Hydrogen diffusion mechanisms in quartz: insights from H–Li, ²H–H and ²H–H–Li exchange experiments

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Abstract

The diffusivity and diffusion mechanisms of hydrogen together with with deuterium and lithium, parallel to the c axis of quartz, were investigated experimentally at 800°C, 0.1 GPa with the activity of H₂O or ²H₂O ≈ 1 [²H is used throughout this work to describe deuterium rather than D, to avoid confusion with the diffusion coefficient, D]. The pH was set using mixtures of H₂O (or ²H₂O) and HCl. Three types of experiment were conducted: (1) H-in/Li-out; (2) ²H-in/H-out; and (3) ²H-in/H + Li out, using three different natural quartz crystals as starting materials. Profiles of H, ²H and Li were measured using Fourier-transform infrared (FTIR) spectroscopy and laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS). H, ²H and Li are charge-compensated by Al³⁺ replacing Si⁴⁺, or by excess O²⁻. The total atomic concentration of monovalent cations appears to remain constant over the duration of the experiments. The resulting diffusion profiles are different for the three experimental designs and three starting materials, and some show complex shapes inconsistent with simple diffusion. A multi-site diffusion–reaction model is developed, with the theory based on previous models that have been derived mainly on the basis of conductivity measurements. In these models, the monovalent cations move away from their charge-balancing ion then diffuse rapidly to another site. The mobility of the monovalent cations is described by both a diffusion coefficient and an equilibrium constant that enables dissociation of the immobile charge-balanced defects. This model can describe complex step-shaped profiles formed in H-in/Li-out experiments, profiles with local maxima (‘humped’ profiles) in ²H-in/H + Li out experiments, and error function-shaped profiles in ²H-in/H-out and previously published Li-in/H-out experiments. Our data provide support for models previously proposed for quartz. Studies of the lengths and forms of diffusion profiles from such experiments provide a useful complement to assertions from conductivity experiments.

Keywords: quartz, diffusion, Fourier transform infrared spectroscopy, point defects

Introduction

Understanding how monovalent cations substitute into quartz, as well as how rapidly, and how, they move, is important for considerations including electrical conductivity (Verhoogen, 1952; Hughes, 1975; Jain and Nowick, 1982), diffusion chronometry or more generally quartz compositional heterogeneity in volcanic systems (Biró et al., 2016; Tollan et al., 2019), the fidelity of quartz hosted melt and fluid inclusions (Rottier et al., 2017; Myers et al., 2019), quartz rheology (Griggs and Blacic, 1965; Mackwell and Paterson, 1985) and purification or controlled alteration of quartz by electro-diffusion ‘sweeping’ (King, 1959; Brown et al., 1980).

For these reasons, much experimental work has concentrated on obtaining independent determinations of the diffusion coefficients of the monovalent cations, mainly focussed on H and its isotopes (Kats, 1962; Shaffer et al., 1974; Kronenberg et al., 1986; Rovetta et al., 1989; Bachheimer, 1998; Jollands et al., 2020a) though also including Li, Na and K (Verhoogen, 1952; Rybach and Laves, 1967; Frischat, 1970; Shaffer et al., 1974; Sarthävea et al., 2004; Rottier et al., 2017), as well as their substitution mechanisms (Staats and Kopp, 1974; Sibley et al., 1979; Halliburton et al., 1981; Hosaka and Taki, 1981; Aines and Rossman, 1984; Paterson, 1986; Stalder and Konzett, 2012; Baron et al., 2015; Frigo et al., 2016; Müller and Koch-Müller, 2018; Potrafke et al., 2019; Jollands et al., 2020b). The fact that the system remains the target of active study >130 years after the first published conductivity measurements in quartz (Tegetmeier and Warburg, 1887; Curie, 1889), which were more-or-less contemporaneous with the first observations of the presence of O–H groups within quartz (Merritt, 1895; Königsmberger, 1897), is testament to its importance. For more information regarding H incorporation and diffusion in quartz, as well as various physical, chemical and geological implications, see the recent review by Stalder (2021).

