Original and pyrometamorphically altered Bentheimer sandstone; petrophysical properties, surface and dielectric behavior

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ABSTRACT
Bentheimer sandstone is a quartz-rich permeable hard sedimentary rock used for core flooding experiments. When fired to stabilize clays (subjected to high temperatures), pyrometamorphical phase changes induce texture and pore framework alteration. As a consequence the new dielectric response may influence wettability. The literature regarding pyrometamorphical behavior during and after thermal treatment is ambiguous, so we evaluate desirable effects (fixation of clay minerals) and undesirable effects (dielectric surface changes) in the matrix. Porosity, permeability, surface charge, specific surface area and dielectric respond were measured before and after firing of samples up to ~1000 °C under oxidizing and non-oxidizing conditions. The matrix properties were determined using X-ray diffraction and X-ray fluorescence, scanning electron microscope imaging, and thermomechanical- and thermogravimetric analysis with differential scanning calorimetry.

Firing causes dehydration, dehydroxylation and irreversible transformation of original clays, organic matter, and carbonates to glass, oxides and feldspars. During heating quartz transfers from α- to β-quartz and back during cooling. This changes the grain volumes and consequently reduces the matrix integrity. The sandstone has a slight porosity and permeability increase (~5%). Further, a shift in the point of zero charge toward a higher pH may result in wettability alteration from strongly water-wet to oil-wet. Additionally, a decrease in the permittivity value and marginal dispersion of the dielectric constant (~5%) between the high and the low frequencies was observed. Due to firing and related dispersion of the iron oxides within the matrix framework, Bentheimer sandstone becomes a weaker insulator.

1. Introduction

Bentheimer sandstone (BS) is widely used as a model rock for studying fluid flow in porous media and preparing 3D pore network models for theoretical studies (De Boever et al., 2013; Klein and Reuschlé, 2003; Ruedrich and Siegesmund, 2007). It is an easy to obtain, inexpensive sample material that has a matrix composed of well-sorted, mostly rounded to sub-rounded grains with a regular distribution of pore bodies and pore throats diameter (Peksa et al., 2015). The presence of accessory clay, organic matter and oxides or hydroxides is often ignored. Nevertheless, they influence petrophysical behavior such as permeability and electrical conductivity behavior.

1.1. Motivations for firing

Sandstone samples are heated to temperatures exceeding 500 °C as a common step in core preparations to limit the risk of permeability reduction during displacement experiments (Potts and Kuehne, 1988; Sydansk). The objective is to remove the chemically bounded water and to transform and stabilize clays. A strong influence of clays in contact with water on sandstone porosity/permeability, pore size geometry and electrical behavior was reported in the literature (Al-Mjeni et al.; Civan and Knapp, 1987; Gabriel and Inamdar, 1983; Gray and Rex, 1966; Huntley, 1986; Klimentos and McCann, 1990; Neasham, 1977; Sharma and Yortsos, 1987; Wildenschild et al., 1999). The high cation exchange capacity of clays, the large surface area and strong interaction with interstitial fluids result in swelling and/or migration. In addition, permeability reduction can be expected because of pore clogging downstream, which causes serious injectivity problems (Gray and Rex, 1966; Jones; Mohan and Fogler, 1997; Mohan et al., 1993; Norrish, 1954). Clay migration was even found to clog high permeable sandstones (~0.5Darcy) (Gray and Rex, 1966). Note that besides clay fines, other components such as organic matter, very fine quartz and feldspars particles, are considered as mobile fines.

