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ORIGINAL PAPER

Modification of fluid inclusions in quartz by deviatoric stress. II: experimentally induced changes in inclusion volume and composition

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Abstract Fluid inclusions in guartz are known to modify their densities during shear deformation. Modifications of chemical composition are also suspected. However, such changes have not been experimentally demonstrated, their mechanisms remain unexplained, and no criteria are available to assess whether deformed inclusions preserve information on paleofluid properties. To address these issues, quartz crystals containing natural CO₂-H₂O-NaCl fluid inclusions have been experimentally subjected to compressive deviatoric stresses of 90-250 MPa at 700°C and ~ 600 MPa confining pressure. The resulting microcracking of the inclusions leads to expansion by up to 20%, producing low fluid densities that bear no relation to physical conditions outside the sample. Nevertheless, the chemical composition of the precursor inclusions is preserved. With time the microcracks heal and form swarms of tiny satellite inclusions with a wide range of densities, the highest reflecting the value of the maximum principle

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Present Address: A. Tarantola UMR-G2R, Université Henri Poincaré, 54506 Vandoeuvre-lès-Nancy Cedex, France stress, σ_1 . These new inclusions lose H₂O via diffusion, thereby passively increasing their salt and gas contents, and triggering plastic deformation of the surrounding quartz via H₂O-weakening. Using microstructural criteria to identify the characteristic types of modified inclusions, both the pre-deformation fluid composition and syn-deformation maximum stress on the host mineral can be derived from microthermometric analysis and thermodynamic modelling.

Introduction

Much of the success in applying fluid inclusion studies to understand processes in the Earth's Crust has derived from the fact that many minerals, notably quartz, behave as rigid, chemically impermeable hosts over the pressure– temperature range of interest. Thus, parameters such as paleofluid composition (X) and molar volume ($V_{\rm m}$), or density, can be derived by treating the inclusions as isochoric-isoplethic thermodynamic systems.

Nevertheless, numerous cases are known in which natural fluid inclusions have not preserved their original $V_{\rm m}$ -X properties since entrapment. Several processes may be responsible for such post-entrapment modifications: at high temperature, diffusion through the host mineral may become fast enough to effect mass transport between the inclusion and its surroundings in response to gradients in chemical potential; changes in rock temperature and in the dependent internal pressure of the inclusions may drive chemical reactions between the inclusion and the host mineral; and changes in external stresses may

irreversibly modify inclusion volumes. Extraction of useful information from such inclusions relies on criteria to recognize post-entrapment modifications a posteriori and on methods to reconstruct by calculation the net exchanges of chemical constituents and the changes in molar volume. Among the experimentally investigated modifications of fluid inclusions in quartz are the diffusive loss or gain of H_2 and H_2O (e.g. Morgan et al. 1993; Mavrogenes and Bodnar 1994; Hall and Sterner 1995; Bakker and Jansen 1991, 1994; Sterner et al. 1995). These results have been applied to studies of natural samples (e.g. Ridley and Hagemann 1999). Plastic deformation of the host crystal is also known to affect the $V_{\rm m}-X$ properties of fluid inclusions (Sterner and Bodnar 1989; Vityk et al. 2000; Schmidt et al. 2003). For example, homogenization temperatures, which are dependent on both composition and molar volume, typically become spread over a wide range according to the state of intracrystalline deformation. Küster and Stöckhert (1997) argue that rates of plastic strain as low as 10⁻¹⁵ s⁻¹ may cause fluid inclusions to adapt their volumes to the ambient stress regime. It has been postulated that crystal-plastic deformation of quartz drives preferential loss of H₂O from homogeneous H₂O-CO₂ fluid inclusions, thereby generating the pure CO₂ fluid inclusions found in high-grade metamorphic rocks (Hollister 1988, 1990; Bakker and Jansen 1990; Lamb and Moecher 1992; Johnson and Hollister 1995) and in ore deposits hosted by shear-zones (e.g. Schwartz et al. 1992; Klemd et al. 1997; Schmidt Mumm et al. 1997).

To understand the effects of crystal-plastic deformation on fluid inclusion properties, we have subjected natural, fluid-inclusion-bearing quartz crystals to uniaxial compressive (deviatoric) stresses at 700°C and ~600 MPa. Under these conditions quartz deforms by crystal plasticity. In a companion paper (Tarantola et al. 2010), the resulting modifications of inclusion shapes and microstructures are reported and interpreted. In the present paper, the focus is on modifications of inclusion composition and molar volume (density).

This article first presents the microthermometric and Raman spectroscopic characteristics of the selected fluid inclusions prior to the experiments. The thermodynamic model that is subsequently used to assess the V_m -X modifications is explained. The observed changes in microthermometric and Raman spectroscopic properties after the experiments are then compared to the original values to quantify the modifications. A discussion of the collective microstructural, compositional and volumetric evidence sheds light on the possible mechanisms of the modifications. These in turn suggest criteria by which natural fluid inclusions in plastically deformed quartz can be interpreted.

Methods

Piston-cylinder experiments

Inclusion-bearing quartz samples were placed in a solid medium, Griggs-type piston-cylinder apparatus and held at 700°C. Several preparatory experiments were run under hydrostatic confining pressure at ~600 MPa (Hyd-5 and Hyd-6, Fig. 1) and at ~800 MPa (Hyd-4, Fig. 1). Six deformation experiments were performed by compressing the samples uniaxially with a constant load for periods of 12–136 h. The four of these experiments that were buffered with respect to H₂ fugacity (Def-1, Def-2, Def-3 and Def-6, Fig. 1) are treated in this article. Details of the experimental method are given in the companion paper (Tarantola et al. 2010).

Fluid inclusions

Quartz plates for the experiments were cut from single, euhedral, freely grown crystals from hydrothermal veins



Fig. 1 *P*–*T* conditions of hydrostatic experiments (squares labelled Hyd-) and deformation experiments involving deviatoric stresses (square labelled Def-). *Stepped paths* show compression-heating and decompression-cooling trajectories of experiments. Fan of isochores delineating entrapment conditions of all inclusions from Brusson sample locality is taken from Diamond (1990). Extrapolation of fan yields hypothetical internal pressures of all inclusions at 700°C. Shaded fan of isochores represents subset of inclusions used as "precursors" in experiments. Inclusions in experiment Hyd-4 (800 MPa) experienced negative differential pressure ($P_{inc} \ll P_{conf}$). Inclusions in Hyd-5,6 and Def-1,2,3,6 (600 MPa) experienced negligible differential pressure ($P_{inc} \approx P_{conf}$). Quartz phase boundary from Hosieni et al. (1985). Approximate position of fluid solvus interpolated from data in Gehrig (1980) and Krüger and Diamond (2001)

at the Fenilia gold mine, Brusson, NW Italian Alps, as described by Diamond (1990; sample LD888). All the inclusions used in the experiments were originally trapped from a homogeneous (1 phase) fluid. Although crystals from the same veins also contain heterogeneously trapped inclusions, namely CO_2 -rich "vapour" inclusions coexisting with immiscible CO_2 -poor "liquid" inclusions (Diamond 1990), these were purposefully excluded from the present experiments. This point is important, as changes in the volume fractions of inclusion phases are induced during the experiments. Any confusion with originally gas-rich inclusions must therefore be avoided.

At room temperature, the inclusions consist of three fluid phases: a weakly saline aqueous liquid containing 4.0–5.5 mass% salts (dominantly NaCl) and a bubble of carbonic liquid enclosing a bubble of carbonic vapour. The carbonic phases are dominated by CO₂ with minor N₂ (3–4 mol%) and trace CH₄ (<0.07 mol%). Some inclusions contain accidentally captured crystallites of muscovite or dolomite. The bulk composition is 90.4–92.2 mol% H₂O and 6.4–4.6 mol% CO₂, the remainder being chloride salts and minor gases. The quartz crystals were formed in open cavities at temperatures between 225 and 300°C. Further details are given by Diamond (1990).