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Notwithstanding the considerable amount of research, absent from the literature are descriptions of experiments wherein H is diffused into quartz at elevated pressure and temperature, and the resulting profiles of H content versus distance are measured using Fourier-transform infrared (FTIR) spectroscopy. This technique has now become widely accepted as a tool for considering H diffusion in other nominally anhydrous minerals (e.g. Demouchy and Mackwell, 2006). Not only can diffusion coefficients be determined using this method, but the contributions of individual H-bearing defects to total H can be resolved, and the specific shapes of profiles in concentration vs. distance space can be used to elucidate diffusion mechanisms.

The diffusion of H in quartz has been proposed to occur by the movement of H along some fast pathway, followed by immobilisation of that H when it moves into association with stationary Al (Hughes, 1975; Jain and Nowick, 1982; Kronenberg and Kirby, 1987). This is discussed in more detail below, though, importantly, the corollary of invoking such a diffusion mechanism would be diffusion profile forms that do not necessarily correspond to error-function shapes (e.g. Dohmen et al., 2010; Bloch et al., 2020). However, in our previous study (Jollands et al., 2020a), where H was diffused out of quartz, coupled with in-diffusion of Li or Na, we consistently observed profiles that could be fitted to error-function forms. This discrepancy between prediction and observation forms the motivation for the present study, which describes experiments that can be considered broadly as the reverse of those presented by Jollands et al. (2020a) i.e. wherein H, or \(^2\)H\(^+\), is diffused into quartz crystals rather than being diffused out.

Herein, experiments are described wherein different natural quartz crystals containing trace concentrations of H and Li were annealed at 800°C and 0.1 GPa together with H\(_2\)O or \(^2\)H\(_2\)O at controlled pH. These experiments show H or \(^2\)H gain into, and Li loss out of, the quartz crystals. Regardless of the relatively simple experimental design, the diffusion profiles of H (and \(^2\)H, Li) are complex, and rarely conform to error-function shapes. These results can be explained by invoking the aforementioned model (Hughes, 1975; Jain and Nowick, 1982; Kronenberg and Kirby, 1987), which are also capable of explaining qualitatively why simple error-function forms were observed by Jollands et al. (2020a).

**Methods**

**Starting materials**

Cubes cut from three of the crystals used and characterised in a previous study considering H diffusion in quartz (Jollands et al., 2020a) were again used for this study. These were two sections of a single quartz slab cut from two Tibetan crystals (TIB2 and TIB6), and a crystal of Brazilian quartz (BRA3). In the

\[^2\]H is used throughout to describe deuterium rather than D, to avoid confusion with the diffusion coefficient, D.
previous study, TIB2 is denoted Q2, and TIB6 is Q6, both cut from crystal ‘TIB’. BRA3 was denoted Q5. These were chosen to represent a variety of H and Li contents. TIB2 has the highest H + Li, TIB6 the lowest and BRA3 is intermediate.

The only notable trace elements in these materials, as measured in Jollands et al. (2020a), are Al, Li, B, Ti and H. Generally Na and K were below detection limits. It is not possible to give initial concentrations in the exact crystals used for experiments, though concentration ranges for detected trace elements in the starting materials overall (wt. ppm) were: TIB2: 2–8; TIB6 the lowest and BRA3 and TIB6 is intermediate.

Using the Thomas et al. (2019) approach, TIB2 is denoted Q2, and TIB6 is Q6, both cut from crystal before the experiment. Such experiments must be undertaken in two steps, firstly to equilibrate (or metastabili- tate) exchange experiment, the only exchange should be between H and2H. Thus, Li or other monovalent cations should be eliminated by exchange with2H (e.g. Kurka et al., 1992). Values obtained were extrapolated using the density correlation approach (Eugster and Baumgartner, 1987; Manning 1994; Dolejs and Manning, 2010). The program Soluble (Roselle and Baumgartner, 1995) was used to calculate pH, using a Debye-Hückel activity model. Values of the pH in the text herein, refer to pH at room temperature (i.e. 1, 3, 5 and 7).

