LETTER

Electrical impedance measurement of hydrous microcrystalline quartz

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High temperature impedance spectra, up to 450 °C, were measured for a microcrystalline quartz aggregate (chalcedony), which initially contained 0.3 wt% of liquid-like water (H₂O) dispersed in grain boundaries and fluid inclusions together with 0.3 wt% of hydroxyl (Si-OH) in the crystal structure. Infrared spectra obtained after heating, showed dominant dehydration of liquid-like water, while much hydroxyl remained stable. Electrical conductivities before (wet) and after heating to 450 °C (dry) gave linear Arrhenius relations with apparent activation energies of 11 ± 1 kJ/mol for initial heating of the wet sample versus 32 ± 3 kJ/mol for the subsequently dry sample. Compared with previously reported Arrhenius relations for α-quartz single crystals, our activation energies are much lower, and the absolute conductivities we obtained range from similar values to three orders of magnitude higher. We infer that the presence of grain boundaries and/or triple junctions containing liquid-like water greatly influences the electrical conductivity.

Keywords: Liquid-like water, Microcrystalline quartz, Electrical conductivity, Dehydration, Chalcedony, Impedance spectroscopy

INTRODUCTION

It is well known that natural rocks contain molecular water confined in grain boundaries, triple junctions and fluid inclusions (hereafter referred to as liquid-like water; Nakashima et al. 2004). Liquid-like water has significant influences on the rheology of the Earth’s crust through deformation processes such as pressure solution, fluid-assisted recrystallization, grain growth, and fluid-assisted cataclasis, as summarized in Dysthe and Wogelius (2006). Of key importance in controlling these processes and the associated rheology is the distribution and properties of the liquid-like water phase. Electrical conductivity is a useful tool for evaluating the distribution and properties of liquid-like water in rocks, since it is highly sensitive to conduction through high diffusivity, fluid-filled pathways (including hydrous fluids and melts).

Numerous data on electrical conductivity have been reported for various materials, including single crystals, glass and grain aggregates, many of which are summarized in Watanabe (2005). Recently, electrical conductivity has been measured for natural crustal rocks dependent on rock texture, under high pressure-temperature conditions (e.g., Fuji-ta et al., 2004; Fuji-ta et al., 2007). Also, several attempts to evaluate the effects of liquid-like water on electrical conductivities have been reported for a few simple granular aggregates (e.g., Watanabe and Peach, 2002 for salt rock; Wang et al., 2006 for olivine aggregates) and for bi-crystal contacts of salt rock undergoing pressure solution (De Meer et al., 2005; Van Noort et al., 2007). However, materials used in experiments and data are thus limited and the influence of liquid-like water on electrical conductivities in low permeability crystalline rock and fine grained microporous material is poorly understood.

We report the changes of electrical conductivity associated with dehydration of liquid-like water in microcrystalline quartz, viewing it as an analogue for quartz-rich rocks in the Earth’s crust. Electrical impedance spectra were continuously measured at several high temperatures up to 450 °C. The difference in electrical con-
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DUCTIVITY BETWEEN INITIALLY WET AND SUBSEQUENTLY DRIED SAMPLES IS ALSO ADDRESSED AND AN ATTEMPT IS MADE TO UNDERSTAND THE DISTRIBUTION OF WATER IN THE SAMPLES.

MATERIALS AND EXPERIMENTAL METHODS

Microcrystalline chalcedonic quartz samples were collected from Chayagawa, Oshamanbe, South-West Hokkaido, Japan. The water content and stability at high temperature of such a sample was roughly determined by in-situ infrared (IR) spectroscopy in our previous study, which demonstrates that liquid-like water gradually dehydrates at less than 400 °C (Fukuda and Nakashima, 2008). Disc-shaped samples of 13 mm diameter and between 1.1–1.2 mm thick were used for the impedance measurement. To ensure good electrical contact, disc-shaped electrodes of 8 and 12 mm diameter were deposited onto the two opposite faces of the samples by gold sputtering in a partial vacuum. A guard ring of 1 mm was also sputtered around the 8 mm electrode to ensure conduction directly through the sample.