Characterization of fluid inclusions prior to experiments

Between 50 and 720 fluid inclusions were selected per sample to represent a range in shape, size and distance to the sample surface. The inclusions were numbered and individually photographed and their positions were mapped to permit relocation after the experiments. Two characteristic phase transitions were measured by microthermometry in 12-84 fluid inclusions in each sample before the experiments. The temperature of clathratehydrate dissociation, $T_{\rm m}$ (cla), was measured as a monitor of salinity (Diamond 1994). Values lie between 7.5 and 8.5°C and dissociation always occurs in the presence of the so-called Q_2 phase assemblage: clathrate + aqueous liquid (L_{aq}) + carbonic liquid (L_{car}) + carbonic vapour (V_{car}). The temperature of homogenization of the carbonic phases, $T_{\rm h}({\rm car})$, was measured as a monitor of bulk density (Diamond 2003). Homogenization occurs between 25.1 and 28.5°C via all three possible modes: bubble-point transitions $(L_{aq}L_{car}V_{car} \rightarrow L_{aq}L_{car})$, critical transitions $(L_{aq}L_{car}V_{car} \rightarrow L_{aq}SCF_{car})$, where SCF denotes supercritical fluid) and dew-point transitions $(L_{aq}L_{car}$ $V_{car} \rightarrow L_{aq}V_{car}$). For convenience in the presentation of the experimental results, the temperatures of these three transitions are denoted $T_{bub}(car)$, $T_{crit}(car)$ and $T_{dew}(car)$, respectively. According to the specific inclusion generation, the critical point of the carbonic mixture lies between 28.1 and 28.5°C. To avoid any deformation or leakage of the inclusions prior to the experiments, the samples were held below 35°C. Therefore, the total homogenization temperatures, known from Diamond (1990) to lie in the range 225–270°C, were not measured.

Microthermometry and Raman microspectroscopy were carried out in the Fluid Inclusion Laboratory of the Institute of Geological Sciences, University of Bern. A LinkamTM THMSG heating-cooling stage was employed, and calibrated using the following phase transitions in synthetic fluid inclusions in quartz: the triple point of CO₂ at -56.6°C, the triple point of H₂O at 0.0°C and the critical point of H₂O at 374°C. The stage was mounted on an OlympusTM BX51 microscope equipped with $40 \times$ and $100 \times$ long-working distance objectives. Particular attention was paid to the precision to which phase transitions could be measured before and after the experiments. Measurements were initially made on inclusions within the \sim 2-mm thick plates. After the experiments, most of the plates were polished down to as little as 150 µm thickness to remove the superficial extension fractures and thereby enhance visibility of the target inclusions.

The LinkamTM microthermometry stage operates by vertical heat conduction and the thermocouple is embedded in the heating block. Therefore, samples of different thickness may potentially experience different thermal gradients and hence display different apparent temperatures of phase transitions. To minimize this effect, phase transitions were measured at a heating rate of 2° C min⁻¹, and for measurements of $T_{\rm m}$ (cla) a pause of 20–30 s was made before cooling steps during cycling (Diamond 1992). The overall uncertainties in microthermometry were quantified by measuring 28 fluid inclusions in a standard sample of 2.10 mm thickness. The sample was then thinned to 1.00 mm, and the remaining 27 inclusions were measured again. Finally, the sample was thinned further to 0.25 mm and the surviving 16 inclusions measured a third time. Figure 2 illustrates that, despite the variation in sample thickness, the $T_{\rm h}({\rm car})$ measurements are reproducible to within $\pm 0.1^{\circ}$ C and the $T_{\rm m}$ (cla) measurements to within ± 0.2 °C. In view of this excellent reproducibility, no additional corrections were applied to the microthermometric data.

The carbonic phases of individual inclusions were analysed for H₂, N₂, CH₄ and CO₂ at a few degrees above $T_{\rm h}({\rm car})$ using a Jobin–YvonTM LabRAM HR800 laser-Raman spectrometer. Spindle-stage measurements using the method of Bakker and Diamond (2006) showed that the volume fraction of the carbonic phase, $\varphi({\rm car})$, lies within the narrow range 0.18–0.21 at $T_{\rm h}({\rm car})$. An example of the measurements is given in "Appendix 1". Fig. 2 Reproducibility of microthermometric measurements of $T_{\rm h}({\rm car})$ and $T_{\rm m}$ (cla) as a function of thickness of quartz sample. Data from 28 inclusions measured in a plate 2.10 mm thick (x-axis) are compared to data from the same inclusions measured after thinning the plate successively to 1.00 and 0.25 mm (y-axis)



Theoretical basis to monitor changes in inclusion density and composition

Thermodynamic model

An appropriate thermodynamic model is required to calculate the bulk molar volume (density) and composition of the inclusions from the various measurable parameters. As the 4 species N₂, CO₂, H₂O and NaCl account for more than 99.8 mol% of the inclusions chosen for the experiments (Table 4 in Diamond 1990), the low-T behaviour of the inclusions can be confidently modelled within the N₂- CO_2 -H₂O-NaCl system. Estimation of the bulk V_m -X properties thus requires a minimum input of 4 measured parameters. Here, we follow the approach of Diamond (1990, 2003), using $T_{\rm m}$ (cla) and $T_{\rm h}$ (car) obtained by microthermometry, $\varphi(car)$ estimated optically at $T_{\rm h}(car)$, and the molar ratio of N₂/CO₂ determined by Raman spectroscopy. It turns out that the first three parameters undergo changes during the experiments, whereas in the hematitebuffered experiments the N₂/CO₂ ratio remains constant.

Figure 3a shows the relevant phase diagram constructed according to the method of Diamond (1994). The mean observed critical temperature of the carbonic mixture is 28.4°C. According to the EoS of Peng and Robinson (1976), this temperature implies that the homogenized carbonic phases consist of 96.2 mol% CO2 and 3.8 mol% N₂, a composition which agrees well with the actual Raman analyses. Figure 3a includes the corresponding $L_{aq}L_{car}V_{car}$ envelope, bounded by the dew- and bubble curves and projected from excess Laq. The amount of aqueous phase in the inclusions can thus vary without influencing the P-Tlocus of the envelope.

The thin curves traversing the envelope are isochores (calculated with the Peng-Robinson EoS) that span the observed range of densities of the carbonic phases following the experiments (see "Results"). The shaded band of

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isochores encompasses the observed variation in carbonic phase densities prior to the experiments. To illustrate how changes in molar volume of the carbonic phase, $V_{\rm m}({\rm car})$, are reflected by changes in $T_{\rm h}({\rm car})$, consider an inclusion that originally homogenizes at 28.4°C at the critical point (point 5, Fig. 3a), at which $V_{\rm m}({\rm car}) = 102 \text{ cm}^3 \text{ mol}^{-1}$. Any experimentally induced reduction in the molar volume of the carbonic phase will depress $T_{\rm h}({\rm car})$, such that homogenization occurs via a bubble-point transition. For example, reduction of $V_{\rm m}({\rm car})$ to 57.2 cm³ mol⁻¹ leads to homogenization of the carbonic phases at point 1, where $T_{\rm bub}({\rm car}) = 12^{\circ}{\rm C}$. It is important to appreciate that any experimentally induced increase in $V_{\rm m}({\rm car})$ will also depress $T_{\rm h}({\rm car})$, but in this case the carbonic phases will homogenize via a dew-point transition. For example, an increase in $V_{\rm m}({\rm car})$ to 172.3 cm³ mol⁻¹ leads to homogenization at point 7, where $T_{dew}(car) = 24^{\circ}C$. To aid display of the microthermometric measurements of $T_{\rm h}({\rm car})$ in the "Results", the hoop-shaped $L_{ao}L_{car}V_{car}$ envelope, consisting of the bubble curve, critical point and dew curve, is straightened out to serve as a one-dimensional axis, as shown along the x-axis of Fig. 3b. Although the direction of increasing temperature switches along this axis, it has the advantage of correlating directly with a monotonic increase in molar volume of the carbonic phase.

Also shown in Fig. 3a is the steep dissociation curve of the N_2 -CO₂ clathrate-hydrate solid-solution, which is in equilibrium with the stipulated composition of the carbonic phase. The curve (labelled $ClaL_{aq}L_{car}$ at high pressure and labelled "Q2 segment" where it cuts the L_{ag}L_{car}V_{car} envelope) was calculated using the curves code of Bakker (2003) for an aqueous NaCl concentration of 3.7 mass%. Progressive increase in aqueous NaCl concentration drives this curve to lower temperatures. Therefore, the aqueous salinity of the inclusions is calculated from $T_{\rm m}$ (cla) by incrementing the modelled NaCl concentration until the temperature at which the inclusion