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Nomenclature

| Symbol | Definition |
|--------|------------|
| $a$    | Average coefficient of expansion, [-] |
| $A$    | Area of the electrode, m$^2$ |
| $\Delta \epsilon_r$ | Dielectric dispersion magnitude |
| $\Delta n$ | Dissipation factor, [dimensionless] |
| $C_0$  | Capacitance of space in a parallel-plate capacitor, pF |
| $C$    | Capacitance of dielectric in a parallel-plate capacitor, pF |
| DAR    | Dielectric Absorption Ratio, [-] |
| $\epsilon'$ | Complex electric permittivity in the frequency domain, F/m |
| $\epsilon_0$ | Permittivity of space, $\epsilon_0 = 8.85*10^{-12}$ F/m |
| $\epsilon''$ | Imaginary part of the electric permittivity, F/m |
| $\epsilon_{r'}$ | Real value of the relative permittivity, [-] |
| $\epsilon_{r''}$ | Imaginary value of the relative permittivity, [-] |
| $E$    | Electric field vector, V·m$^{-1}$ |
| $f$    | Frequency, Hz |
| $F$    | Faraday constant, C/mol |
| $G$    | Conduction, S |
| $G_{AC}$ | AC conduction, S |
| $G_{DC}$ | DC conduction, S |
| $g(t)$ | Dielectric response function of Bentheimer sandstone sample |
| $I$    | Total current density vector, A/m$^2$ |
| $I_c$  | Capacitive part of the current density vector, A/m$^2$ |
| $I_{c'}$ | Capacitive part of the electric current, A |
| $I_{L}$ | Leakage part of the current density vector, A/m$^2$ |
| $I_{L'}$ | Leakage part of the electric current, A |
| $I_p$  | Polarization current through the sample, A |
| $I_r$  | Insulation resistance, $\Omega$ |
| $L$    | Sample length, mm |
| $L_0$  | Initial sample length, mm |
| $\delta L/\delta T$ | Slope of the expansion curve at the temperature, mm$^2$/°C |
| $M$    | Mass, g |
| $P$    | Polarization, C/m$^2$ |
| PI     | Polarization index, [dimensionless] |
| $\varphi$ | Effective porosity, [dimensionless] |
| $\rho$ | Bulk density, kg/m$^3$ |
| $R$    | Lossy part of the dielectric, $\Omega$ |
| $\tau_r$ | Relaxation time, s |
| $S_s$  | Specific surface, m$^2$/g |
| $\sigma_0$ | Charge, C/m$^2$ |
| $\sigma$ | Electrical conductivity, S/m |
| $T$    | Temperature, °C |
| $t_p$  | Polarization duration, min |
| $\tan\delta$ | Dissipation factor, [dimensionless] |
| $\tan\delta_{tot}$ | Polarization losses on the dielectric, [dimensionless] |
| $\tan\delta_{cond}$ | Conduction losses on the dielectric, [dimensionless] |
| $U$    | Voltage, V |
| $V_m$  | Matrix volume, m$^3$ |
| $\omega$ | Angular frequency, Hz |
| $\omega$ | Angular frequency, Hz |
| $\Delta$ | Alternating current |
| $\alpha$ | Angle of refraction, [-] |
| $\beta$ | Angle of reflection, [-] |
| $\gamma$ | Angle of incidence, [-] |
| $\delta$ | Angle of extinction, [-] |
| $\theta$ | Angle of diffraction, [-] |
| $\phi$ | Angle of polarization, [-] |
| $\psi$ | Angle of rotation, [-] |
| $\chi$ | Angle of absorption, [-] |
| $\rho$ | Density, kg/m$^3$ |
| $\mu$ | Permeability, N·s/m$^2$ |
| $\kappa$ | Conductivity, S/m |
| $\lambda$ | Thermal conductivity, W/m·°C |
| $\nu$ | Poisson’s ratio, [-] |
| $\sigma$ | Surface tension, N/m |
| $\theta$ | Contact angle, degree |
| $\gamma$ | Surface energy, J/m$^2$ |
| $\Delta H$ | Enthalpy change, J/mol |
| $\Delta S$ | Entropy change, J/K·mol |
| $\Delta G$ | Free energy change, J/mol |
| $\Delta F$ | Gibbs free energy change, J/mol |
| $\Delta V$ | Volume change, m$^3$/mol |
| $\Delta n$ | Refractive index change, [-] |
| $\Delta T$ | Temperature change, °C |
| $\Delta h$ | Height change, mm |
| $\Delta W$ | Work done, J |

1.2. Firing controls on wettability

Firing of the sandstone caused disintegration of clay minerals, a partial transformation of feldspar and quartz into glass and iron oxides. Moreover, according to Barclay and Worden (2009), if the specific surface of the iron oxides increases, a positive correlation of oil-wetting iron mineral content in the sandstones is observed. González and Moreira (1991) reported iron oxides to be strongly oil-wet due to substantial adsorption of asphaltene. They observed a significantly higher maximum of asphaltene adsorption for hematite than for clay minerals (up to 1.5 times) (González and Moreira, 1991). According to Wang and Guidry (1994), the mechanisms of the strong oil-wetness of iron oxides can be explained by (1) ferric ions, located on the mineral surface. They represent possible adsorption sites for carboxylic acid anions which can change the wettability of the mineral surface; (2) ferric ions, mainly on the surfaces. They can cause oxidation of the petroleum components using catalysts and produce polar wettability-altering compounds; (3) ferric ions in solution. They may behave as bridges between petroleum and a silica surface. In addition, it must be noted that the wettability state of the reservoirs containing ferric ions (Fe$^{3+}$) is indirectly controlled by the chemistry i.e. redox state, of the reservoir fluids (Barclay and Worden, 2009).

1.3. Firing characterization

Following previous studies (Ma and Morrow, Shaw et al.; Wu and Firoozaabadi, 2010) that high temperatures cause changes in mineralogical composition of sandstones and in physical and electrical properties, we decided to fire the Bentheimer samples to temperatures where all silica and carbonate minerals but quartz are dehydrated, dehydroxylized, decarbonized and disintegrated. For Bentheimer sandstone, the information on pyrometamorphical and electrical properties has been found available in the literature is rather limited (Loahardjo et al., 2012; Maloney et al., 1990; Olafuyi et al., 2010). In particular we are not aware of published works on the role of high-temperature minerals, resistant to chemical weathering, within its framework after firing and their effect on wettability. To fill the gap in the understanding of these systems, we decided to perform a series of thermal treatment at temperatures ≥600 °C and petrophysical, petrological and dielectric experiments.
2.2.1. Characterization of unfired and fired samples and thermal analysis