The open ends of the capsules were then flat crimped, welded closed, weighed, and left for >12 hr at ~110°C, then reweighed to verify no mass loss. Charges were loaded into cold-seal pressure vessels which were pressurised (0.1 GPa, N2 pressure medium), and pre-heated to 800°C, with temperatures monitored using type-K thermocouples (further information is provided in the Supplementary Appendix, see below for details). The capsule was then introduced into the hot spot and left for 1 hr using a magnetic system (e.g. Matthews et al., 2003). At the end of the runs, the samples were removed from the hot spot and the bombs were cooled down by blowing with compressed air – thus cooling from run temperature to <200°C in a few minutes. The vessels were then depressurised and charges were recovered and re-weighed to ensure that water was retained through the experiment. Capsules were then opened and the crystals, none of which were cracked, were recovered and mounted in 1 inch epoxy discs, with the c axis of the crystal in the plane parallel to the face of the disc.

The epoxy discs were then prepared as thick sections (~400 μm thickness) for FTIR spectroscopy, ensuring that the section represented approximately the core of the quartz cube. One side was polished (1–3 μm diamond) and the other side was flattened with alumina slurry (>15 μm grit) only. It was found that the latter preparation gave more consistent ablation behaviour when conducting laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) analyses with our system.

**Experimental methods**

*Hydrogen in-, lithium-out ([H-in]/[Li-out]) experiments*

These and all other experiments were conducted at the University of Lausanne. Gold tubing of 4.5 mm O.D. and 4.1 mm I.D. was cut into 20–35 mm lengths, which were cleaned using acetone, then triple crimped and welded at one end. The open capsules were then annealed over a propane flame, then cleaned with boiling concentrated HCl, acetone then ethanol and dried at 60°C. They were then packed with SiO2 powder, together with ~1.5 mm cubes of quartz and ~20 μl of milliQ or HCl–milliQ mixes to give room temperature and pressure pH of 1, 3, 5 or 7. One experiment was run at each pH condition. The experiments at pH1, pH5 and pH7 contained only one crystal of BRA3 – these samples are denoted HQHP-pH1, HQHP-pH5 and HQHP-pH7, respectively. The experiment at pH3 contained one crystal each of BRA3, TIB2 and TIB6, with the samples denoted HQHP-pH3, HQHP-TIB2 and HQHP-TIB6, respectively. Further experimental details are provided in Table 1. At run conditions, the pH values are 5.39, 6.38, 6.88 and 6.90, respectively (see column “pH (expt. P, T)” in Table 1), with log10K for the dissociation reactions of H2O (~13.8) and HCl (~9.79) calculated using SUPCRT (Johnson et al., 1992). Values obtained were extrapolated using the density correlation approach (Eugster and Baumgartner, 1987; Manning 1994; Dolejs and Manning, 2010). The program Soluble (Roselle and Baumgartner, 1995) was used to calculate pH, using a Debye-Hückel activity model. Values of the pH in the text herein, refer to pH at room temperature (i.e. 1, 3, 5 and 7).

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**Deuterium-in, hydrogen-out ([H-in]/[H-out]) experiment**

This experiment is denoted HQHP-H2H. For a true H–H exchange experiment, the only exchange should be between H and 2H. Thus, Li or other monovalent cations should be eliminated from the crystal before the experiment. Such experiments must be undertaken in two steps, firstly to equilibrate (or metastable equilibrate) fully the crystal with H at the run conditions, followed by exchange with 2H (e.g. Kurka et al., 2005; Novella et al., 2017).

To this end, crystals of BRA3 were placed into Au capsules with a H2O–HCl mix at pH3, welded, as above, then annealed