Fig. 3 a Low *P*–*T* phase relations of the fluid inclusions used in the experiments, projected from excess aqueous liquid (L_{aq}). *Steep curve* is upper stability limit of clathrate. *Fine lines* are isochores. *Grey field* encompasses observed range of isochores prior to experiments; other isochores represent observed range after experiments. The carbonic phases (L_{car} and V_{car}) homogenize on the boundary of the phase envelope made up of the *bubble curve*, critical point (marked 5), and *dew curve*. **b** Effect of addition of H₂ on measurable phase transitions. Precursor inclusions that originally homogenized on the *bubble curve* inclusions that originally homogenized on the right. **c** Phase

isochore intersects the dissociation curve (e.g. point 8) matches the observed $T_{\rm m}$ (cla) (e.g. 8.0°C) (Diamond 1994). Any experimentally induced loss of H₂O from the inclusions will increase the residual salinity and shift $T_{\rm m}$ (cla) to lower values. Similarly, any addition of H₂O to the inclusions will shift $T_{\rm m}$ (cla) to higher values. Figure 3a shows that the Q₂ segment of the clathrate dissociation curve is extremely steep. When the precision of $T_{\rm m}$ (cla) measurements is taken into account (±0.2°C), the Q₂ segment can be treated as essentially vertical (isothermal). This means that any experimentally induced

relations showing how increased content of CH₄ (owing to high-*P*–*T* influx of H₂ into the inclusions) shifts the phase envelope and the clathrate stability field. *Bold dashed* phase boundaries show states without CH₄ (as in Fig. 3a); *continuous curves* show states when carbonic phases contain 10.4 mol% CH₄. **d** Vertical cross-section (to scale) of the sample capsule during the experiments. To test the effectiveness of the H₂ buffer, Fe₂O₃ powder was packed around the lower alumina piston as shown. Resulting microthermometric values of the inclusions are plotted in **b** (*diamonds* show inclusions near buffer; *circles* show inclusions far from buffer)

changes in $V_{\rm m}({\rm car})$ will not affect $T_{\rm m}({\rm cla})$. Shifts in $T_{\rm m}({\rm cla})$ therefore serve as a strict monitor of changes in aqueous salinity. Thus, values of $T_{\rm m}({\rm cla})$ can be recast in terms of aqueous NaCl concentration to serve as a parallel axis in the plots of microthermometric data (e.g. y-axis in Fig. 3b).

The bulk V_m -X properties of the inclusions may be calculated by first extracting the following 3 parameters from the above phase diagram: the pressure inside the inclusions at $T_h(car)$, denoted $P_h(car)$; the molar volume of the carbonic phases, $V_m(car)$ at $T_h(car)$; and the mole

fractions of H₂O and NaCl in the H₂O–NaCl binary subsystem (here termed "brine"), $X_{H_2O}^{brine}$ and X_{NaCl}^{brine} . The mole fractions of N₂ and CO₂ in the carbonic phase at $T_h(car)$, $X_{N_2}^{car}$ and $X_{CO_2}^{car}$, are directly known from Raman analysis. The concentrations of N₂ in the aqueous phase and of H₂O in the carbonic phase can be neglected under these conditions (Akinfiev and Diamond 2003). In contrast, the concentration of CO₂ in the aqueous phase, $X_{CO_2}^{aq}$, is significant and is obtained from the EoS by Akinfiev and Diamond (submitted) along with the molar volume of the aqueous phase, $V_m(aq)$, now treated as a CO₂–H₂O–NaCl ternary. All these parameters are inserted into the following set of volume- and mass-balance equations (Diamond 2001). Thus, the bulk molar volume, $V_m(tot)$, is given by:

$$V_{\rm m}({\rm tot}) = \frac{V_{\rm m}({\rm aq}) \cdot V_{\rm m}({\rm car})}{V_{\rm m}({\rm car}) + \varphi({\rm car})(V_{\rm m}({\rm aq}) - V_{\rm m}({\rm car}))}$$
(1)

The mole fraction of the homogeneous carbonic phase is found from the relation:

$$X_{\rm car}^{\rm tot} = \frac{V_{\rm m}({\rm tot}) \cdot V_{\rm m}({\rm aq})}{V_{\rm m}({\rm car}) \cdot V_{\rm m}({\rm aq})},\tag{2}$$

which allows calculation of the bulk composition in terms of the 4 selected components:

$$X_{N_2}^{\text{tot}} = X_{\text{car}}^{\text{tot}} \cdot X_{N_2}^{\text{car}},\tag{3}$$

$$X_{\rm CO_2}^{\rm tot} = X_{\rm car}^{\rm tot} \cdot X_{\rm CO_2}^{\rm car} + (1 - X_{\rm car}^{\rm tot}) X_{\rm CO_2}^{\rm aq}, \tag{4}$$

$$X_{\rm H_2O}^{\rm tot} = \left(1 - X_{\rm car}^{\rm tot}\right) X_{\rm H_2O}^{\rm aq},\tag{5}$$

$$X_{\text{NaCl}}^{\text{tot}} = \left(1 - X_{\text{car}}^{\text{tot}}\right) X_{\text{NaCl}}^{\text{aq}}.$$
(6)

Thermodynamic model of H₂ gains

After analysing a first set of experiments, it was realized that hydrogen diffuses from the piston-cylinder assembly through the quartz into the inclusions, where it reacts with CO_2 as follows (e.g. Morgan et al. 1993):

$$4\mathrm{H}_2 + \mathrm{CO}_2 \to \mathrm{CH}_4 + 2\mathrm{H}_2\mathrm{O}.\tag{7}$$

Therefore, in subsequent experiments Fe_2O_3 powder was added to the sample capsules to buffer $f(H_2)$ at a negligible level:

$$3Fe_2O_{3(s)} + H_{2(g)} \rightarrow 2Fe_3O_{4(s)} + H_2O_{(1)}.$$
 (8)

For the unbuffered experiments, it is still possible to interpret the microthermometric results meaningfully by taking account of reaction (7) in the thermodynamic model. Thus, the change in gas composition and in H₂O concentration induced in the fluid inclusions by reaction (7) causes $T_{\rm h}({\rm car})$ and $T_{\rm m}({\rm cla})$ to become interdependent. The effect of this reaction on the two phase transitions is modelled in Fig. 3c using the same approach as mentioned earlier. With the concentrations of C, O, N₂ and NaCl held constant, the reaction of CO₂ to CH₄ decreases $T_{\rm h}({\rm car})$, while the mode of homogenization remains unchanged. At the same time, the clathrate dissociation curve becomes less steep and it is shifted to distinctly higher temperatures at marginally lower salinity (owing to dilution by the H₂O produced via reaction (7)). Thus, a decrease in $T_{\rm h}({\rm car})$ is coupled to an increase in $T_{\rm m}({\rm cla})$.

Consider, for example, an inclusion initially with no CH₄ and with $T_h(car) = 28.4^{\circ}C$ at the critical point labelled 0 in Fig. 3c. Production of 10 mol% CH₄ in the carbonic phase will depress $T_h(car)$ to 19.7°C at the critical point labelled 10.4. This change in gas composition will also shift the clathrate dissociation temperature from 8.0°C at point 8 to 9.7°C at point 9.

The quantitative model in Fig. 3c allows a reference "H₂ diffusion curve" to be plotted in Fig. 3b, against which the progress of reaction (7) can be monitored. The curve is calculated for a model inclusion initially with $T_{\rm crit}({\rm car}) = 28.4^{\circ}{\rm C}$. Because the mode by which the carbonic phases homogenize does not change as H₂ is added, inclusions that originally underwent bubble-point transitions will shift to the left in Fig. 3b, parallel to the reference curve. Conversely, inclusions that originally underwent dew-point transitions will shift to the right in Fig. 3b as H₂ is added.

Results

The results of the two series of experiments (hydrostaticand deviatoric-stress) are presented separately below. As the inclusions tend to split apart during the experiments and adopt new microstructures (details in Tarantola et al. 2010), the following nomenclature is used to facilitate their description (Fig. 4): "Intact" inclusions are those that survived as single objects during the experiments, regardless of any changes in shape; "Dismembered" inclusions usually generate a "cluster" of inclusions consisting of a recognizable "relict" of the "precursor" surrounded by



Fig. 4 Nomenclature of fluid inclusions modified by deviatoric stresses

smaller "neonate" inclusions that formed entirely during the experiments.

In evaluating the microthermometric results, it is important to bear in mind a shortcoming of our experimental method. Although between 12 and 84 inclusions were analysed by microthermometry prior to each experiment, few of these specific inclusions could be measured again after the experiments. In some cases the samples had broken at crucial points, and in others the new shapes acquired by the relict inclusions were so irregular (including narrow tubes and tails) that reliable measurement of $T_{\rm m}$ (cla) was impossible (Diamond 1992). However, many of the inclusions that had been simply photographed prior to the experiments could be successfully relocated and analysed. Consequently, the microthermometric measurements collected after the experiments largely pertain to different sets of inclusions from those referred to below as the "precursors". This shortcoming reduces the resolution to which changes in $V_{\rm m}$ -X properties can be ascertained, but it will be seen that the changes are in any case large enough to be defined unequivocally.

Hydrostatic scoping experiments

Table 1 lists the experimental conditions and key observations pertaining to the hydrostatic experiments. Table 2 shows the microthermometric results.