2.2.1.1. X-ray diffraction (XRD)/X-ray fluorescence (XRF) and texture composition

Samples characterized for XRD and XRF analysis were pulverized to a powder with a particle size of less than 2 µm and analyzed for their mineral composition and element-oxide distribution. The XRD was recorded in a Bragg-Brentano geometry, equipped with a Huber incident beam monochromator and a Bruker PXI diffractometer with a Huber incident beam monochromator and Bruker PXI diffractometer equipped with a Huber incident beam monochromator and a Bruker PXI diffractometer. The XRF measurements were conducted with a Panalytical Axios Max WD-XRF spectrometer measuring the element oxides and elements (>0.5 wt%) with an accuracy of 2%. Details on XRD and XRF and corresponding setups are provided in literature (Morse, 2013; Moore and Reynolds, 1989).

2.2.2. Scanning electron microscope (SEM)

To identify changes in the matrix structure, selected samples were examined using a scanning electron microscope (JEOL 8800 M JXA SuperprobeTM). The methodology of SEM and further image analysis can be found elsewhere (Krisley et al., 2005).

2.2.3. Porosity and permeability measurements

Porosity and permeability were determined on cylindrical samples having a diameter of 30 mm and length of 36 mm. Dry porosity tests were conducted with a Gas Expansion UltraPycnometer 1000 (Quantachrome InstrumentsTM) using Boyle’s gas expansion law (Selley and Sonnenberg, 2014). The results are the matrix volume ($V_m$), the bulk density ($\rho$) and the porosity ($\phi$). The permeability of dry cores was measured with $\text{N}_2$ gas by using a Ruska gas permeameter (Ruska InstrumentsTM) (Tiab and Donaldson, 2015).

2.2.4. Thermogravimetry (TGA) and differential scanning calorimetry analysis (DSC)

For thermal analysis, the samples were crushed to powders with particles smaller than 50 µm. The thermos-analyzer Netzsch STA F3 JupiterTM with QMS403C AéolosTM gas analysis system was simultaneously used to conduct thermogravimetric analysis (mass change ± 1 µg) and differential scanning calorimetry (heat difference ± 0.1 µW). The instrument measures the mass changes and provides endo-/exotheermic results as a function of temperature against a reference measurement in an empty crucible (Gabbott, 2008). The measurements were performed in both argon and air conditions at atmospheric pressure. The standard firing procedure was done in the air atmosphere, although to acquire more information about thermal mineral behavior in a neutral or reducing environment, argon was introduced. The gas velocities were 20 ml/min, with a dynamic temperature program at rate of 10 °C/min. In order to detect possible drying of the sample, the measurement started at 30 °C and went to a maximum temperature of ~1000 °C (± 1.5 °C).

2.2.5. Thermal expansion

To measure the changes in the physical properties of sandstone such as coefficient of thermal expansion ($\alpha$) and linear thermal expansion ($D/L$), mechanical analysis techniques in thermal analysis were used (Haines, 2002). The cuboid sample of 5.34 mm ± 0.01 mm in length and of a cross-sectional area 4 mm x 4 mm was placed in the PerkinElmerTM thermomechanical analyzer to measure small changes in sample dimensions. The sample was evenly heated over a programmed temperature range of 10 °C/min up to maximum 600 °C and kept constant for 2 h. Thereafter, it was cooled down at the same rate. The thermomechanical analyzer converts movements of the probe into an electrical signal and generates the thermal response curves from which the coefficient of thermal expansion was derived. The average coefficient of thermal expansion (CTE) is represented by $\alpha$, that is a function of temperature:

$$\alpha(T) = \frac{1}{L_0} \frac{\partial L}{\partial T}$$

where $\frac{\partial L}{\partial T}$ is the change in the sample length ($L$) over a temperature range expressed as a slope of the expansion curve at temperature $T$; $L_0$ is the initial sample length.

2.2.6. Surface charge and specific surface area

Potentiometric titrations were performed in a nitrogen atmosphere to the change in surface charge of Bentheimer sandstone due to the firing process. Before starting the experiments a sample of Bentheimer (0.5, 5 and 10 g) was dispersed in the electrolyte (100 ml $\text{H}_2\text{O}$ with 9 ml of 0.1 M NaOH) and then stirred vigorously with a Teflon magnetic stirrer for 24 h. The potentiometric titrations were performed by adding 0.1 M HCl droplet by droplet, while continuously stirring the
solution. The pH data were recorded every 5 min. The pH-dependent surface charge (σ0) was calculated as Janusz (1999):

\[ \sigma_0(pH) = \frac{F \cdot (\Delta n_{tot, dry} - \Delta n_{loc})}{M_s} = \frac{f_{ads}(pH)}{S_s}, \]

where \( F \) is the Faraday constant (\( F=96500 \text{ C/mol} \)); \( M \) is sample mass; \( S \) surface area and \( \Delta n_0 \) the pH-dependent amount of acid/base consumed. The dissolution effect (\( \Delta n_{tot, dry} \)) was obtained by comparison of the balance of protons and hydroxyl ions in the potentiometric titration of Bentheimer samples with blank potentiometric titration results:

\[ \Delta n_{tot, dry}(pH) = \Delta n_{loc, base}(pH) - \Delta n_{loc, base/blank}(pH). \]