| Table 1. Experimental conditions. Each entry in the ‘Experiment’ column represents a single run. Some of the runs contained multiple crystals, these are each given individual sample IDs (final column). The experiment labelled ‘Pre-eq.’ was the pre-equilibration step before the H–H exchange experiment ([H/H]). HCl was 1 M. |
| --- | --- | --- | --- | --- | --- | --- | --- |
| H-in/Li-out | Experiment | T (°C) | P (GPa) | t (hr) | Fluid composition | Nominal pH | pH (expt. P, T) | Crystals used | Sample ID |
| 'pH1' | 800 | 0.1 | 1 | HCl–milliQ | 1 | 5.39 | BRA3 | HQHP-pH1 |
| 'pH3' | 800 | 0.1 | 1 | HCl–milliQ | 3 | 6.38 | BRA3 | HQHP-pH3 |
| 'pH5' | 800 | 0.1 | 1 | HCl–milliQ | 5 | 6.88 | BRA3 | HQHP-pH5 |
| 'pH7' | 800 | 0.1 | 1 | milliQ | 7 | 6.90 | BRA3 | HQHP-pH7 |
| 2H-in / H-out | Experiment | T (°C) | P (GPa) | t (hr) | Fluid composition | Nominal pH | pH (expt. P, T) | Crystals used | Sample ID |
| 'DQHP' | 800 | 0.1 | 1 | HCl–2H2O | 3 | 6.38 | BRA3 | DQHP-H2 |
| 'DQHP' | 800 | 0.1 | 53.5 | HCl–2H2O | 3 | 6.38 | BRA3 | N/A |
at 800°C, 0.1 GPa for 53.5 hr. The crystals were then recovered, as above. One crystal was doubly polished and analysed for homogeneity by FTIR spectroscopy (further information is provided in the Supplementary Appendix).

One other crystal was then placed into an Au capsule with nearly pure \( ^2 \text{H}_2\text{O} \) and welded closed, as with the H-in/Li-out experiments. To set the pH, and to avoid large changes in the H/\( ^2 \text{H} \) ratio of the \( ^2 \text{H}_2\text{O} \), 0.6 \( \mu \text{L} \) of 1 M HCl was added to a 0.6 \( \mu \text{L} \) vial of \( ^2 \text{H}_2\text{O} \) (99.99%, Sigma Aldrich) as it was opened. Capsules were filled and welded within ~2 hr of opening the vial of \( ^2 \text{H}_2\text{O} \), which was not re-used, so \( \text{H}^– - \text{H}^+= \) exchange with the atmosphere should be negligible. This experiment was then run as with the H-in/Li-out experiments, for 1 hr at 800°C, 0.1 GPa. Following the experiment, the crystal was mounted in epoxy and prepared for FTIR and LA-ICP-MS analyses, as above.

**Deuterium-in, hydrogen and lithium-out \((^2\text{H}-\text{in}/\text{H}+\text{Li} \text{ out})\) experiments**

The experimental method was a hybrid of the H-in/Li-out and \(^2\text{H}-\text{in}/\text{H} \text{-out} \) experiments. Crystals of TIB2, TIB6 and BRA3 were welded into an Au capsule with SiO\(_2\) powder and pH3 \(^2\text{H}_2\text{O} \) then run at 0.1 GPa, 800°C for 1 hr, and the crystals were recovered and prepared as above. The samples are denoted DQHP-TIB2, DQHP-TIB6 and DQHP-BRA3. For reasons described above, these experiments cannot be considered as \( \text{H}^- - \text{H}^+ \) exchange experiments given the presence of Li in the starting material.

**Analytical methods**

**FTIR spectroscopy**

FTIR spectra were recorded at the University of Bern using a Bruker Tensor II spectrometer coupled to a Bruker Hyperion 3000 microscope, equipped with an XYZ imaging stage and a dry-air-purged analytical chamber. Spectra were recorded (32–128 scans, 8 cm\(^{-1}\) resolution) using a single mercury cadmium telluride (MCT) detector, with a 25–40 \( \mu \text{m} \times 150–200 \mu \text{m} \) on-sample aperture, with the long axis oriented parallel to the crystal edge. Such FTIR analyses sample a Gaussian volume within the sample (Ni and Zhang, 2008) with a full width at half maximum that may be considerably larger than the apparent sampling width determined by the nominal aperture size. Therefore, the diffusion coefficients reported herein can be considered maxima due to convolution effects (e.g. Jibamitra et al., 1988; Jollands, 2020). Spectra were recorded every ~10–20 \( \mu \text{m} \) along a 1D profile parallel to the c axis. Whilst some profiles were also measured perpendicular to the c axis, the effective diffusion distances were generally only several tens of micrometres. This is to be expected, given the extreme anisotropy of diffusion for monovalent cations in quartz (e.g. Verhoogen, 1952; Frischat, 1970; Jollands et al., 2020a), where diffusion parallel to the c axis is considerably faster than diffusion perpendicular to c. This effectively precludes quantitative descriptions of diffusivity or diffusion mechanisms perpendicular to the c axis – such short profiles will be strongly affected by convolution artefacts. Qualitatively though, this is in line with previous demonstrations of the diffusive anisotropy of H.