Experiment Hyd-4 (unbuffered) was conducted at 800 MPa ($\Delta P = P_{inc} - P_{conf} \sim -200$ MPa) for 16 h. Phase transition temperatures showed shifts (Fig. 5a; Table 3): T_m (cla) had dropped from an average of 8.1°C before to 7.1°C after the experiment, and the dew points of the carbonic phases fell from 27.5 to 26.8°C before to 25.2–22.8°C after the experiment. The carbonic phases were found by Raman to be enriched in CH₄ produced via reaction (7) upon influx of H₂. While this accounts for the decrease in T_{dew} (car), it is inconsistent with the observed reduction in T_m (cla), as the inclusions lie far below the H₂-diffusion band in Fig. 5a. It follows that the aqueous salinity of the inclusions had increased during the experiment.

Experiment Hyd-5 (unbuffered) was conducted at 600 MPa ($\Delta P \sim 0$) for 16 h. Raman analysis showed an increase in the concentration of CH₄ in the carbonic phase. The phase transition temperatures are accordingly more scattered (Fig. 5b), but all the inclusions lie within or very

close to the model H_2 -diffusion band, consistent with reaction (7).

Experiment Hyd-6 (buffered by hematite at base, as shown in Fig. 3d) was performed at 600 MPa ($\Delta P \sim 0$) for 139 h. After the run, the initially reddish-brown buffer powder had turned greenish-black, and Raman analysis showed it to be an intergrowth of hematite and magnetite. The inclusions that were situated near the hematite powder display only slight shifts in phase transition temperatures (diamonds in Fig. 5c) and they contained no new CH₄ according to Raman analysis. In contrast, the inclusions near the upper, unbuffered face of the sample (Fig. 3d) acquired abundant CH₄ and they plot as expected along the H_2 diffusion curve (circles in Fig. 5c). These observations demonstrate the effectiveness of the hematite buffer via reaction (8). However, no trace was found of the H_2O generated by this reaction, for example in the form of new fluid inclusions within the adjacent quartz.

The aforementioned microthermometric evidence supports the conclusion in Tarantola et al. (2010), based on fluid inclusion shapes and microstructures, that differential hydrostatic pressure on the inclusions in our particular samples is negligible under the conditions of experiment Hyd-6. Only a slight reequilibration of the H₂-buffered inclusions is revealed by the shift in $T_{bub}(car)$ to a mean of 27.4°C (compare precursor field with diamonds, Fig. 5c). Accordingly, the subsequent deformation experiments were all performed at 700°C with $P_{conf} = \sigma_3 \approx 600$ MPa, under the assumption that this fulfils the condition ($P_{inc}-P_{conf}$) = 0.

Uniaxial compression experiments

The run conditions and microthermometric data for the four hematite-buffered deformation experiments involving deviatoric stress are listed in Tables 3 and 4. The results are presented in the following order of increasing strain: Def-1 (lowest strain), Def-2 (moderate strain) and Def-3 (medium strain). Experiment Def-6 is presented thereafter because its strain could not be determined, owing to sample disintegration. Following the details of each experiment, the volume fractions of the carbonic phases, $\varphi(car)$, and the calculated bulk V_m -X properties are presented for all the experiments simultaneously. The final section describes the fluid inclusions found in vertical mode-I fractures.

Table 1 Experimental conditions and main observations of hydrostatic experiments at 700°C

Exp.	Basel lab no.	Fe ₂ O ₃ buffer	P (MPa)	Duration (hours)	Comments
Hyd-4	129AT	No	800	16	Formation of CH ₄ ; irregular-shaped inclusions; 3D-haloes
Hyd-5	130AT	No	600	16	Formation of CH ₄ ; euhedral inclusions
Hyd-6	139AT	Yes	600	139	No formation of CH ₄ ; euhedral inclusions



Lowest strain experiment Def-1

After being subjected to $\Delta \sigma = 92$ MPa for 110 h, the strain in this quartz sample was very low (<0.2%). None of the inclusions changed their gas compositions. The intact inclusions show no significant modifications of their micro-thermometric properties (triangles in Fig. 6a) versus the precursor inclusions (shaded field in Fig. 6a), despite their changes in shape (Fig. 5 in Tarantola et al. 2010). The

Fig. 5 Microthermometric results, $T_{\rm m}$ (cla) and $T_{\rm h}$ (car), for fluid inclusions before and after hydrostatic experiments. Axes are explained in Fig. 3b. Dark-shaded fields denote precursor inclusions. Light-shaded bands show modelled effects of H2 influx for inclusions that contain CH₄ according to Raman analysis. a Experiment Hyd-4, conducted at 800 MPa without an H2-buffer. Data (circles) lie far below model H2-diffusion curve, indicating loss of H2O from the inclusions in addition to addition of H₂. b Experiment Hyd-5, conducted at 600 MPa without an H2-buffer. Most data fall in band indicating influx of H₂. Contents of CH₄ in carbonic phases are up to 7 mol%. c Experiment Hyd-6, conducted at 600 MPa with H₂ buffer (Fe₂O₃ powder) situated below sample (setup in Fig. 3d). Data for inclusions near unbuffered top surface of sample (circles) show increases of up to 5 mol% CH₄ in carbonic phase due to H₂ influx, as confirmed by Raman analysis. Inclusions near buffered lower surface of sample showed no CH₄ in Raman analysis. Their data (diamonds) show shifts to left of precursors owing to slight decrease in bulk molar volume. This relationship illustrates the effectiveness of the H₂ buffer

relicts of dismembered inclusions (circles in Fig. 6a) display somewhat scattered microthermometric values, but no consistent trends in $V_{\rm m}({\rm car})$ or salinity are visible. In contrast, the neonate inclusions (black squares in Fig. 6a) have significantly lower $V_{\rm m}({\rm car})$ and higher aqueous salinity, as reflected by the strong shifts to lower $T_{\rm bub}({\rm car})$ and $T_{\rm m}({\rm cla})$ values.

Moderate strain experiment Def-2

After 136 h with $\Delta \sigma = 135$ MPa, the gas compositions of the intact and relict inclusions were unchanged. Raman analyses of monophase neonates revealed the presence of dissolved CO₂ in aqueous solution.

Intact inclusions show no change in $V_{\rm m}({\rm car})$ but a notable increase in aqueous salinity (Fig. 6b). The microthermometric values of the relict inclusions are quite distinct from those of their satellite neonates. Whereas the relicts now display dew-points at temperatures as low as 24.0°C, the neonates undergo bubble-point transitions over a broad range, at temperatures down to 18.6°C, well below those of the precursor inclusions. Clathrate dissociates consistently as part of the Q₂ assemblage, as it did prior to the experiment, but at systematically lower temperatures. Relict inclusions show an increase in aqueous salinity of about 1 mass%, whereas all the neonate inclusions have increased by about 2.5 mass%. As shown by the tie-lines in Fig. 6b, in any given cluster of dismembered inclusions, the relict always displays higher $V_{\rm m}({\rm car})$ and lower aqueous salinity than the surrounding neonates (Table 3).

Medium strain experiment Def-3

After being subjected to $\Delta \sigma = 162$ MPa for 110 h, most of the inclusions had split into clusters of neonates. Raman microspectroscopy showed that the gas compositions remained unchanged, and that the flat, monophase neonates

Exp.	Hom. mode of car ^a	Befor	re experiment		After	experiment	
		N	$T_{\rm m}({\rm cla})^{\rm b}$ (°C)	$T_{\rm h}({\rm car})^{\rm c}$ (°C)	N	$T_{\rm m}({\rm cla})^{\rm b}$ (°C)	$T_{\rm h}({\rm car})^{\rm c}$ (°C)
Hyd-4	Dew point	12	7.9-8.2 (8.1)	26.8-27.5 (27.3)	4	7.0–7.1 (7.1)	22.8-25.2 (23.5)
Hyd-5	Bubble point				5	8.5-8.9 (8.7)	22.7-26.1 (24.6)
	Critical				4	7.9-8.1 (8.0)	26.8-27.1 (27.0)
	Dew point	14	7.8-8.4 (8.1)	28.1-28.4 (28.3)	31	8.0-9.3 (8.6)	23.0-26.9 (25.8)
Hyd-6 buffered ^d	Bubble point	32	7.9-8.3 (8.1)	27.4-28.1 (28.0)	29	7.7-8.3 (8.1)	26.9-27.7 (27.4)
Hyd-6 no bufferr ^d	Bubble point				39	8.3-8.9 (8.5)	24.7-26.9 (26.1)