For correct interpretation of the electrical behavior of the powdered fresh and fired Bentheimer sandstone, the specific surface area was measured by adsorption of liquid nitrogen, at low temperatures (−196.5 °C). The detailed methodology is provided elsewhere (Peksa et al., 2015). The specific surface data was calculated using the BET sorption theory (Brunauer et al., 1938). The obtained specific surface area represents the “external” surface area, since the penetration through the interlayer of smaller clay particles surfaces by weakly adsorbed nitrogen is not entirely possible.

2.3. Dielectric spectroscopy

2.3.1. Principle and apparatus

The main objective of the dielectric spectroscopy studies was to study differences in between unfired and fired samples through the frequency dependence of the permittivity and dissipation factor. The secondary objective is to conduct the dielectric adsorption test under DC conditions to recognize a change in the conductive behavior. For the tests, flat circular discs with a diameter of 140 mm and thickness, \( h \) of 8–10 mm were placed between two electrodes to arrange a parallel-plate capacitor of plate area, \( A \sim 15.9 \times 10^6 \text{ mm}^2 \). The contact surface of the samples and the electrodes was sufficiently smooth for the current transfer to occur.

The apparatus used for the measurement of the dielectric properties of the samples consisted was a three-terminal parallel-plate capacitor connected to a Megger™ capacity meter (Fig. 2) consisting of: (1) a bottom electrode directly connected to a high voltage unit, and (2) a top electrode connected to a low voltage terminal. The internal part of the upper electrode is used for measurements while the external part acts as a guard and eliminates the influence of surface leakage currents from the measurement results (Fig. 2a). The electrode terminal is covered by a tight glass shade to prevent the entrance of moisture from the atmosphere.

2.3.2. Sample preparation

Prior to the experiments, several techniques are employed to remove the water including heating in the oven at 105 °C for 48 h, and further saturation of the cores placed in the three-terminal parallel-plate capacitor with dry air. The drying procedure in the capacitor was implemented three times for each sample. Samples were subsequently: saturated with 90 °C dry air, evacuated, re-saturated with dry air, and then stored in a dry air atmosphere for 24 h. Measurements were later made with the cores in vacuum. The above procedure ensured the removal of free water from the samples. To reduce and quantify the effect of the electrode polarization (ionic charge accumulations at the electrodes) (Ishai et al., 2013) the following methods have been used when measuring dielectric properties: (1) varying the thickness of the samples to quantify the polarization effect; (2) using aluminum films as spacers to reduce the effect as much as possible.

2.3.3. Dielectric permittivity test

The test cell was connected to a capacity meter that measures the capacity and the dissipation factor of a sample as a function of frequency at a desired voltage of \( U=0.5 \text{ kV} \). A non-ideal dielectric like BS supports a varying charge, absorbs and dissipates the electric energy in the form of heat. When a potential difference is applied to a capacitor, energy is dissipation dissipated due to two main factors: (1) a flow of free charges through the material, called conduction loss (DC term), and (2) a displacement of electric charges that gives rise to the creation of dipoles in the sample (AC term). The second mechanism, called polarization \( P \), arises as a result of an electronic and an ionic polarization, a dipole orientation and an interfacial polarization (Howell and Licastro, 1961). The polarization of heterogeneous materials like Bentheimer sandstone depends on the individual properties of minerals, their proportion and distribution across the sample volume. This mechanism is related to the space charge accumulation along the interface in the presence of the electrical field. For this reason, it can be expected that the relative permittivity of a mixture of minerals may exceed that of a single mineral. Different types of polarization have varying time response abilities to an applied field
frequency. Therefore, the real part of electric permittivity is frequency dependent (Fig. 3). The frequency dependence of the polarization process on the relative permittivity value is presented elsewhere in greater detail (Dakin, 2006). Measurements conducted that are described in this paper cover the interval of the frequency domain from 0.1 to 15 kHz. Application of an electric field \( \mathbf{E} \) gives rise to the electric displacement field \( \mathbf{D} \) flowing the relation (Laj and Channell, 2007):

\[
\mathbf{D} = \varepsilon^* \mathbf{E}.
\]

(4)

The interactions of a material with an electric field \( \mathbf{E} \) are described by the complex electric permittivity \( \varepsilon^* \):

\[
\varepsilon^* = \varepsilon' - i\varepsilon''.
\]

(5)

where the expressions \( \varepsilon' \) and \( \varepsilon'' \) define the real and non-negative component of the permittivity and the energy dissipation due to polarization, respectively. The real part of the complex permittivity can be calculated at the various frequencies by using the measured capacitance values of a sample in a parallel-plate capacitor between the high and low voltage electrode (\( C_s \)) at the strong accumulation region from the relation (Bartnikas, 1987)

\[
C_s = \varepsilon_r C_0
\]

(6)

where \( C_0 \) is the capacitance without the sample, \( \varepsilon_r \) denotes the real part of the relative permittivity, represented by the ratio \( \varepsilon_r / \varepsilon_0 \), where \( \varepsilon_0 \) is the permittivity in vacuum.