The complex impedance spectra were measured using a Solartron 1260A frequency response analyzer with a Solartron 1296A dielectric interface for an increased impedance range up to $10^{14}$ Ω and capacitance down to $10^{-12}$ F. Systematic measurement errors with the combined apparatus are less than 2%. The conductivity cell, constructed of refractory alumina (Ten Grotenhuis et al., 2004), was equipped with 0.5 mm thick Pt electrodes and guard ring, of similar areal dimensions to the sputtered sample electrodes, to ensure good connection via shielded platinum wires. This was situated inside a 60 mm internal diameter tube furnace, with temperature controlled and monitored using Pt/Pt – 10%Rh thermocouples. The real $Z'$ and imaginary $Z''$ components of the impedance were obtained in the frequency range of 0.1 Hz to 1 MHz with a periodic sinusoidal r.m.s. voltage of 200 mV. The complex impedance spectra were continuously measured at constant temperatures of 250, 350, and 450 °C under open

Figure 1. (a) Complex impedance spectra series; an example of continuous heating at 350 °C. The changes of arcs are shown by time-line. One single arc grows less circular and larger with exposure time. These arcs contain contributions from sample interior (bulk and grain boundaries (GB in the figure) or triple junctions/inclusions containing liquid-like water) and predominant charge-up of the electrodes (seen at lower frequencies; most of visible data points are <1 kHz). Changes in arc diameter above 1 kHz are due solely to dehydration whilst those at lower frequencies include charging of the electrodes. This influence becomes larger with exposure (heating) time, excessively stretching the arcs towards larger apparent real impedances (resistances). (b) Changes of corresponding relative permittivities. Permittivities are frequency independent above 1 kHz, but grow rapidly with decrease of frequency below that value (phenomenon of low frequency dispersion; possible ionic-charge buildup on electrodes in layers with associated high capacitance). (c) Some fitted arcs to simple RC circuit below 1 kHz (30 data points per arc).
atmosphere conditions for periods up to 120 hours, and spectral changes associated with dehydration were obtained as a function of time.

Before and after the experiments, the liquid-like water (H$_2$O) and hydroxyl (Si-OH) contents were separately determined using IR spectroscopy (Jasco FT-IR 470 + IRT-30 instruments). The IR apparatus is equipped with a Globar (silicon carbide) IR source, a Ge-coated KBr beamsplitter, and a mercury-cadmium-telluride detector. The spectra were obtained by multiple scanning (256 times) with a resolution of 4 cm$^{-1}$.

RESULTS AND DISCUSSION

Changes of complex impedance spectra and relative permittivity spectra at 350 °C are shown in Figures 1a and 1b, respectively, as an example. Roughly, a single arc grows larger with increased exposure time to heating. Since the solid-state conductivity of dry quartz is very low at these conditions (Jain and Nowick, 1982), conductivity is expected to be dominated by other, mainly intercrystalline pathways. The arcs can be considered to be due to the behavior of the sample interior and sample electrode interface effects. The sample interior components will contain contributions from bulk quartz, and its grain boundaries, triple junctions or inclusions containing liquid-like water. Electrode interface effects will be related to charge transfer (from/to ions in solution to/from electrons in metal connectors, as redox reactions) and capacitive storage charge/discharge (e.g., via Gouy–Chapman double-layers). At low frequencies, electrode interface effects are confirmed by the presence of low frequency dispersion or LFD (Jonscher, 1991), seen as a significant increase in cell-capacitance towards lower frequencies. This is most likely due to the charge-up of the electrode by accumulation of transported ions in double-layer structures. Above 1 kHz, the cell capacitance (that corresponds to cell permittivity) is frequency independent (Fig. 1b) and so only this part of the arcs was used to estimate the conduction related impedances from a simple parallel RC circuit model, where capacitance C, must be assumed constant to extract the resistance R, related to ionic transport, from the arc diameter. The size of the arcs grows with continued dehydration as can be seen by measuring continuously (Fig. 1a), but the high frequency changes in arc diameter are less than this figure suggests since the majority of the visible data is for frequencies below 1 kHz and is also affected by a frequency dependent growth in capacitance and interface resistance (LFD). By only using the constant cell capacitance, LFD free data above 1 kHz, we avoid the complications of the electrode charging effects. The results of arc fitting are shown in Figure 1c. Electrical conductivity, $\sigma$ originating from the sample interior is calculated by $\sigma = d/SZ'$, where d is the sample thickness, S is the measured area (8 mm diameter electrode), and Z$'$ is the real part of the impedance or the resistance, R. This was determined from the intersection of the arc, projected from just the data segment above 1 kHz, with the real impedance axis (pure resistance) by numerically fitting a simple parallel RC circuit model (Fig. 1c), as a best fit semicircle. Some fitted arcs are shown in Figure 1c. When the data above 1 kHz is viewed more closely then there is no sign of additional arcs that could suggest any separate pathways or additional mechanisms for conduction, but the data do fall over a more circular single arc, with very little depression, allowing a good fit to the simple parallel RC model. Changes of electrical conductivity at each temperature against the duration of heat treatment (heat exposure time) are shown in Figure 2.

