has suggested that electrode polarization can be represented by the equivalent circuit shown in figure 73. The effects of electrode polarization are represented by a resistor and by a variable capacitor in series and shunted with a second resistor. The capacitance is assumed to decrease exponentially as frequency increases, the factor $s$ representing the rate of decrease. Such an equivalent circuit would result in an apparent dispersion of the dielectric constants as shown by the curves presented in figure 74. For comparison, one of the observed dielectric-dispersion curves is shown. It is apparent that the observed dielectric constant decreases less rapidly with frequency than any of the cases where the dispersion might be the result of electrode polarization.

Further evidence that the high dielectric constants are not caused by electrode polarization is given by extrapolating dielectric constants to zero frequency. The apparent dielectric constant that would result from an electrode polarization capacity of 10 microfarads per square centimeter is shown as a function of water content by the solid curve in figure 75. The observed values of static dielectric constant are 2 to 5 orders of magnitude smaller than would be expected if the dispersion were caused by electrode polarization. This indicates that in the frequency range used in the present work, the effects of electrode polarization are relatively unimportant.

Actually, all solid and liquid dielectrics possess abnormal or anomalous properties at low frequencies beyond those that can be explained by electrode polarization (Hartshorn, 1926). In addition to the normal charging current and conduction current, such dielectrics exhibit an anomalous charging current when subjected to a steady-state
Figure 74.—Dispersion curves showing effect of electrode-polarization capacity decaying as an inverse power of the frequency.

Figure 75.—Static dielectric constant versus water content.
voltage. When a static voltage is applied to a material with anomalous behavior, the current flowing may be expressed as

\[ I_t = i_o + i_n(t) + i_a(t) \]  

where

\[ i_o = \text{the ohmic current flowing in the medium} \]
\[ i_n(t) = \text{the normal capacitative charging current} \]
\[ i_a(t) = \text{the anomalous charging current} \]

This anomalous charging current is comparatively large at the instant the voltage is applied but decreases rapidly with time. However in some materials an appreciable part of this current may persist for months. As the anomalous current is proportional to the applied voltage, its characteristics may be expressed by the equation

\[ i_a(t) = C_o E \beta F(t) \]

where

\[ C_o = \text{the geometric capacity of a system} \]
\[ E = \text{the applied voltage} \]
\[ \beta = \text{a proportionality parameter specifying the magnitude of the anomalous current in a particular material} \]
\[ F(t) = \text{the characteristic time function, describing the manner in which the anomalous current decreases with time} \]

The form of \( F(t) \) is a matter of some dispute. Many early investigators found that within the limits of their measurements the characteristic time function was a simple power function

\[ F(t) = \beta t^{-m}, m < 1 \]

It is obvious, though, that this expression cannot be physically correct, because if one considers zero time, the current is infinite, and if one considers the accumulated charge during an infinite charging period, this is also infinite.

In an alternating field the anomalous current will increase both the apparent capacity and the loss; the amount of this increase depends on the length of a cycle. According to Hartshorn (1926) Schweidler has shown that the fractional increase in capacity caused by the anomalous current is

\[ \Delta C/C = \beta \int_0^{\infty} F(t) \cos \omega t dt \]

where

\[ \omega = \text{the angular frequency of the alternating field} \]

An exponential form for \( F(t) \)

\[ F(t) = \beta e^{-\alpha t} \]

has been suggested on theoretical grounds by Maxwell and by Pellat (Hartshorn, 1926). Maxwell arrived at this expression by considering
the discharge current from a material composed of alternating layers of different electrical properties, and Pellat arrived at the same result by considering a viscous displacement of charge following the initial elastic deformation in an applied field.

Unfortunately, observed data do not ordinarily follow simple exponential time functions. Schweidler has extended Pellat's analysis by considering a characteristic time function consisting of a sum of exponential terms

$$F(t) = \sum \beta_n e^{-\alpha_n t}$$  

(7)

where there is an infinite variety of arbitrary coefficients $\beta_n$, $\alpha_n$.