Due to the existence of the loss current, the BS in the parallel-plate capacitor can be defined as a “leaky” dielectric and can be represented by the equivalent circuit (Fig. 4a). The losses in the sample can be represented by a conductance \( G \). If an AC voltage source \( U \) is applied on the capacitor, the total current \( \mathbf{I} \) propagating in the sample consists of two components:

\[
\mathbf{I} = \mathbf{I}_c + \mathbf{I}_L
\]

(7)

A charging current \( \mathbf{I}_c \) leads the in-phase component current by 90°(Fig. 4b) and can be expressed by

\[
\mathbf{I}_c = \omega \mathbf{C}_0 U,
\]

(8)

where \( \omega \) denotes the angular frequency (\( \omega = 2\pi f \)). A loss current \( \mathbf{I}_L \) that is in phase with the applied voltage includes the AC conduction from the inertial resistance \( G(\omega)C_r \) and the DC conduction \( G_{DC} \):

\[
\mathbf{I}_L = (G(\omega)C_r + G_{DC}) U
\]

(9)

with \( G(\omega)C_r = \omega \mathbf{C}_r \).

(10)

The DC conductance \( G_{DC} \) of the samples can be represented in terms of the electrical conductivity \( \sigma \) as \( G_{DC} = \sigma A/d \). Thus, implementing (8), (9) and (10) into (7), the total current for a BS sample can be expressed as

\[
\mathbf{I} = (i\omega c_r C_0 + \omega c_r C_0 + G_{DC}) U = (i\omega c_0 C_r + G_{DC}) U
\]

(11)

where \( c_r \) is the imaginary value of the relative permittivity, represented by \( c_r / c_0 \) ratio. From the above expressions and from the parallel circuit presented in Figs. 4a and b, the total dissipation factor value of the sample can be derived as Arora and Mosch (2011), Nabighian (1987)

\[
\tan \delta = \frac{I_L}{I_c} = \frac{G_{DC} + \omega c_r C_0}{\omega c_r C_0} = \frac{G_{DC}}{\omega c_0 c_r C_r} + \frac{\varepsilon_r}{\varepsilon_0}
\]

(12)

The total dissipation factor of a dielectric is characterized by the sum of: a) the polarization losses, \( \tan \delta_{pol} \), and b) the losses caused by conduction, \( \tan \delta_{cond} \) when the resistance of the dielectric is sufficiently small. Hence, the dissipation factors related to the polarization and conduction are equal to:

\[
\tan \delta_{pol} = \frac{\varepsilon_r}{\varepsilon_0},
\]

(13)

\[
\tan \delta_{cond} = \frac{\sigma}{\omega c_0 c_r}
\]

(14)

Assuming that the DC contribution of loss current is much smaller than the AC contributions (\( G_{DC} \ll \omega c_r C_0 \)), the total dissipation factor simplifies to

\[
\tan \delta = \frac{\varepsilon_r}{\varepsilon_0}
\]

(15)
2.3.4. Polarization current measurements and dielectric absorption test

The main concept of this test is the increase in insulation resistance due to the polarization of the atoms in sandstone over time. The DC voltage ($U = 0.2$ kV) is applied to samples placed in the same three-terminal parallel-plate capacitor during a polarization duration, $t > 10$ minutes. Thus, due to the application of a DC voltage $U$, a pulse-like polarization current flows through the sample. During the polarization duration, the current decreases to a certain value corresponding to the conduction current. This is assuming that the sample in the parallel-plate capacitor is totally discharged prior to the measurement and that a step voltage is applied with the following characteristics (Jonscher, 1984)

\[
U(t) = \begin{cases} 
0 & t < 0 \\
U_0 & 0 \leq t \leq t_p \\
0 & t > t_p 
\end{cases}
\]  

The above relation gives zero current for times before $t=0$, and polarization currents for times $0 \leq t \leq t_p$. The polarization current through the sample can be expressed as Jonscher (1984)

\[
I_p = U_0C_0 \left[ \frac{\sigma}{\epsilon_0} + \epsilon_0\gamma(t) + g(t) \right].
\]

It contains three parts: (1) process independent of any polarization being in relation to the conductivity of the sample; (2) the gamma function, arising from the suddenly applied step voltage, at $t=0$ and
\[ I_p = \frac{U_0 C}{\epsilon_0 + g(t)} \]

**NOTE:** The dimension of \( g(t) \) is \( \text{l/s} \) and its magnitude is tied to \( C_p \). Moreover, the simultaneous measurements of the insulation resistance \( (I_p) \) at different time intervals allow the calculation of: a) dielectric absorption ratio (DAR) that is the ratio of the insulation resistance at 60 s to the insulation resistance at 30 s, and b) the polarization index \( (P) \) that is the ratio of the insulation resistance value measured at 600 s to the insulation resistance value measured at 60 s. All reported polarization current measurements were performed at ambient temperature.