Data were processed by baseline subtraction using a concave rubberband algorithm (built into Bruker OPUS software) with 256 baseline points and 3 iterations. Further information is provided in the Supplementary Appendix. Correction to 1 cm thickness was done on baseline uncorrected data, using the orientation-independent relationship valid for both polarised and unpolarised light derived by Jollands et al. (2020a), based on the height of the 2675 cm\(^{-1}\) band from a linear baseline drawn between 2548 and 2750 cm\(^{-1}\).

Most analyses were done using polarised light, with each profile measured twice, once with E\( || \text{c} \) and once with E\( \perp \text{c} \), giving total absorbance \( \sum \text{Abs} = 2 \times \text{Abs}_{(\text{E} || \text{c})} + \text{Abs}_{(\text{E} \perp \text{c})} \). Some measurements were done unpolarised, where we assume \( \sum \text{Abs} \approx 4 \times \text{Abs}_{(\text{unpol})} \). The latter approximation is only valid for quartz measured on a plane including the c axis, because almost all absorbance in the OH stretching region occurs when E\( \perp \text{c} \) (e.g. Brunner et al., 1961; Baron et al., 2015; Potrafke et al., 2019), thus, in this plane, \( \text{Abs}_{(\text{unpol})} \approx 0.5 \times \text{Abs}_{(\text{E} || \text{c})} \). Any spectra showing clear contamination by epoxy resin were removed. The distance between the first un-contaminated spectrum and the crystal edge was measured using a photomicrograph for each profile.

The remaining spectra were resolved into a series of pseudo-Voigt peaks, as described in the Supplementary Appendix. Then, the resolved peaks were integrated and grouped together into defect associations (Table 2) based mainly on previous work (Sibley et al., 1979; Halliburton et al., 1981; Kronenberg, 1994; Bachheimer, 1998; Stalder and Konzett, 2012; Baron et al., 2015; Frigo et al., 2016). Herein, defects are described in the text body using ‘short-hand’ notation, with curly braces (the first column of Table 2). These and other defects referred to in this study are presented in end-member and Kröger-Vink notation in Table 2. Broadly, there are two classes of defects – those that are associated with Al\(^{13+}\) replacing Si\(^{4+}\) (e.g. \{AlH\}, \{AlLi\}), and those probably associated with excess O\(^{2–}\) (e.g. \{LiOH\}). Visualisations of the atomic structures of these defects can be found in Jollands et al. (2020b), their figures 4–7. The resolved peaks at 3190, 3205 and 3296 cm\(^{-1}\) (expressed as two bands due to overlap between the former two) are assigned to a Si–O vibration (Kats, 1962). This assignment remains the subject of debate – Biró et al. (2016) suggests they might be related to surficial water, though this will not affect the results of this study given that neither represents structurally bound OH groups. The possibility that \{HOH\} is represented by a band at 3475 cm\(^{-1}\) is considered in the discussion.

**LA-ICP-MS**

LA-ICP-MS analyses were conducted using an Atlex 193nm laser housed inside an Australian Scientific Instruments RESOLUTION
laser ablation system, equipped with a dual-volume Laurin Technic cell (S155), at the University of Lausanne. A 6×7 square grid of 38 μm laser spots, with ∼70 μm spacing between points, was defined, then sheared by ∼30° into an approximate parallelogram, such that each spot was slightly further from the interface than the last. Analyses were done between ∼13–17 J cm⁻² on-sample fluence and 30–40 Hz. The sample was ablated for 20 s, with 10–20 s background before and after each point. A standard (NIST SRM 610) was analysed after every 12 points – this was done at 10 Hz, 5 J cm⁻² fluence. 29Si (internal standard), 7Li, 11B, 23Na, 27Al, 39K and 49Ti were counted for 0.01–0.1 s per sweep, giving a total sweep duration of 0.5–0.8 s. Data were processed using Iolite (Paton et al., 2011). The first 4 s of each analysis were deleted to reduce the potential for surface contamination. Further information is provided in the Supplementary Appendix.