Table 2 Microthermometric data of hydrostatic experiments

^a Homogenization mode of carbonic phases

^b Dissociation temperature of N₂-CO₂ clathrate in Q₂ phase assemblage. Values listed as range and mean (in brackets)

^c Homogenization temperature of carbonic phases. Values listed as range and mean (in brackets)

^d Buffered inclusions at base of sample near Fe₂O₃ powder, as in Fig. 3d; Inclusions near top of sample are unbuffered

contain CO₂ dissolved in the metastable aqueous liquid. Only a few intact inclusions were found and, except for one case in which salinity had increased, the inclusions had undergone no changes in microthermometric values (Fig. 6c; Table 4). Relict and neonate inclusions show the same microthermometric trends as observed in experiment Def-2, but more pronounced. Here again, within each inclusion cluster, with one exception, $V_{\rm m}({\rm car})$ of the relicts is always higher and aqueous salinity is always lower than in the associated satellite neonates (tie-lines, Fig. 6c). In the relict inclusions, $T_{\rm h}({\rm car})$ is as low as 25.8 (dew-point) or 24.9°C (bubblepoint), and the aqueous salinity shows a wider range than the precursors. Values of $T_{\rm h}({\rm car})$ in the neonate inclusions scatter over a huge range from 11.9 to 28.1°C, the phase transition always being a bubble point. Clathrate always dissociates within the Q₂ assemblage, revealing a systematic increase in aqueous salinity by about 1 mass%.

Experiment Def-6

The general trends in microthermometric values after the experiment (Fig. 6d) are comparable to those in Def-1, Def-2 and Def-3. Intact inclusions do not vary from the precursor properties. All the inclusions show $T_{\rm m}$ (cla) values over a somewhat wider range but still centred on the field of the precursor inclusions. Also the $T_{\rm h}$ (car) values are more widely scattered: the relict inclusions (the latter by as much as 3°C below the average precursor), but salinity remains approximately constant. Neonate inclusions define a clear trend towards lower $T_{\rm bub}$ (car) at constant salinity.

Volume fractions of the carbonic phases, $\varphi(car)$

In all the deformation experiments, the intact inclusions retained the $\varphi(\text{car})$ values of the precursors (~0.18–0.21)

as measured at room temperature. The neonate inclusions are difficult to measure with the spindle stage and so in the cases where flat inclusions were found, their area-fractions were measured and taken as directly equivalent to volume fractions. Overall, the values were found to be very similar or slightly lower than the precursors: $\varphi(car) = 0.15-0.20$. Phase volume fractions were also difficult to measure in the relict inclusions owing to their highly irregular shapes (for which the method of Bakker and Diamond 2006 is not valid). Subjective optical estimates indicate a range of $\varphi(car)$ between 0.20 and 0.35. The range is not random but rather follows a distinct correlation with $V_{\rm m}({\rm car})$ values of the relicts. Inclusions with $V_{\rm m}({\rm car})$ around 80 cm³ mol⁻¹ (Fig. 6) have $\varphi(car) \sim 0.20$, and as $V_m(car)$ increases to around 170 cm³ mol⁻¹ the $\varphi(car)$ values progressively rise to ~ 0.35 .

Changes in fluid inclusion bulk V_m -X properties

The foregoing results from the deformation experiments permit the bulk $V_{\rm m}$ -X properties of the inclusions to be calculated from Eqs. 1 to 6. Figure 7 illustrates the results for selected inclusions from the four deformation experiments reported in Fig. 6. In general, all four experiments show the same trends: (1) The intact inclusions closely retain the $V_{\rm m}$ -X values of the precursor inclusions; (2) Most of the relict inclusions preserve the composition of the precursors, except for some examples in Def-2 and Def-3 in which H₂O contents have slightly decreased and the salt and gas contents have increased. Most notable is the wide spread in molar volumes of the relict inclusions, spanning the range from somewhat lower values than the precursors up to much higher volumes; (3) Compared to the precursors, the neonates all display a clear trend towards lower molar volumes, lower H₂O contents and higher NaCl and gas contents. The neonates mostly have

Fig. 6 Microthermometric results, $T_{\rm m}$ (cla) and $T_{\rm h}$ (car), for fluid inclusions before and after deformation experiments involving deviatoric stresses $(\Delta \sigma)$. Temperature axis correlates non-linearly with molar volume of carbonic phases, $V_{\rm m}({\rm car})$, shown in top scale (explained in Fig. 3a). Vertical line at 28.4°C denotes critical temperature of carbonic phases in precursor inclusions. Vertical $T_{\rm m}$ (cla) axis correlates inversely with aqueous salinity (right scale). Shaded field encompasses all data for precursor inclusions. a Experiment Def-1, conducted with $\Delta \sigma = 92$ MPa for 110 h. **b** Experiment Def-2, conducted with $\Delta \sigma = 135$ MPa for 136 h. c Experiment Def-3, conducted with $\Delta \sigma = 162$ MPa for 110 h. d Experiment Def-6, conducted with $\Delta \sigma = 217$ MPa over short duration, 17 h



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Exp.	Basel lab no.	Fe ₂ O ₃ buffer	Duration ^a (hours)	Strain ^b (%)	Strain rate (s^{-1})	Confining pressure (σ_3) (MPa)	Compressive stress (σ_1) (MPa)	Nominal dev. stress $\Delta \sigma$ (MPa)
Def-1	163AT	Yes	110	< 0.2	$<5 \times 10^{-9}$	613	705	92
Def-2	149AT	Yes	136	0.1-0.3	$<6 \times 10^{-9}$	615	750	135
Def-3	161AT	Yes	110	0.3-0.6	$<2 \times 10^{-8}$	616	778	162
Def-6	152AT	Yes	17	?	?	630	847	217

Table 3 Experimental conditions of deformation experiments at 700°C

^a Time at target P-T conditions of experiment

^b Plastic strain measured from difference in sample height before and after experiment

lower H_2O and higher NaCl contents than their adjacent relict inclusions (indicated by tie-lines in Fig. 7).

Fluid inclusions in vertical mode-I fractures

Typically, the vertical loading fractures are "dry", but some are partly healed and contain myriads of tiny fluid inclusions. All fractures that host such inclusions appear to have cut through a pre-existing inclusion in the sample.

In sample Def-1, the fracture-hosted neonate inclusions are large enough to permit identification of two or three fluid phases at room temperature, and Raman analysis confirms the presence of a carbonic phase. The volume fraction of the carbonic phase varies widely between 0.4 and 0.8 in sample Def-1 and can exceed 0.9 in sample Def-2. The microthermometric results (Table 4) show that the aqueous phase has essentially the same salinity as the precursor inclusions (average $T_m(cla) = 3.9^{\circ}C$; clathrate dissociates within the Q₂ or ClaL_{aq}L_{car} assemblages). Values of $T_{bub}(car)$ vary over an extremely wide range, from -9.1 to 21.9°C, much wider than the spreads of neonate inclusions in Fig. 6.

The range in volume fractions and the microthermometric features of the inclusions are diagnostic of heterogeneous entrapment (e.g. Diamond 2003). The cause and timing of the phase separation in the mode-I cracks is explained in the Electronic Supplement.

Discussion

The results presented earlier are internally consistent and reproducible. The most obvious differences between the experiments are due to the different durations and magnitudes of applied deviatoric stress. The results thus provide a basis to generalize the modifications of fluid inclusion composition and density induced by small degrees of uniaxial shortening. The purpose of the following discussion is to elucidate the mechanisms of these changes with a view to formulating criteria to interpret naturally deformed inclusions. Changes in fluid inclusion composition

In general, the intact inclusions faithfully retain the bulk composition of the precursors. Only in experiment Def-2 is a slight systematic increase in salinity visible. Also the relict inclusions mostly preserve the composition of the precursors (again Def-2 shows an increase in salinity). In contrast, the neonate inclusions show quite dramatic, systematic shifts to lower H₂O contents and higher gas and salt contents. Moreover, the higher the imposed deviatoric stress and the longer the duration of the experiment, the greater are these changes compared to the precursors. This correlation supports the idea that the application of deviatoric stress over extended periods of time is the driving force for the compositional changes. In most cases, the neonates have lower H₂O and higher gas and salt contents than the relict inclusions that they surround. These systematic features raise the questions of how and why the compositions of the neonates change, while the intact and relict inclusions remain unchanged.

The first task is to identify which of the chemical components is mobile. This becomes evident by replotting the results in Fig. 7 into a ternary diagram (Fig. 8). The shift of the neonates from the precursor fields in all four experiments can be well explained by loss of H_2O (vector marked with arrow in Fig. 8a), without the need to invoke mobility of the gas or salt components.