### 3. Results and discussion

#### 3.1. XRD/XRF analysis and SEM interpretation: general results

The pyrometamorphic alterations up to 970 °C have been evaluated by XRD/XRF-analysis and SEM. The primary constituents of the Bentheimer sandstone are quartz (91.7 wt%), feldspars (4.9 wt%), clay minerals (2.7 wt%), carbonates (0.4 wt%), pyrite and iron hydroxides (0.2 wt%). Note that the wt% are average values from the analysis of our previous study (Peksa et al., 2015). Additionally, SEM analyses show the presence of kaolinite, illite, and mixed-layer clays in Bentheimer sandstone.

Fig. 5 mainly shows the reduction of intensity corresponding to kaolinite in the XRD patterns between unfired and fired Bentheimer sandstone. In samples fired to 970 °C the structure became amorphous, and might only represent traces of meta-kaolinite and mullite (Figs. 6c–d) (Friolo et al., 2005; Ismailov, 1981). Quartz and feldspars were transformed to a lesser extent in comparison to other components, as most of the sandstone components have a lower melting point than quartz. They melt rst creating vitrified textures and consequently, the quartz grains dissolve into the glassy matrix (Figs. 6c–d). In the case of the feldspar grains, a disappearance of clay minerals is observed. It results in irregular grain shapes, and in small amounts of glass at the grain boundaries (Fig. 6d). Therefore, it indicates an initial integration of the feldspars into melts. Moreover, distinct interfaces between the grains were recognized in the SEM-photos (Fig. 6b). Before firing, the boundary between the detrital quartz grains was merged by overgrowths and/or pressure solutions.

Furthermore, hematite was identified as a minor component in the unfired sample and as well in an increased amount as a secondary phase. Decomposition of most of iron (hydro) oxides/carbonates (goethite, siderite) and iron rich cement resulted in conversion to mainly hematite and probably wüstite, giving the rock a distinctive reddish color (supplement Fig. S1). The destruction of the cementing material by firing made Bentheimer more friable. An interesting observation was SEM identification of rutile as an accessory component, occurring both as a precursor and as a polymorph phase after anatase (Figs. 6e–f).

### 3.2. Thermal analysis and phase transformations

TGA and DSC analyses were carried out in argon and air environment, and the results are illustrated in Fig. 7. The dehydration of sandstone components is observed in the DSC traces with an initial endothermic peak observed between 40 °C and 170 °C, and between 20 °C and 90 °C in the air and argon environment, respectively. The process is related to free water being removed from the minerals. In the range 200–350 °C, small endothermic peaks demonstrate the dehydration of iron-oxides and -hydroxides and the crystallization of iron oxide, mainly hematite (Prasad and Sitakara Rao, 1984; Sharma et al., 1994). The slight mass losses observed in the thermal data in the region 470–650 °C in each of the TG curves indicate dehydration processes and thermal decomposition resulting into oxidation in air and reduction with oxygen release in an argon environment. In region 470–650 °C (in each of the DSC curves) peaks centered at 580 °C corresponding to the dehydration of kaolinite are observed (Grim, 1968). For argon atmosphere, an additional endothermic reaction took place above about 750 °C, while endothermic disintegration and exothermic oxidation occurred in the air. In the air environment, the exothermic peak can be associated with mullite formation (Carroll, 1970; Friolo et al., 2005; Ismailov, 1981).

The results of thermal expansion measurements are presented in Figs. 8a and b. It can be observed that the slope is unlikely to be constant, varying from 10.9·10^{-6}/°C to 35.2·10^{-6}/°C (1–4 boxes in

![Fig. 7. Thermogravimetry and differential scanning calorimetry analysis of Bentheimer sandstone under the air and argon atmosphere, at heating rate 10 °C/min.]()
Fig. 8a). The most visible expansion occurs above 500 °C. The structure of the kaolinite breaks down in the endothermic reaction results into meta-kaolinite and a consequent distortion of structures (Fig. 6 and Table 1). Quartz undergoes a trigonal \( \alpha \)-quartz to hexagonal \( \beta \)-quartz transformation and above 800 °C disintegrated feldspars create a vitrified textures with melting illite (Fig. 6d and Table 1). The cooling curve of Bentheimer differs significantly from the heating curve; this difference is due to the elongation of quartz grains during heating.

To a lesser extent, the change in sample length between the beginning and the end of the experiment can be attributed to the greater force acting on the sample at the final stage of the experiment. For comparison, the linear thermal expansion of quartz is included in Fig. 8b. A similarity in the heating curve trend between Bentheimer and quartz is observed. As shown above, the deformation and expansion of the sample is mainly governed by quartz; however, at temperatures above 573 °C (where the transition from \( \alpha \)- to \( \beta \)-quartz occurs), the coefficient becomes negative (Fig. 8b), which suggests that more constituents than quartz control the expansion process in Bentheimer sandstone. The thermal expansion curves prove again that there are no high-temperature forms of quartz.