This equation may be applied in a practical case by specifying a law of distribution for these arbitrary factors. Several such laws have been suggested (Fuoss and Kirkwood, 1941; Macdonald, 1952; and Davidson and Cole, 1950), but the law which is most applicable to our data is a log-normal distribution law suggested by Wagner (1913). Wagner's equation for the characteristic time function is

$$F(t) = \int_0^\infty \gamma(T)e^{-\frac{t}{T}}dT$$  

(8)

where

$$\gamma(T)dT = \text{the sum of the amplitudes of all terms in the range from } T \text{ to } dT,$$

with the distribution function replacing the individual amplitude parameters $\beta_n$ of equation 7

$T =$ relaxation time, replacing the reciprocals of the parameters $\alpha_n$ in equation 7

For a log-normal distribution, the distribution function is

$$\gamma(T)dT = \frac{\gamma b e^{-b z^2}}{\sqrt{\pi}} dz; z = \frac{T'}{T_0}$$  

(9)

where

$T_0 =$ the principle value of $T$, the time constants

$b =$ a parameter expressing how closely the values of the time constants are grouped about their principle value

Curves showing the distribution of time constants for various values of $b$ are presented in figure 76.

Wagner has shown that with such a distribution of time constants, the capacity and phase angle of a material are given by the following equations:

$$\frac{\Delta C}{C} = \frac{\gamma b e^{-b z^2}}{\sqrt{\pi}} \int_0^\infty e^{-u^2} \frac{\cosh (2b^2z_0 - 1) u}{\cosh u} du$$

$$(1 + \frac{\Delta C}{C}) \tan \delta = \frac{\gamma b e^{-b z^2}}{\sqrt{\pi}} \int_0^\infty e^{-u^2} \frac{\cosh b^2z_0 u}{\cosh u} du$$  

(10)
where

\[ z = \ln \frac{T}{T_0} \]
\[ z_0 = \ln \omega T_0 \]
\[ z + z_0 = u \]

Curves for this form of dielectric dispersion are presented in figure 77, and curves for the variation of phase angle with frequency are presented in figure 78.

An adaptation of a graphical method described by Yager (1936) was used to determine the parameters \( b, \gamma, \) and \( T_0 \) for the dispersion data obtained with the natural-state Morrison cores. In this technique a set of curves relating a factor \( \delta \) (defined as the ratio of two values of dielectric constant, one at 10 times the frequency of the other) a factor \( z = \ln \frac{T_0}{T} \) (\( T_0 \) is the principal relaxation time, and \( T \) is the relaxation time for the frequency at which \( \delta \) is defined) and the distribution parameter \( b \) are used. These curves are presented in figure 79.

In analyzing a particular group of data, the ratio of two dielectric constants a decade apart in frequency is formed; and by using the curves in figure 79, a curve relating \( z \) and \( b \) is constructed. This process is repeated with other pairs of dielectric constants taken at other frequencies. The final result is a series of curves of \( z \) versus \( b \) for a particular set of dielectric constants, as shown by the example

![Figure 76](image-url)
Figure 77.—Wagner's equation for dispersion of the real dielectric constant in a substance with a normal distribution of relaxation times.

Figure 78.—Equation for dispersion of the imaginary dielectric constant in a substance with a normal distribution of relaxation times.
in figure 80. The intersection of these trial curves determines the correct values of \( b \) and \( z_0 \) for the data. The principal relaxation frequency may be obtained from the definition

\[
z_0 = \ln \omega_r / \omega_0
\]

where

\( \omega_r \) = the frequency at which the first dielectric ratio was formed
\( \omega_0 \) = the principal relaxation frequency
The second stage in the graphical evaluation of the data is the determination of the low-frequency or static value of dielectric constant. For this purpose a group of curves showing the relation between $b$ (the distribution parameter), $z = \ln \omega / \omega_0$ (the frequency), and the dielectric constant (expressed as a fraction of the maximum value attained at low frequencies) is used. These curves are presented in figure 81.

As $b$ and $z$ are already known from the analysis described in the preceding paragraph, it is a simple matter to determine a theoretical value for the dielectric constant from figure 81 for each of the frequencies at which an experimental value was determined. When the experimental values are plotted as a function of these theoretical values, a straight line relation with a slope equal to the low-frequency dielectric constant is obtained provided the data satisfy Wagner's equations. An example is shown in figure 82.

This procedure was used to determine the dispersion parameters for each of the cores studied. The results are given in table 1.
An example of the agreement between the experimental data and Wagner's equations is given in figure 83. The observed values of resistivity are plotted in terms of an imaginary dielectric constant, defined as

$$\varepsilon'' = \frac{i\omega \rho}{\varepsilon_0}$$

(12)

where

$\omega = \text{the frequency in radians per second}$

$\rho = \text{the electrical resistivity in ohm-centimeters}$

$\varepsilon_0 = \text{the dielectric constant of a vacuum in farads per centimeter}$
Graphical determination of the maximum dielectric constant and the parameter $k$ of Wagner's equation.