Results

FTIR spectra

H-in/Li-out experiments
Crystal core and rim spectra from these and all other experiments are shown in Fig. 1. As expected for a H-in experiment, spectra from the rims show consistently greater absorbance in the OH stretching region. In general, these are dominated by bands associated with [AlH]. The 3481 cm⁻¹ peak, attributed to [LiOH], which is present in some of the core spectra, is consistently absent in rim spectra. The near-rim spectra of sample HQHP-TIB2 show an additional band at 3475 cm⁻¹, which was not present in the starting material. Both the rim and core spectra of sample HQHP-TIB6 show low absorbance of [AlH], which is barely resolvable. There is no apparent effect of experimental pH on rim spectra, i.e. the rim spectra from experiments HQHP-pH1, -pH3, -pH5 and -pH7, all of which used the same starting material (crystal BRA3) are similar.

2H-in/H-out experiment
Following the initial long (53.5 hr) anneal prior to the diffusion experiment, the spectra showed only bands associated with [AlH], with constant absorbance across the crystal (more information is provided in the Supplementary Appendix). Following the short (1 hr) diffusion experiment (HQHP-H²H), as with the 2H-in/H + Li-out experiments, the interface spectrum shows a decrease, but not total loss of, absorbance in the O–H stretching region. The only O–H stretching bands in the core are the triplet associated with [AlH] (Fig. 1). At the rim, the main band associated with [Al²H] at 2505 cm⁻¹ is clearly visible. The other band of the doublet at 2473 cm⁻¹ is not clearly visible, but is resolvable when the spectra are fitted. Spectra from this experiment show contributions from neither [LiOH] nor [LiO²H].

2H-in/H + Li-out experiments
Crystal rim spectra (Fig. 1) show near-complete loss of absorbance in the OH stretching region relative to the core spectra. The peaks associated with [AlH] and [LiOH] are absent within the resolution of our technique (experiments DQHP-TIB6, DQHP-BRA3) or nearly so (DQHP-TIB2). This loss of absorbance in the O–H stretching region is associated with a gain in the O–²H stretching region. The peaks associated with O–²H bonds overlap with Si–O overtones, thus quantifying their absorbance requires spectra to be resolved into separate peaks, some of which are assigned to O–²H and others to Si–O. The [Al²H] defect is expressed as a doublet at 2505 and 2473 cm⁻¹. The [LiO²H] defect is expressed as a peak at 2572 cm⁻¹.

We note that the presence of these peaks associated with O–²H does not appear to affect the validity of the thickness correction method, even though this used a band (2675 cm⁻¹) in an overlapping wavenumber region.

Profile lengths and shapes
Spectra as a function of distance from the crystal edge for two experiments are presented in Fig. 2. Profiles of defect-specific
Fig. 3. FTIR profiles resolved into individual defect associations, together with the Li content measured by LA-ICP-MS. H-in/Li-out experiments are shown in panels (a)–(e) and (g), (i) shows the $^2$H-in/H-out experiment, and all others (f, h, j) show profiles from the $^3$H-in/H + Li-out experiments.
Al$^{3+}$ is charge-compensated by an interstitial Li, i.e. $\{\text{AlLi}\}$. Next, (b) the $\{\text{AlH}\}$ dissociation with the Al from the previously dissociated $\{\text{AlH}\}$, similar to (c), but with the Li now mobile. In (f), the mobile Li moves into coordination with the O ions forming the c axis parallel double helix. In (d) there is an exchange reaction where the mobile H displaces the immobile Li. The cartoon shows this occurring simultaneously, though this may not be the case. Image (e) is the cartoon of the diffusion plus reaction concept modelled in this study. The initial condition is in (a), where one OH$^-$ group exists adjacent to an Al$^{3+}$, and another Al$^{3+}$ is charge-compensated by an interstitial Li, i.e. $\{\text{AlLi}\}$. Next, (b) the $\{\text{AlH}\}$ dissociates, thus mobilises the H. (c) The mobile H moves. The model shows it moving towards the other Al$^{3+}$; this would be better represented as a random walk, possibly hopping between O ions forming the c axis parallel double helix. In (d) there is an exchange reaction where the mobile H displaces the immobile Li. The cartoon shows this occurring simultaneously, though this may not be the case. Image (e) is similar to (c), but with the Li now mobile. In (f), the mobile Li moves into coordination with the Al from the previously dissociated $\{\text{AlH}\}$.