3.3. Porosity and permeability

Permeability and porosity of the Bentheimer Sandstone cores can be expected to vary as a consequence of firing along with texture and mineral alteration. Firstly, a differential thermal expansion may cause cracking of the grain contacts and by that increase of the pore volume. Secondly, the disintegration and recrystallization of minerals that have a lower melting point than quartz may slightly increase the quartz grain volume. As well the differential expansion of minerals separates grains at the interfaces (Fig. 6b) and thus, creates an irreversible volume increase (Kühnel, 1998; Wolf, 2006; Somerton, 1992; Bhargava et al., 2009). Fig. 9 shows that the absolute gas permeability against porosity hardly changed before and after thermal treatment. The porosity and permeability both increased by less than 5%, which is within the experimental error for usual core flood tests. The slight change in the porosity and permeability is most likely due to textural and compositional changes. As stated by Klimentos and McCann (1990) clays, even in small amounts, may strongly influence flow parameters of the rock. Due to exposure to high temperatures (\( \sim \)970 °C), clays undergo decomposition and degradation, and, as a result, change the sandstone matrix structure. On the other hand quartz undergoes reversible reaction at 573 °C from \( \alpha \)- to \( \beta \)-quartz. That is related to the increasing vibrations and slight displacement of the atoms in the crystal lattice, and further volume increase (Ghiorso et al., 1979). Moreover, the firing process leads to weakening and releasing of the cementing material, which results in dispersion and relocation of fine particles within the sandstone pore matrix (Ma and Morrow).

3.4. Surface charge

The potentiometric titration for establishing surface charge behavior was started in alkaline pH (\( \sim \)12). Comparing the titrated surface charge curve obtained from fired Bentheimer sandstone samples, Fig. 9b, to unfired (Peksa et al., 2015) shows that protons bind to surfaces at the highest protons affinity sites. Subsequently, with an addition of HCl the sites with lower proton affinity come to be protonated.

For unfired and fired samples the change in the pH value, with addition of 0.1 M HCl titrant, showed a similar trend for the same mass of the samples, however with a steeper gradient for fired samples so that the point of zero charge (PZC) for unfired and fired samples are respectively at \( \text{pH} = 8 \) and \( \text{pH} = 10.6 \). The shift in the PZC can be attributed to a higher concentration of iron oxides present in the sample after firing.

In the Bentheimer sandstone, the iron oxides coatings on quartz grains can give large local specific surface areas. At the low pH, they may reduce a negative charge due to (1) the physical blockage of negative charges by iron oxides coatings on e.g. quartz grains, (2) the mutual neutralization of negatively charged particles (Sumner, 1963). It proves the previous remark that as a result of firing iron becomes grain coating material, which can significantly influence the surface response of the sample. Iron oxides might be strongly oil-wet and result in a change of the wettability of Bentheimer sandstone. Nevertheless, in the terms of the volume, the iron oxides can be neglected.

3.5. Dielectric spectroscopy

The frequency dependence on the electric parameters for fired and unfired samples was examined at pressure of 2 bars, temperature of 30 °C. The obtained values were compared with predictions made using the model of Lichtenecker and Rother (1931).

3.5.1. Dielectric permittivity

The variation of the real part of electric permittivity \( \varepsilon_r \) and the dissipation factor \( \tan \delta \) as a function of frequency are depicted in Figs. 10a and b. The results represent average values over three consecutive tests. It is observed that the real part of electric permittivity decreases after the samples are thermally treated. Fig. 10a shows that over the frequency range of 0.1–10 Hz where the counter ion diffusion polarization occurs, the dielectric dispersion magnitude is \( \Delta \varepsilon_{\text{max}} = 0.5 \) (for sample 13A) and \( \Delta \varepsilon_{\text{max}} = 3.27 \) (for sample 12A). It decreases with increasing frequency reaching \( \Delta \varepsilon_{\text{max}} = 0.13 \) (for sample 13A) and \( \Delta \varepsilon_{\text{max}} = 0.39 \) (for sample 12A) over the frequency range of 10 Hz to 15 kHz. This phenomenon in the low frequency regimes is common for rocks and is attributed to the variation in polarization, being a result of the charge accumulation at the grain boundaries, and/or grain imperfections, and further, the composition heterogeneity in the rock matrix (Saint-Amant and Strangway, 1970; Sengwa and Soni, 2006). This has been explained previously by a Maxwell-Wagner model (Howell and Licastro, 1961). For high frequencies (>MHz) this effect will be negligible (Rao and Smakula, 1965).