Observed real dielectric constant

Observed imaginary dielectric constant

Theoretical real dielectric constant

Theoretical imaginary dielectric constant

Theoretical curves are Wagner's equations with the following numerical parameters:

$f_c = 1.35$ cycles per second

$b = 0.205$

$\varepsilon^\infty = 66,000$

$\varepsilon^\infty = 6.6$

$k = 10,000$

Figure 83.—Dielectric constants in core Mo-18.
The agreement between the real dielectric constant and Wagner's equation was excellent in all cases, but the agreement between imaginary dielectric constant (resistivity) and theoretical curves is not good.

### Table 1.—Characteristics of the dielectric dispersion of natural-state cores

<table>
<thead>
<tr>
<th>Core No.</th>
<th>Water content</th>
<th>Relaxation frequency</th>
<th>Distribution parameter $b$</th>
<th>Low-frequency dielectric constant</th>
<th>High-frequency dielectric constant</th>
<th>Dielectric increase-ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>0.088</td>
<td>61.3</td>
<td>0.24</td>
<td>$7.0 \times 10^4$</td>
<td>10.0</td>
<td>$7.0 \times 10^4$</td>
</tr>
<tr>
<td>2</td>
<td>0.059</td>
<td>1.75</td>
<td>0.26</td>
<td>$8.0 \times 10^3$</td>
<td>6.9</td>
<td>$1.1 \times 10^5$</td>
</tr>
<tr>
<td>3</td>
<td>0.012</td>
<td>254</td>
<td>0.44</td>
<td>$9.0 \times 10^5$</td>
<td>3.5</td>
<td>$1.6 \times 10^4$</td>
</tr>
<tr>
<td>4</td>
<td>0.014</td>
<td>18.1</td>
<td>0.42</td>
<td>$5.0 \times 10^4$</td>
<td>3.6</td>
<td>$4.3 \times 10^4$</td>
</tr>
<tr>
<td>5</td>
<td>0.031</td>
<td>1.57</td>
<td>0.35</td>
<td>$8.0 \times 10^4$</td>
<td>7.1</td>
<td>$3.0 \times 10^5$</td>
</tr>
<tr>
<td>6</td>
<td>0.0079</td>
<td>0.60</td>
<td>0.25</td>
<td>$6.0 \times 10^4$</td>
<td>7.9</td>
<td>$1.8 \times 10^4$</td>
</tr>
<tr>
<td>7</td>
<td>0.0049</td>
<td>0.87</td>
<td>0.20</td>
<td>$8.0 \times 10^4$</td>
<td>4.6</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>8</td>
<td>0.0013</td>
<td>0.68</td>
<td>0.20</td>
<td>$8.0 \times 10^4$</td>
<td>4.1</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>9</td>
<td>0.0009</td>
<td>1.02</td>
<td>0.20</td>
<td>$8.0 \times 10^4$</td>
<td>4.0</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>10</td>
<td>0.0189</td>
<td>0</td>
<td>0.13</td>
<td>$8.0 \times 10^4$</td>
<td>4.2</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>11</td>
<td>0.0106</td>
<td>1.59</td>
<td>0</td>
<td>$8.0 \times 10^4$</td>
<td>3.6</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>12</td>
<td>0.0910</td>
<td>0.67</td>
<td>0</td>
<td>$8.0 \times 10^4$</td>
<td>4.3</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>13</td>
<td>0.0076</td>
<td>0.01</td>
<td>0.13</td>
<td>$8.0 \times 10^4$</td>
<td>3.7</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>14</td>
<td>0.0013</td>
<td>1.74</td>
<td>0.175</td>
<td>$8.0 \times 10^4$</td>
<td>4.0</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>15</td>
<td>0.0540</td>
<td>1.45</td>
<td>0.44</td>
<td>$8.0 \times 10^4$</td>
<td>4.6</td>
<td>$6.0 \times 10^4$</td>
</tr>
<tr>
<td>16</td>
<td>0.0366</td>
<td>11.2</td>
<td>0.36</td>
<td>$8.0 \times 10^4$</td>
<td>9.0</td>
<td>$2.3 \times 10^5$</td>
</tr>
<tr>
<td>17</td>
<td>0.0513</td>
<td>55.3</td>
<td>0.35</td>
<td>$8.0 \times 10^4$</td>
<td>9.0</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>18</td>
<td>0.0180</td>
<td>1.36</td>
<td>0.25</td>
<td>$8.0 \times 10^4$</td>
<td>7.4</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>19</td>
<td>0.0110</td>
<td>0.45</td>
<td>0.20</td>
<td>$8.0 \times 10^4$</td>
<td>5.6</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>20</td>
<td>0.0730</td>
<td>1,510</td>
<td>0.60</td>
<td>$8.0 \times 10^4$</td>
<td>5.7</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>21</td>
<td>0.0445</td>
<td>670</td>
<td>0.50</td>
<td>$8.0 \times 10^4$</td>
<td>7.5</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>22</td>
<td>0.0505</td>
<td>2,470</td>
<td>0.76</td>
<td>$8.0 \times 10^4$</td>
<td>5.3</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>23</td>
<td>0.0363</td>
<td>4,580</td>
<td>0.72</td>
<td>$8.0 \times 10^4$</td>
<td>6.9</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>24</td>
<td>0.0465</td>
<td>2,277</td>
<td>0.25</td>
<td>$8.0 \times 10^4$</td>
<td>6.6</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>25</td>
<td>0.0151</td>
<td>12.2</td>
<td>0.20</td>
<td>$8.0 \times 10^4$</td>
<td>5.3</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>26</td>
<td>0.135</td>
<td>308</td>
<td>0.16</td>
<td>$8.0 \times 10^4$</td>
<td>6.1</td>
<td>$2.1 \times 10^5$</td>
</tr>
<tr>
<td>27</td>
<td>0.0306</td>
<td>2.23</td>
<td>0.27</td>
<td>$8.0 \times 10^4$</td>
<td>6.7</td>
<td>$2.1 \times 10^5$</td>
</tr>
</tbody>
</table>