IR spectral analysis shows the changes in water content before and after the experiment (Fig. 3). Water content, including liquid-like water (H$_2$O) and hydroxyl (Si-OH), can be calculated separately from each coupling vibration mode of stretching and bending, which can be recognized around 5200 and 4500 cm$^{-1}$, respectively (Graetsch et al., 1985). This calculation gives the true water content to the first decimal place, but the relative value can be more closely compared within the same sample as Figure 3. Initially, water content, as we described, was 0.32 wt% of liquid-like water and 0.28 wt% of hydroxyl in Figure 3. After the experiment, dominant dehydration of liquid-like water is confirmed, while much hydroxyl remains stable. In the region of the water stretching band around 3800–3000 cm$^{-1}$, a new peak at 3740 cm$^{-1}$ can be recognized after the heating experiment, especially after exposure to higher temperatures. This peak is considered
to be the isolated surface silanol, which is not hydrogen bonded with liquid-like water and appears after dehydration of liquid-like water starts (Yamagishi et al., 1997). According to the results of IR spectroscopy, the changes of resistance (Fig. 1) and conductivity (Fig. 2) with heat treatment duration or exposure time must mainly reflect the dehydration of liquid-like water.

In Figure 2, the initial values of conductivity can only reflect the temperature dependence of the hydrous samples, since liquid-like water trapped in grain boundaries or triple junctions, or structurally incorporated as hydroxyl, has not yet been driven out of the samples. The initial value at each temperature is $4.0 \times 10^{-7}$ S/m at 450 °C, $3.3 \times 10^{-7}$ S/m at 350 °C, and $2.1 \times 10^{-7}$ S/m at 250 °C. Separate from the dehydration experiments at these three temperatures, initial conductivity at 200 °C for a fourth wet sample was determined and the value was $1.5 \times 10^{-7}$ S/m.

With exposure to high temperatures, liquid-like water dominantly dehydrates as described above. In the case of 450 °C, conductivity quickly decreased from $4.0 \times 10^{-7}$ S/m to an almost constant value (ca. $2.7 \times 10^{-7}$ S/m) from 5 hours onwards. Coupled with the result of IR spectroscopy (Fig. 3), these constant values correspond to the state at which the sample has no liquid-like water. Therefore, these values come from grain boundaries and quartz itself. At temperatures of 250 and 350 °C, although both conductivities reached similar values during the experimental heat treatment, the conductivities obtained depend on both the water contents and temperatures. That is specifically considered as follows. Calculated water contents after the experiments show dominant and different degree of dehydration of liquid-like water depending on heating temperatures; 0.14 and 0.27 wt% for after 350 °C and 250 °C, respectively (Fig. 3). These differences of water contents have influences on values of electrical conductivities. On the other hand, the conductivity increases with temperature increase even in the same sample composition. The combination of water content and temperature reflect the obtained similar value of conductivities at 350 and 250 °C.