Absorbance (following resolution of spectra into pseudo-Voigt curves) and Li from each sample are presented in Fig. 3. For clarity, every plot in Fig. 3 is also presented in the Supplementary Appendix with expanded scales.

**H-in/Li-out experiments**

Profiles measured from experiments using the same starting material (BRA3) though with different pH (HQHP-pH1, -pH3, -pH5, -pH7) are similar, with no systematic variation in H concentration or profile form as a function of experimental pH. They all show $\{\text{AlH}\}$- and $\{\text{LiOH}\}$-versus-distance curves that do not correspond to an error-function shape (Fig. 3a, c, g, e). Rather, they show stepped shapes, with a relatively high absorbance, gently dipping section near the interface towards the core, then a sharp decrease towards background absorbance. Li profiles show the inverse trend: Li below detection limits near the rim, then an increase in Li towards the core. Whilst the effective profile lengths (600–700 μm) are not identical for the four experiments conducted at different pH conditions, there does not appear to be any systematic relationship between profile length and pH. The absorbance–distance curve associated with the $\{\text{LiOH}\}$ defect shows a similar form to that associated with Li, albeit at much lower absorbance.

The $\{\text{AlH}\}$ profile in experiment HQHP-TIB2 (Fig. 3b) shows a gradual decrease in absorbance from rim to core, with an approximate effective diffusion distance of ~600 μm. However, this profile is not consistent with an error-function form, as indicated by the near linear decrease in absorbance away from the crystal edge. Again, the Li profile approximately mirrors the $\{\text{AlH}\}$ profile. The $\{\text{LiOH}\}$ profile shows a stepped shape, with relatively high absorbance in the core, then a decrease towards the rim, and a flat, low absorbance section in the ~100 μm closest to the rim. The profile of the band assigned to [HOH] mirrors that of $\{\text{LiOH}\}$ – this is the main reason for assigning the band to [HOH]. This is discussed in more detail below with regards to modelling the profiles and associated caveats.

The $\{\text{AlH}\}$ and Li profiles in experiment HQHP-TIB6 appear to be relatively flat (Fig. 3d), however the absorbance, and Li contents, are so low that it is difficult to confidently describe these profiles, so they are not discussed further.

$^{2}$H-in/H-out experiments

Profiles of absorbance versus distance of the bands attributed to $\{\text{AlH}\}$ and $\{\text{Al}^2\text{H}\}$ from experiment HQHP-H$^2$H show forms apparently describable as error functions (Fig. 3i), with $\{\text{AlH}\}$ showing out-diffusion and $\{\text{Al}^2\text{H}\}$ showing in-diffusion, as expected given the experimental design. A series of spectra from this experiment, showing the variation from core to rim in both the O–H and O–$^{2}$H stretching regions, is presented in Fig. 2. The profile lengths associated with both defects are equal at ~300 μm, which is shorter than the profile lengths from the H-in/Li-out experiments, which were run at the same P-T-t conditions.

$^{2}$H-in/Li + H out experiments

These experiments consistently show profile forms and length scales that are considerably more complex than both the $^{2}$H-in/H-out and H-in/Li-out series. The absorbance of experiment DQHP-TIB2 (Fig. 3f) is dominated by the bands attributed to $\{\text{Al}^2\text{H}\}$, which shows a near-linear decrease from rim to core in the first ~250 μm of the crystal, followed by a gradual decrease from ~250–450 μm from the rim. The profile of absorbance attributed to $\{\text{AlH}\}$ generally shows out-diffusion, but with maximum absorbance at ~300 μm from the rim, then a decrease in absorbance, appearing to reach background values at ~600 μm from the rim. $\{\text{LiOH}\}$ and $\{\text{LiO}^2\text{H}\}$ profiles show forms approaching step shapes, and both are approximately 400 μm long. The Li profile is ~500 μm long showing out-diffusion, with a concave-up form in its near rim section, and a concave down section near the core, the latter being approximately consistent with an error function.