Wagner's equations have provided a satisfactory description of dispersion of the dielectric constant and resistivity, but the problem of constructing a pragmatic model that can explain the physical processes involved remains.

Maxwell (1892) has shown that anomalous phenomena can result from a particular megascopic structure of the dielectric material. If a material is composed of alternating layers of two materials with different electrical properties, these layers may be represented by the equivalent circuit shown in figure 84. The increase in low frequency dielectric constant over the high frequency constant for such a system is given by

$$ K = \frac{(T_1 + T_2 - \tau) T - T_1 T_2}{T_1 T_2} $$

where

$$ \tau = R_1 C_1 $$

$$ r_2 = R_2 C_2 $$

$$ \tau = R_1 R_2 (C_1 + C_2)/(R_1 + R_2) $$

and $R_1$, $R_2$, $C_1$, $C_2$ are the resistances and capacities of the alternate layers.
The ratio of the increase has been plotted as a function of water content for a hypothetical rock consisting of layers of quartz and water (fig. 85). Although Maxwell's interfacial polarization can account for large dielectric constants, these constants are found only at very large water contents.

Von Hippel (1954) points out that there is a tacit assumption in Maxwell's derivation that the migration of charge carriers (ions) through the layers is unimpeded. McCardell (McCardell and Winsauer, 1953) has suggested that mudstones or clay-bearing sandstones actually act as ion sieves. If a rock is considered to be a semipermeable membrane with fixed negative charges scattered through it, then it is expected that positive ions may flow through the rock more readily than negative ions. The characteristics of such selectively permeable membranes have been discussed by Sollner (1945).

A typical pore structure in a clastic detrital rock might be considered as a relatively large, possibly equidimensional void, connected
Figure 85.—Dielectric constant for a laminated mixture of quartz and water.

Figure 86.—Idealized pore structure.
to other such voids by thin fibrous channels. The essential features of such a pore structure are shown by the idealized "porelet" in figure 86. An example of the actual pore structure in a sandstone of the Morrison formation is shown in figure 87.