This deduction is consistent with previous studies. To our knowledge, no published experimental work has demonstrated that N_2 , CO_2 or NaCl may be lost from fluid inclusions in quartz via diffusion. In contrast, several high-*T* experimental studies (Bakker and Jansen 1990, 1991; Hall and Sterner 1993; Sterner et al. 1995) have concluded that molecular H₂O may diffuse into or out of fluid inclusions in quartz in response to gradients in the fugacity of H₂O. Whereas the physicochemical interaction between small, polar H₂O molecules and the various dislocations and fracture surfaces in quartz is known to facilitate its mobility via pipe- or microcrack diffusion (e.g. McLaren et al. 1983, 1989; Kronenberg et al. 1986; Cordier and

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Exp.	Homog. mode of car ^a	Bef	fore experiment		Afte	rr (intact inclusi	(suo)	Afte	sr (relict inclusi	(suo	After	· (neonate inclu	sions)
		и	$T_{\rm m}({\rm cla})^{\rm b}~(^{\circ}{\rm C})$	$T_{\rm h}({\rm car})^{\rm c}~(^{\circ}{ m C})$	и	$T_{\rm m}({\rm cla})^{\rm b}~(^{\circ}{\rm C})$	$T_{\rm h}({\rm car})^{\rm c}~(^{\circ}{\rm C})$	и	$T_{\rm m}({\rm cla})^{\rm b}~(^{\circ}{\rm C})$	$T_{\rm h}({\rm car})^{\rm c}~(^{\circ}{\rm C})$, u	$T_{\rm m}({\rm cla})^{\rm b}~(^{\circ}{ m C})$	$T_{\rm h}({\rm car})^{\rm c}~(^{\circ}{\rm C})$
Def-1	Bubble point	21	7.8-8.1 (8.0)	28.2-8.5 (28.3)	23	7.8-8.9 (8.4)	26.9-28.2 (27.6)	5	7.8-8.6 (8.2)	27.2-28.3 (27.6)	22 (6.9-8.2 (7.8)	17.8-28.1 (26.0)
	Critical	2	8.2-9.1 (8.7)	28.3-28.4 (28.4)									
	Dew point	30	7.6–9.0 (8.2)	27.3-28.5 (28.1)				7	8.5-9.2 (8.9)	27.9-28.2 (28.1)			
	Bubble point ^d		n.a	n.a	20	1.5-5.0 (3.9)	-9.1 to 21.9 (7.4)						
Def-2	Bubble point	31	7.8-8.4 (8.1)	24.2-28.4 (27.4)	6	7.1–7.7 (7.4)	23.9-26.2 (25.1)				8	6.2-6.8 (6.6)	18.6-25.0 (22.8)
	Dew point	Ζ	7.9-8.2 (8.1)	28.2-28.4 (28.3)	-	7.4	27.9	З	7.5-8.1 (7.8)	24.0-26.9 (25.3)			
Def-3	Bubble point	13	7.9–9.1 (8.2)	25.3-28.3 (27.9)	4	6.8-8.5 (7.9)	26.2-27.6 (26.8)	8	7.6–9.0 (8.1)	24.9-27.7 (26.6)	. 89	7.2-8.6 (7.7)	11.9-28.1 (21.4)
	Critical	5	8.1-8.5 (8.3)	28.1-28.5 (28.2)									
	Dew point	37	7.9-8.5 (8.2)	28.1-28.4 (28.2)				4	7.6–9.3 (8.4)	25.8-28.0 (27.2)			
Def-6	Bubble point	29	7.8-8.8 (8.2)	27.7-28.3 (28.2)	12	7.9–8.7 (8.2)	28.2-28.4 (28.4)	4	8.2-8.8 (8.4)	26.9-27.9 (27.4)	25	7.7-8.4 (8.0)	22.7-27.3 (26.0)
	Critical	0	8.2-8.3 (8.3)	28.3									
	Dew point	9	8.3-8.4 (8.4)	28.2-28.4 (28.3)	4	7.5-8.1 (7.7)	28.0-28.3 (28.2)	5	8.1-8.9 (8.4)	26.0-28.0 (27.4)			
^a Homoe	enization mode of carbo	onic r	phases										

 Table 4
 Microthermometric data of deformation experiments involving deviatoric stresses

^b Dissociation temperature of N₂-CO₂ clathrate in Q₂ phase assemblage. Values listed as range and mean (in brackets)

^c Homogenization temperature of carbonic phases. Values listed as range and mean (in brackets)

^d Fluid inclusions in vertical mode-I fractures



Fig. 7 Calculated bulk compositions, $X(CO_2 + N_2)$, $X(H_2O)$ and X(NaCl) and molar volumes, $V_m(tot)$, of representative fluid inclusions in deformation experiments involving deviatoric stresses. Values of $\varphi(car) = 0.25-0.35$ have been used for the relict inclusions, and $\varphi(car) = 0.20$ has been used for the precursor, intact

Doukhan 1989; Bakker and Jansen 1990, 1991, 1994; Kronenberg 1994), N_2 , CO_2 and NaCl molecules appear to be immobile in quartz under conditions of overall electrical neutrality. Accordingly, the changes in composition of the neonate inclusions may be attributed solely to loss or gain of H₂O.

Loss of H₂O via diffusion requires a significant gradient in H₂O fugacity between the neonates and a specific local sink. Figure 9 shows $f(H_2O)$ calculated using the EoS of Bowers and Helgeson (1985) for the same Def-2 inclusions that are displayed in Fig. 7d–f. The bulk molar volumes, V_m (tot), have been adjusted as described in the next paragraph (Eq. 9). It can be seen that the low molar volume (high density) of the neonates implies much higher $f(H_2O)$ values at 700°C than those of the intact inclusions ($\Delta f(H_2O) \sim 80$ –200 MPa) and of the relict inclusions ($\Delta f(H_2O) < 400$ MPa), despite the lower H₂O contents of the neonates. As the neonates have already lost

and neonate inclusions. The effect of the uncertainty in $\varphi(car)$ between 0.15 and 0.20 for some neonates is illustrated by one example *error bar*. *Tie-lines* join data for inclusions that belong to the same cluster of relict and neonate inclusions

considerable H_2O compared to their precursors, the high $f(H_2O)$ values in Fig. 9 must be residuals of even higher fugacities that presumably prevailed during the early stages of the experiments.

Now with clear evidence for high $f(H_2O)$ in the neonates, the potential sinks for H₂O need to be identified. Many of the adjacent relict inclusions and the nearby intact inclusions evidently had lower $f(H_2O)$ at 700°C, therefore these two types of inclusions may have acted as sinks for the H₂O lost from the neonates. However, because both inclusion types are vastly larger than the neonates (typically by a volume factor of 10^2-10^3), any gain of H₂O may be undetectable. In fact, none of the intact or relict inclusions show measurable gains of H₂O (Fig. 7).

A second sink for H_2O could be the quartz host-crystal itself, but current understanding of H_2O in quartz defects (e.g. Cordier et al. 1994; Cordier and Doukhan 1995) suggests that the amount of H_2O that can be stored is



Fig. 8 Ternary CO₂–H₂O–NaCl compositions (mole fractions) of fluid inclusions after deformation experiments involving deviatoric stresses. *Black area* in *inset triangle* shows region of expanded diagrams. Thin reference lines emanate from H₂O apex, defining directions of possible loss and gain of H₂O. Positions of neonate inclusions indicate loss of H₂O relative to precursor inclusions. **a** Def-1, **b** Def-2, **c** Def-3 and **d** Def-6

minute—far lower than that needed to account for the observed changes in the neonate inclusions.