For the thermally treated samples, the dielectric constants measured at low frequencies are up to \( \sim \)5% larger than those at the high frequencies, which is a much smaller variation than that observed in
the unﬁred samples. Differences in permittivity results between ﬁred and unﬁred samples are mainly due to the transformations of clay due to ﬁring. The high speciﬁc surface area minerals (i.e. clays) usually exhibit higher dielectric dispersion magnitudes than low speciﬁc surface area minerals (Arulanandan, 2003). The low dielectric constant for ﬁred samples that contain iron oxides (the large speciﬁc surface and capillary retention capacity) was discussed by Van Dam et al. (2002). They demonstrated that the iron oxides do not directly alter the relative permittivity of the solid phase in the sediment. However, the correlation exists between the larger speciﬁc surface and capillary retention capacity of iron oxides, as compared with quartz grains. Thus, iron oxides can have a profound inﬂuence on the relative permittivity if they occur in larger quantities than in the Bentheimer sandstone (TGA measurements and XRD).

3.5.2. Dissipation factor

Fig. 10b shows the dissipation factor vs. frequency. At low frequencies, losses are much higher than at higher frequencies for both ﬁred and unﬁred samples. For the ﬁred samples, at frequencies larger than 10 Hz, the dissipation factor is almost independent of frequency. This can be explained by a Maxwell-Wagner effect for a widespread distribution of conductivity of the accessory minerals and thermos-mineralogical alterations within Bentheimer (Saint-Amant and Strangway, 1970). With increasing frequency, a slightly different approach that inﬂuences the dissipation factor behavior, the Garton’s mechanism (Garton, 1941), is observed. More tests in higher frequencies should be conducted to entirely conﬁrm that statement. Note that this mechanism of invariance with frequency is a special case for quartz (Argall and Jonscher, 1968).

The observed general dispersion in the obtained results can be attributed to local heterogeneities due to varying clay concentrations in the Bentheimer samples. The variation in the dielectric constant is directly related to differences in the mineral composition, the crystal structure and the relation and interaction of the various constituents of the rocks to one another, besides clay.

The trend of a decrease in dielectric permittivity and the corresponding change in the dielectric properties have been ascertained by the extended drying process and repeated measurements. Attention was taken to eliminate any possible moisture and instrumental source causing variations in the measurements. While conducting the measurements, we found that there is a difference between the results obtained at certain stages of the drying. Hence, the measurements account as “dry rock” were obtained after the third drying when the value of the loss tangent varied <0.5% between repeated measurements. Fig. 11b shows the inﬂuence of water content on the dielectric constant of unﬁred sample no. 14 with different moisture content in the frequency domain covering the interval from 1 Hz to 1 kHz. The trend of spectrum variation is close to the dry rock samples, although a noticeable role of moisture is observed.

3.5.3. Polarization current and the dielectric absorption test

For the dielectric absorption test, the DC voltage $U=0.2$ kV was used. As a result, a polarization current, $I_p$, ﬂows through the dielectric. Subsequently with time, the transient phenomena “polarization current decay” heads to a small steady state conduction current (Fig. 12).

For both samples, the application of a DC electric ﬁeld results in a current jump as an effect of fast electronic, ionic and orientation polarization. Subsequently, the current decreases gradually due to slow polarization processes. A good ﬁt of the Curie-von Schweidler model (Curie, 1888; Schweidler, 1907) with the measured data was obtained.
The observed decay phenomena can be described by a power function, where the space charge formation creates an inner electric field of the opposite direction which progressively decreases the current. Fig. 12 shows the variation of the polarization current with time and sample treatment (unfired and fired).

The application of DC voltage for fired samples showed that changes in mineral composition and their conductivity tend to affect the tail of polarization currents. Due to firing and related dispersion of the iron oxides within the matrix framework, Bentheimer sandstone becomes a weaker insulator (Table 2).

### 4. Conclusions

We have investigate the firing of the Bentheimer samples, a commonly used procedure in core preparation, to make the internal rock properties uniform and improve reproducibility of displacement experiments. The petrophysical, petrological pyrometamorphical and electrical properties of unfired and fired samples were measured using various complementary methods. The following main conclusions can be drawn:

In unfired samples, ion exchange between minerals (mainly clays) and injected fluids caused clay migration (kaolinite and illite) and swelling (illite and smectite) leading to partial or total plugging. The local clay concentrations, differences in the crystal structure and the interaction of the various constituents were validated by dispersion in the dielectric permittivity over frequencies.

Firing of Bentheimer sandstone above 900 °C led to gradual changes in the samples mineral composition, color, surface roughness and physical properties, through transformation, disintegration and pyrometamorphism of the components.

Fine loose particles were created by thermal disintegration and still may migrate during liquid flow experiments. The conducted XRD study and the decrease in the permittivity value with its insignificant dispersion (∼5%) between the high and the low frequencies indicated the clay transformations.

Thermal stresses also caused variation in the pore geometry, change in grain boundaries and contacts and particle cracking, i.e. as a result, the porosity and permeability increased by up to 5% and samples became more friable.

The mechanical strength of cementing material decreased due to dehydration-rehydration and heterogeneous thermal expansion. The boundary between the detrital quartz grains became detached by better connected, larger and wider throats.

Firing resulted in decomposition of minerals with Fe content and caused formation of source of amorphous oxides that influence the wettability. The iron oxides do not directly alter the relative permittivity of the solid phase. However, the correlation exists between the clay transformations.

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### Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.petrol.2016.10.024.