Clay particles positioned in constrictions in connecting channels will effectively block migration of negative ions because of the small size of the channels and the force of repulsions from the negative charges usually present on the clay particles. In such a situation the positive ions are free to move through the rock under the influence of an external field, but anions can migrate only until they come in contact with one of these selectively permeable pore constrictions. As a result there is an accumulation of negative charge as the anions

Figure 87.—Tracing made from a photomicrograph of a stained thin section, showing the pore structure of a typical sandstone in the Salt Wash member.
migrate from one side to the other of the free area in a pore. The
average anion would migrate a distance equal to

$$d_\text{=} = \mu_\text{=} \int_{t=0}^{t=t_\text{=}} \frac{V_\text{(t)}}{D} \, dt$$  \hspace{1cm} (14)$$

where

- $\mu_\text{=} =$ the mobility of anions
- $V_\text{(t)} =$ the voltage drop across a given pore
- $D =$ the diameter of that pore
- $t =$ the length of time that voltage is applied

The voltage drop across a pore would be a function of time because
of the counter electromotive force set up by the concentration differ­
ences as anions collect at the barriers. The diffusion potential set
up by the anion concentrations will cause an accumulation of positive
ions and an associated polarization; but as long as the diffusion
potential is small compared to the applied voltage drop, the polariza­
tion due to cation concentration will be small compared to the polar­
ization of the anions.

If it is assumed that the cations are relatively unaffected by the
anion concentrations, the charge stored by the anions in one porelet
will be

$$q = \frac{d_\text{=} - I_\text{=}}{D}$$  \hspace{1cm} (15)$$

where

- $I_\text{=}$ = the total quantity of electricity passing through the pore by means of anion
  motion
- $\frac{d_\text{=} - I_\text{=}}{D} =$ the fraction of that motion which is not compensated by motion of cations
  in the opposite direction.

The fraction of the total current carried by the anions is a function
of the ionic mobilities and the relative concentrations of cations and
anions. The clay particles act as immobile anions, so the number of
mobile anions will be less than the number of cations. The part of
the total current carried by anions will be

$$I_\text{=} = \frac{\mu_\text{=} - a - a_\text{=} \alpha}{\mu_\text{=} + \mu_\text{=} - 2a - a_\text{=} \alpha} \, I$$  \hspace{1cm} (16)$$

where

- $\mu_\text{+}$ and $\mu_\text{=} =$ the mobilities of the cations and anions, respectively
- $\alpha =$ the concentration of cations
- $a_\text{=} =$ the equivalent concentration of the clay anions

Values of $a_\text{=}$ measured on a group of typical sandstones of the Morrison
formation, fully water saturated, are presented in figure 88.
The volume of the central part of a pore, as presented in the idealized form in figure 86, would be $\pi D^3/8$, and the volume of the small connecting channels can be called $k\pi D^3/8$, so that the number $n$ of pores in a unit volume of rock having a porosity, $\phi$, is

$$n = \frac{8\phi}{(1+k)\pi D^3}$$  \hspace{1cm} (17)

If the connecting channels of only a fraction, $N$, of these pore structures are blocked by selectively permeable membranes, the charge stored in a unit volume will be

$$Q = nNq$$  \hspace{1cm} (18)

Under the application of a unit voltage, the capacity of this unit volume will be equal to the charge, $Q$. Ordinarily, most of the resistance of a rock is contributed by the small pores because of their small cross section and great length. The larger, central pores will have a smaller voltage gradient, smaller by a factor, $m$, because of their smaller resistance per unit length.
Combining all of these considerations, it is possible to estimate the dielectric constant of a typical, water-saturated sandstone

$$\varepsilon = \frac{C}{C_0} = \frac{Q}{8.85 \times 10^{-14}}$$  \hspace{1cm} (19)

where

$C_0 =$ the capacity of a cubic centimeter of a vacuum, or $8.85 \times 10^{-14}$ farads

Using equations 14 through 18 to replace $Q$, we obtain

$$\varepsilon = \frac{9 \times 10^{13}}{(1 + k) \pi D} \left( N \right) \frac{\mu^2 I t (\alpha - \alpha_0)}{(\mu_+ + \mu_-) (2 \alpha - \alpha_0)} \int_{t=0}^{t=t} V_1(t) \, dt$$  \hspace{1cm} (20)

It is necessary to know how the diffusion potential caused by the concentration of anions depends on charging time and pore diameter to evaluate the integral precisely. However, in a qualitative approach, the integral may be evaluated as

$$\int_{t=0}^{t=t} V_1(t) \, dt = \frac{A(t, D) t E}{m}$$  \hspace{1cm} (21)

where

$E =$ the externally applied voltage gradient, taken as 1 volt per centimeter for the purposes of this discussion

$A =$ a constant depending on $t$ and $D$

Also, Archie’s law (Archie, 1947) may be used to express the current, $I$, in equation 20 in terms of porosity and water resistivity

$$I = \frac{E \phi^2 S^2}{\rho_w}$$  \hspace{1cm} (22)

where

$\rho_w =$ the resistivity of the water in the pores

$S =$ the fraction of the pores filled with water

Assuming that the pores are fully saturated, equation 20 becomes

$$\varepsilon = \frac{9 \times 10^{13} \phi^2 A N D \mu^2 (\alpha - \alpha_0)}{(1 + k) \pi D \rho_w m (\mu_+ + \mu_-) (2 \alpha - \alpha_0)}$$  \hspace{1cm} (23)

An evaluation of this expression for several hypothetical sets of conditions is presented in table 2.