A third sink for H_2O could be the exterior of the quartz samples, which are tightly encased in a gold jacket during the experiments. Although the H_2 -buffer reaction (8) generates water on the surfaces of the two alumina pistons inside the gold jacket (Fig. 3d), none of this water seems to have reached the surfaces of the quartz samples (the quartz is separated from the alumina by unwelded gold discs; Fig. 3d). Most of the mode-I fractures that formed in the



Fig. 9 Fugacity of H₂O at 700°C in two clusters of dismembered inclusions from deformation experiment Def-2 ($\Delta \sigma = 135$ MPa, moderate strain). See text for "adjustment" of bulk molar volume, $V_{\rm m}(\text{tot})$. Calculations made with EoS of Bowers and Helgeson (1995) with $\varphi(\text{car}) = 0.20$ for neonate, precursor and intact inclusions, and $\varphi(\text{car}) = 0.30$ and 0.35 for relict inclusions R_5 and R_2 , respectively. *Numbers* with an *asterisk* are unadjusted $V_{\rm m}(\text{tot})$ values (cm³ mol⁻¹) at room temperature. *Tie-lines* join dismembered fluid inclusions within the same cluster. The neonates have much higher $f(\text{H}_2\text{O})$ than the precursor, intact and relict inclusions, and the relicts have the lowest values of all

quartz during heating and pressurization towards the run conditions are open and "dry". Had water wetted the quartz surfaces, the fractures would certainly have healed and trapped fluid inclusions (this process is the basis of the well established method of synthesizing fluid inclusions, e.g. Bodnar and Sterner 1987). Where inclusions did form in the fractures, their compositions show that they derive from rupturing of pre-existing CO2-H2O-NaCl inclusions ("precursors"). The fact that the fracture-hosted inclusions were trapped in a heterogeneous state indicates much lower effective fluid pressures than $P_{\rm conf}$ (discussed in detail in the Electronic Supplement). As well as demonstrating the absence of an external source of water, these same points of evidence also demonstrate that no appreciable H₂O reached the sample surfaces from within the quartz. Therefore, despite the quartz surfaces being a potential sink for H_2O , and despite the nominally huge gradient in $f(H_2O)$ between the fluid inclusions and the unwetted quartz surfaces, it seems that diffusive transport was too slow relative to the duration of the experiments to permit loss of H₂O to the sample exterior.

In conclusion, the most probable explanation is that, given the high fugacity gradient in Fig. 9, H_2O diffused from the neonates into the adjacent relict inclusions at the length-scale of individual inclusion clusters. It is reasonable to assume that the H_2O gained by the relict inclusions is undetectable. The dominant pathways for transport were most likely the abundant microcracks and dislocations that link the two inclusion types within individual clusters.

Changes in molar volume (density) of the fluid inclusions

Changes in molar volume of neonate inclusions

In the foregoing, it is concluded that the difference in molar volume between the neonate and relict inclusions generates the gradient in $f(H_2O)$ which drives exchange of H_2O . The next question to be addressed is the origin of the differences in molar volume.

Based on the results of previous experimental studies involving hydrostatic underpressures ($P_{inc} < P_{conf}$) (e.g. Sterner and Bodnar 1989), we expect the precursor inclusions to become denser as they adapt to a new mechanical (P-V) equilibrium commensurate with the deviatoric stress state of the host quartz. However, the end-point of the new P-V equilibrium is not obvious a priori. For instance, the fluid may adopt a molar volume compatible with the mean stress at 700°C, or with some other stress value. The bulk $V_{\rm m}$ -X properties calculated for the intact, relict and neonate inclusions (e.g. Fig. 7) all correspond to defined fluid pressures at 700°C. These pressures, $P_{\rm inc}$, can be reconstructed using an appropriate EoS by assuming quasi-isochoric behaviour of the inclusions from $T_{\rm h}({\rm car})$ up to 700°C. Thus, comparison of the $P_{\rm inc}$ values with the σ_1 and σ_3 stresses imposed during the experiments should in principle reveal the end-point of reequilibration of the inclusions.

Calculations of P_{inc} at 700°C were duly made for precursor, intact, relict and neonate inclusions in experiments Def-1, Def-2, Def-3 and Def-6. Nitrogen was lumped with CO₂ to calculate ternary CO₂–H₂O–NaCl isochores with the Bowers and Helgeson (1985) and Duan et al. (1995) EoS. The modification of the Bowers and Helgeson (1985) EoS by Bakker (1999) was also used to take explicit account of N₂, but the pressures were not significantly different for the very small bulk $X(N_2)$ values involved. Corrections were made for the expansivity and compressibility of α -quartz between P_{inc} at 700°C and $T_h(car)$ at $P_h(car)$ using the EoS of Hosieni et al. (1985).

The results of these calculations pose a conflict in our experimental method. The value of $\sigma_3 = 600$ MPa is compatible with the calculated range of pressures extrapolated for the complete set of inclusions reported by Diamond (1990) at the Brusson locality (Fig. 1). However, the $P_{\rm inc}$ values calculated for the specific set of precursor inclusions used in the experiments lie between 750 and 820 MPa, some 150–220 MPa higher than σ_3 (or 100–170 MPa applying the maximum uncertainty in σ_3). Nevertheless, the hydrostatic scoping experiments Hyd-5 and Hyd-6 both showed microstructural equilibration with $P_{\rm conf} = 600$ MPa (Fig. 4 in Tarantola et al. 2010), and it

was this evidence that lead us to assume that $\Delta P = 0$ at 600 MPa and 700°C. We have carefully reexamined our experimental method and the approach and assumptions made in calculating $P_{\rm inc}$ from the isochores, but we have been unable to resolve this conflict. As our prime aim is to evaluate the effects of deviatoric stress, $\Delta \sigma$, in general, rather than the effects of absolute stress, this inconsistency is not crucial to our study. There seems to be more leeway in the EoS extrapolation of the isochores to high temperature than in the measurement of maximum pressure in the Griggs piston-cylinder apparatus. Consequently, we have chosen to accept that $\sigma_3 = 600 \pm 50$ MPa truly represents the experiments.

To evaluate the internal pressures of the modified inclusions, we have adjusted the calculated isochores such that the mean precursor isochore intersects the corresponding confining pressure (σ_3 ; Table 2) at 700°C. The fan of precursor isochores derived from this mean is displayed in light shading in Fig. 10. To preserve the relative difference in pressure between the neonate and precursor inclusions, the EoS pressures calculated for the neonates at 700°C were adjusted as follows:

$$P_{\text{neonate}}(\text{adjusted}) = \frac{P_{\text{neonate}}(\text{EoS})}{P_{\text{precursor}}(\text{EoS})} \cdot \sigma_3[\text{MPa}]$$
(9)

The resulting shifts in internal pressures are indicated in Fig. 10. Points labelled 1 show P_{inc} of the densest neonate inclusions based on EoS calculations and points labelled 2 show the adjusted values. The adjusted pressures were used to construct the fans of isochores of the neonate inclusions, as shown in dark shading in Fig. 10.

In experiments Def-1 and Def-3 (Fig. 10a, c), the isochores of the neonates span the entire range of fluid pressures between σ_1 and σ_3 . In the other long-duration experiment, Def-2 (Fig. 10b), the densest neonates approach σ_1 quite closely, whereas in the short-duration experiment Def-6 (Fig. 10c), the highest pressure in a neonate inclusion lies at about a third of the interval to σ_1 . This suggests that the neonates do not initially form with high densities, but that they increase their densities with time. It also seems likely that the end-point of this density evolution is equilibrium with σ_1 , rather than with the mean stress ($\sigma_1 + 2\sigma_3$)/3.

Changes in molar volume of relict inclusions

The relict inclusions generally have not changed composition with respect to the precursors, but many of them have increased their molar volumes. It follows that these relicts have *expanded* during the experiments. This conclusion agrees with the observed correlation between $\varphi(car)$ and $V_m(car)$. Calculations of volume expansion of a model precursor (Fig. 11; details of calculation in

Fig. 10 Comparison of adjusted isochores of precursor inclusions (light shading) and neonate inclusions (dark shading) in deformation experiments involving deviatoric stresses. Open squares show magnitude of $\sigma_3 \pm 50$ MPa at 700°C; filled squares show magnitude of $\sigma_1 \pm 10\%$ relative to σ_3 at 700°C. Points 1 and 2 show the adjustment of maximum P_{neonate} at 700°C, according to Eq. 9 (see text). The adjusted pressures of neonates span the range between σ_3 and σ_1 , the highest values being approximately equal to the absolute value of σ_1 . a Experiment Def-1. **b** Experiment Def-2. c Experiment Def-3. d Experiment Def-6



"Appendix 2") predict correlations between $\varphi(\text{car})$, $V_{\rm m}(\text{car})$ and $V_{\rm m}(\text{tot})$ that match the observations quite well. The maximum observed $\varphi(\text{car})$ and $V_{\rm m}(\text{car})$ values of around 0.35 and 170 cm³ mol⁻¹, respectively, suggest that some inclusions expanded by as much as 20% (Fig. 11b). Presumably the microcracking observed around the relicts permitted this expansion to occur. Nevertheless, any expansion at all is remarkable considering that the samples were subjected to positive deviatoric stresses ($\sigma_1 > P_{\rm inc}$). The high molar volumes of the relicts correspond to internal pressures as low as 400 MPa during the experiments, i.e. some 200 MPa *below* the value of σ_3 . Thus, contrary to expectation, the final state reached by the relict inclusions during the deformation process lies further from *P*–*V* equilibrium than the initial (precursor) state.