After the experiment at 450 °C, i.e., for the dry sample, conductivity was again measured at several temperatures down to 100 °C. These values together with the initial values (i.e., wet) for the other samples at each temperature described above were plotted in an Arrhenius
Figure 4. Arrhenius diagram of electrical conductivity (S/m) for before (initial) and after prolonged heating (dry) at 450 °C. Water contents in the form of liquid-like water (H₂O) and hydroxyl (Si-OH) are also shown. Conductivity becomes lower and activation energy becomes higher for the dry sample, indicating the additional conduction due to liquid-like water in the wet (initial) sample. The typical value for a natural α-quartz single crystal parallel to crystallographic c-axis reported by Jain and Nowick (1982) is also shown as solid line for their measured data and dotted line for extrapolated data.

Apparent activation energies for the dry and wet sample were determined from the Arrhenius relation; $\sigma = \sigma_0 \exp(-E/RT)$, where $\sigma$ is the conductivity, $\sigma_0$ is the conductivity at infinitely high temperature, $E$ is the apparent activation energy, $R$ is the molar gas constant, and $T$ is absolute temperature. The data for the dry sample were fitted excluding that of 450 °C, since the data at 450 °C seems to deviate from other four temperature data and may represent a conduction process more intrinsic to the bulk quartz rather than conduction along grain boundaries or triple junction tubes. This calculation gives the apparent activation energies and pre-exponential factors $\sigma_0$ of respectively 11 ± 1 kJ/mol and 0.28 ± 10⁻³ for the wet sample, versus 32 ± 3 kJ/mol and 0.38 ± 10⁻⁴ for the dry sample. Good linearity can be seen. However, water on the saturation liquid vapor curve is known to exhibit a maximum conductivity at around 230 °C (IAPWS, 1990), the conductivity at infinitely high temperature, or triple junction tubes. This calculation gives the apparent activation energy, $E = 11 \pm 1$ kJ/mol (H₂O: 0.32 wt%, Si-OH: 0.28 wt%) for the dry sample at 450 °C is best explained by a change of charge carrier over 450 °C from water in grain boundaries to the ionic conduction within quartz itself as given by the Jain and Nowick (1982) data trend in Figure 4.

The distribution of liquid-like water can be roughly evaluated. We consider here the water distribution at 350 °C, for example, using reported conductivity data for pure water from IAPWS (1990). The liquid-like water content of 0.3 wt% translates to 0.77 vol%. The grain size must correspond to the crystallite size of 30 nm (Fukuda and Nakashima, 2008). If the fluid is distributed evenly on grain faces then the water content represents a layer 0.1 nm thick, which is only half a monolayer of water molecules (O–H bond length is approximately 0.096 nm). In this restricted state, bound between adjacent grains, water is not likely to behave as liquid water. However, water is observed to be present in a liquid-like state, (Fukuda and Nakashima, 2008; as inferred from in situ IR spectroscopy) so for the thickness of boundary water to be great enough to allow such freedom to molecules then it must be partially covering (<50% of grain faces) and possibly disconnected. Hence, a distribution in triple junction tubes is more likely. Therefore, we assume that liquid-like water is evenly distributed in fully-connected grain-edge tubes. Taking the electrical conductivity of pure water as $6.0 \times 10^{-3}$ S/m at 350 °C (fluid density = 0.7 g/cm³), a connected grain-edge tube model (Grant and West, 1965; Schmeling, 1986; Ten Grotenhuis et al., 2005) then gives the calculated conductivity as $2.7 \times 10^{-6}$ S/m, which is about one order of magnitude higher than the experimentally determined value of $3.3 \times 10^{-7}$ S/m. Conductivity of pure water has a maximum at 230 °C, a factor of ~15 higher than at 350 °C, as described above. So, the fully-connected grain-edge model overestimates the bulk conductivity by more than 2 orders at the lower temperatures studied. Therefore, we infer that liquid-like water in tubes must either expand, (limited by thickness restrictions argued above), to fill grain boundaries lowering its density and hence conductivity at high temperatures or it becomes lost during heating prior to measurement, or the fluid-filled tubes are partially disconnected before or during heating.

We conclude from our results that below 350 °C it is the presence of grain boundaries and of liquid-like water
within triple junction tubes, or penetrating grain boundaries on heating, that leads to the high electrical conduction seen in comparison with quartz single crystals.

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