**Table 2.—Evaluation of dielectric constant under certain hypothetical conditions**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\phi$</td>
<td>0.10</td>
<td>0.16</td>
<td>0.24</td>
</tr>
<tr>
<td>$t$</td>
<td>100</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>$D$</td>
<td>ppm</td>
<td>0.01</td>
<td>0.05</td>
</tr>
<tr>
<td>$N$</td>
<td>sec</td>
<td>1</td>
<td>20,000</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>ppm</td>
<td>50,000</td>
<td>50,000</td>
</tr>
<tr>
<td>$\mu_-$ (for Cl$^-$)</td>
<td>ppm</td>
<td>$6.76 \times 10^{-3}$</td>
<td>$6.76 \times 10^{-3}$</td>
</tr>
<tr>
<td>$\mu_+$ (for Na$^+$)</td>
<td>ppm</td>
<td>$45.6 \times 10^{-3}$</td>
<td>$45.6 \times 10^{-3}$</td>
</tr>
<tr>
<td>$\rho_w$</td>
<td>ohm-cm</td>
<td>20</td>
<td>34</td>
</tr>
<tr>
<td>$A$</td>
<td>cm</td>
<td>1</td>
<td>20</td>
</tr>
<tr>
<td>$I$</td>
<td>cm/sec</td>
<td>8.05 X 10$^4$</td>
<td>1.19 X 10$^4$</td>
</tr>
</tbody>
</table>

1 Values expressed as cm/sec.

volt/cm
The influence of any single factor on the dielectric constant cannot readily be evaluated because of the qualitative nature of this analysis. However, if values which are geologically probable are assumed for the various parameters, it is obvious that the existence of a membrane effect would lead to high values of static dielectric constant.

One may infer from such considerations that siltstones will probably have larger dielectric constants than any other rock types. In shales all the negative charges may be immobile, so that no polarization can take place; and in clean sandstones, there may be no anion traps to cause polarization. Siltstones will provide a large number of anion traps and at the same time have a sufficient number of mobile anions to make use of these traps.

Other mechanisms for the origin of large polarizations have been suggested (Dakhnov and others, 1952). In a metal-bearing rock, electrode polarization may take place at every interface between metal and electrolytic solution. The sandstones of the Morrison formation contain only small percentages of metallic minerals, so this mechanism is not of interest. A second phenomenon suggested by Dakhnov is the storage of charge by electroosmosis. When a voltage is applied across a rock, the water in the pores may be carried along with the electric current. Following the cessation of current the displaced water will return to its original position, giving rise to an electrokinetic potential. As the water in sandstones of the Morrison formation is highly saline, it is not likely that an appreciable electrokinetic potential could be generated or that an appreciable charge could be stored.

**SUMMARY AND CONCLUSIONS**

The experiments described in this report suggest that the electrical characteristics and probably the nature of electrical conduction in earth materials is considerably more complicated than is commonly assumed in geophysical prospecting. Measurements on water-bearing sandstones and siltstones from the Morrison formation showed a strong dispersion of both resistivity and dielectric constant with frequency. The variation in resistivity is most pronounced in rocks with low water contents; and large dielectric constants are associated with rocks having the higher water contents, such as siltstones and mudstones. If a rock is considered to contain selectively permeable membranes, as has been suggested to explain natural potentials observed in electric logging, it is possible to explain extremely high dielectric constants.

These measurements were made on parts of only 27 cores from 2 neighboring drill holes in a small area of the Colorado Plateau, but there is no reason to believe that these specimens are unique. The high dielectric constants observed with the water-bearing mudstones
(as great as $10^6$) might also be observed with any equally wet mudstone or shale. The dispersion of resistivity observed on the sandstones may not be so common, as it appears only with water contents of less than 3 percent of the rock volume; but there are examples of sandstones in desert areas and in oil-wet oil reservoirs where such low saturations are the rule rather than the exception. For this reason the consideration of these phenomena in electrical prospecting should be of general interest to geophysicists.

LITERATURE CITED