There is no evidence from the experiments that the pressure drop associated with volume expansion was sufficient to cause phase exsolution (also termed heterogenization or unmixing or boiling) in the relict inclusions (in contrast to the vertical mode-I fractures, as described in the Electronic Supplement). In nature, however, it is conceivable that phase exsolution could take place if the quartz



Fig. 11 Test of hypothesis that relict inclusions have expanded during deformation experiments involving deviatoric stresses. *Curves* show predicted relationships for the range of bulk compositions observed in relict inclusions. Calculations made according to "Appendix 2". **a** Relationships between measurable parameters: molar volume of carbonic phase, $V_m(car)$, and volume fraction of carbonic phase, φ (car). **b** Predicted relationship between $V_m(car)$ and amount of expansion of relict inclusions (%). Data points with *error bars* are examples of observations from relicts in experiments Def-2 and Def-3. Agreement between predictions and observations is consistent with expansion hypothesis

deformation occurs at lower lithostatic pressures (shallower levels in the crust), closer to the miscibility boundary of the specific CO_2 –H₂O–NaCl fluid. This process may be relevant to the occurrence of almost pure carbonic fluid inclusions in shear-zone hosted quartz veins (e.g. Schwartz et al. 1992; Klemd et al. 1997; Schmidt Mumm et al. 1997).

Mechanisms of fluid inclusion modifications

Our understanding of the mechanisms that result in the observed modifications of the inclusion shapes and V_m -X

properties is still incomplete. In the following, we summarize our conclusions so far in the form of a conceptual model.

Application of deviatoric stress causes stress concentration radially normal to σ_1 at acute angles in the walls of the precursor fluid inclusions. This generates brittle microcracks that propagate into the surrounding quartz by parting the crystallographic cleavage planes oriented subparallel to the σ_3 plane. Crack propagation stops when the elastic component of the strain is relaxed by a net volume increase (by up to 20%) of the inclusion. The fluid injected into the cracks promotes quartz healing by dissolution at sites of high interfacial energy (e.g. sharp irregularities) and by adjacent reprecipitation. Branches of fluid are first isolated in this crack-healing process and with time myriads of tiny, subequant to equant neonate inclusions become isolated as the branches neck down, locally minimizing interfacial energy. The fluid density of the neonates is initially similar to that of the expanded relict inclusions, but with time the density of the neonates increases towards *P*–*V* equilibrium with the maximum applied stress, σ_1 . The compression of the fluid that elevates the density seems to be facilitated by crystal-plastic dislocation glide of the quartz in the vicinity of the healed cracks. Presumably this plasticity is stimulated by H₂O-weakening of the dislocation-rich "wetted" quartz adjacent to the cracks (Fitz Gerald et al. 1991). The compression of the neonates significantly raises their $f(H_2O)$, which drives diffusion of H₂O out into the surroundings. The salt and gas contents of the neonates thereby increase passively with time. Owing to the low internal pressures of the relict inclusions, they may act as sinks for the diffusing H₂O. The plastic deformation that nucleates around the neonate inclusions progressively spreads out into the surrounding lattice via basal and prism glide systems, leading to bands of discrete c-axis misorientation and undulatory optical extinction in inclusion-bearing domains of the sample (cf. Fitz Gerald et al. 1991).

Conclusions

This experimental study has provided a coherent picture of fluid inclusion modifications induced by low crystal-plastic strain at high temperature. The companion paper (Tarantola et al. 2010) provides arguments that the experimental results are transferable to nature. Thus, several guidelines can be established regarding the information obtainable from fluid inclusions in natural, low-strain, sheared quartz samples. These guidelines are formulated in terms of the nomenclature for deformed inclusions given in Fig. 4.

Very small shear strains impart irregular shapes to the intact inclusions, but their compositions and densities

remain unperturbed. Higher strains may cause a slight increase in salt and gas contents owing to diffusive loss of H_2O adjacent to relict inclusions. Microthermometric analyses of the intact inclusions in low-strain samples should therefore allow the pre-deformation V_m -X properties of the precursors to be reconstructed to a close approximation.

Large irregularly shaped relicts of the precursor inclusions sit at the centre of the clusters of neonates. According to our current understanding, these relicts faithfully record the composition of the precursors but in addition to shape changes they may have undergone significant volume expansion (despite the compressive deviatoric stress). In general, their densities (and hence their isochores) bear no relation to any pressure- or stress-state experienced by the rock sample during the deformation event. Therefore, their densities cannot be used to gain meaningful information on the free paleofluids or on the magnitudes of the principal shear stresses.

The neonate inclusions typically have lost significant H_2O via diffusion and hence their salt and gas contents are passively increased in comparison to the precursor inclusions. As such, the chemical compositions of the neonates are artefacts of the deformation process and they do not record the properties of any free hydrothermal or metamorphic fluid that initially existed at the grain boundaries of the quartz sample. In contrast, the densities of the neonates have the potential to yield useful information. They span a very wide range, the highest of which may be equal to the density in P-V equilibrium with the principal compressive stress, σ_1 , during shear deformation. Therefore, reconstruction of the isochores of the densest neonates may help to estimate the absolute stress of σ_1 if the temperature of deformation is known.

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Appendix 1: Measurement of volume fraction of carbonic phase in a precursor inclusion

Following the method of Bakker and Diamond (2006), the volume fraction of the carbonic phases in an example precursor inclusion can be determined from the two data plots in Fig. 12. Figure 12a displays triplicate measurements of the total projected area of the inclusion as a function of rotation angle of the spindle stage. Figure 12b displays triplicate measurements of the area fraction of the

carbonic phase as a function of rotation angle of the spindle stage. The maximum in the data array in Fig. 12a lies at the rotation angle 12.5° (marked by an arrow). This angle, when intersected with the area-fraction curve in Fig. 12b, indicates the area fraction (0.193) which most closely approximates the numerical value of the volume fraction. The method carries an uncertainty of $\pm 4\%$ relative. Thus, the result for this example is $\varphi(\text{car}) = 0.193 \pm 0.008$.



Fig. 12 Estimation of the volume fraction of the carbonic phase, $\varphi(car)$, at room temperature in an example inclusion using the method of Bakker and Diamond (2006). **a** Projected area of the inclusion versus angle of rotation in the spindle stage. **b** Area fraction of the carbonic phase versus angle of rotation in the spindle stage. See text for explanation

Appendix 2: Calculation of relationship between $V_{\rm m}({\rm car})$ and $\varphi({\rm car})$ for an expanding fluid inclusion

To test the hypothesis that the relict fluid inclusions have expanded their volumes at constant composition, the relationship between the observable parameters $V_{\rm m}(\text{car})$ and $\varphi(\text{car})$ can be calculated as a function of volume expansion. For a fluid inclusion of fixed composition, the mole fraction of carbonic phase can be obtained by rearranging Eq. 5:

$$X_{\rm car}^{\rm tot} = 1 - \frac{X_{\rm H_2O}^{\rm tot}}{X_{\rm H_2O}^{\rm aq}}.$$
 (10)

This value can be inserted into a rearrangement of Eq. 2 to yield the molar volume of the carbonic phase:

$$V_{\rm m}({\rm car}) = \frac{V_{\rm m}({\rm tot}) - V_{\rm m}({\rm aq})}{X_{\rm car}^{\rm tot}} + V_{\rm m}({\rm aq}), \tag{11}$$

where for simplification $V_{\rm m}(\rm aq)$ is held constant. Input values of $V_{\rm m}(\rm tot)$ are generated by multiplying the initial (measured) $V_{\rm m}(\rm tot)$ by an arbitrary factor to simulate expansion of the inclusion volume. For example, 10% expansion increases an initial $V_{\rm m}(\rm tot)$ of 22 cm³ mol⁻¹ to a final $V_{\rm m}(\rm tot)$ of 24.2 cm³ mol⁻¹. Solution of Eq. 10 for a range of $V_{\rm m}(\rm tot)$ yields the model relationship between $V_{\rm m}(\rm aq)$ and the hypothesized volume expansion (curve in Fig. 11b).

Inserting the appropriate constant $V_{\rm m}(aq)$ and the $V_{\rm m}(car)$ value calculated from Eq. 10 into the following expression (Eq. 5 in Diamond 2001):

$$\varphi(\operatorname{car}) = \frac{\left(\frac{V_{\mathrm{m}}(\mathrm{aq})}{V_{\mathrm{m}}(\mathrm{tot})} - 1\right)}{\left(\frac{V_{\mathrm{m}}(\mathrm{aq})}{V_{\mathrm{m}}(\mathrm{car})} - 1\right)},\tag{12}$$

yields the model relationship between $V_{\rm m}({\rm car})$ and $\varphi({\rm car})$ (curve in Fig. 11a), which can be compared with the observations.

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