The profiles from experiment DQHP-BRA3 (Fig. 3b) show similar relative behaviour to DQHP-TIB2, albeit at much lower absorbance. The $\{\text{AlH}\}$ profile shows a local absorbance maximum at ~300 μm from the rim. The main difference in profile forms between DQHP-BRA3 and DQHP-TIB2 is that, in the former, $\{\text{Al}^2\text{H}\}$ shows a near linear decrease from the rim directly towards background values, whereas in the latter, the absorbance approached the background values gradually. As with experiment HQHP-TIB6, DQHP-TIB6 (Fig. 3i) has low H, $^2$H and Li...
contents with considerable scatter, such that the data cannot be described in a meaningful way. This experiment is not discussed further.

**Discussion**

**Developing a multi-site diffusion model**

In the vast majority of studies of diffusion in geologically relevant crystals using in situ analytical methods, profiles of trace-element concentration (C) versus distance (x) from the diffusion interface had forms corresponding to the error function (e.g. Cherniak, 2003). Such profiles can be fitted to equation 1, from which a diffusion coefficient, D, is obtained, when the time (t) is known.

\[ C(x, t) = C_{\text{core}} + (C_{\text{rim}}-C_{\text{core}}) \text{erfc}(\frac{x}{2\sqrt{Dt}}) \]  

(1)

\( C_{\text{rim}} \) and \( C_{\text{core}} \) are the concentrations at the crystal rim and core, respectively and \( \text{erfc} \) is the complementary error function. This relationship assumes 1D, concentration-independent diffusion in a semi-infinite medium, plane-sheet geometry, with constant boundary conditions (Crank, 1975). In this study, the only sample showing such a profile geometry is sample DQHP-H†H, i.e. from the H-in/H-out experiment (Fig. 3i). This implies that, in general, it is not reasonable to describe this system using the above assumptions. However, given the experimental and analytical methods, there is no reason to assume that such geometrical and boundary conditions are inappropriate, and there is also no obvious reason to invoke concentration-dependent diffusion for H, especially at such low concentrations, and tests invoking different forms of concentration-dependent diffusion (linear and exponential) could not reproduce the observed profile shapes.

Rather, some of the profile shapes resemble diffusion coupled to inter-site reactions (Dohmen et al., 2010). This describes a situation where a crystal has at least two sites on which the diffusant can sit, where each of the sites is associated with a certain diffusivity, and where it can move between the sites. Whilst this process can yield a variety of profile shapes, the most characteristic form in experiments with constant boundaries (in terms of both composition and position) is the step shape (e.g. Fig. 3a, c, g, e), though various other deviations from error-function forms have also been observed (Holycross and Watson, 2017; Bloch et al., 2020). This is broadly similar to the ‘trapping’ behaviour that has been well-known in metallurgy since the mid 1900s (Darken and Smith, 1949).

There is an established precedent derived from conductivity studies for invoking such a model to describe H diffusion in quartz. Kronenberg and Kirby (1987), expanding on a model proposed by Hughes (1975) and Jain and Nowick (1982) described the process as follows: “diffusion of H at high temperatures requires the dissociation of Al-H pairs to form free H" followed by transport to neighboring \( \text{Al}_{\text{Si}} \) sites”, referring to the [AlH] defect, the major defect in most natural and some synthetic quartz (e.g. Halliburton et al., 1981; Müller and Koch-Müller, 2018; Tollan et al., 2019). The implication is that the overall mobility of H should be a function of: (1) the rate at which free H can hop through the lattice; and (2) a constant describing dissociation of charge-neutral associations. This is expanded on below, considering each type of experiment in turn. Further information regarding all models, including curve fitting, is provided in the Supplementary Appendix.

**H-in/Li-out experiments**

For these experiments, we first consider that H moves by dissociation of \( \text{H}_{\text{i}}^{\dagger} \) (or \( \text{H}_{\text{Si}}^{\dagger}, \text{Li}_{\text{i}}^{\dagger} \)) from its charge-balancing ion (\( \text{Al}_{\text{Si}}^{\dagger} \)), followed by hopping to a nearby position. The process is illustrated in cartoon form in Fig. 4.

The dissociation of the [AlH] defect, which dominates the FTIR spectra (Figs 1 and 3) will be described by equation 2, where the H is described generically as ‘interstitial’ with an i:

\[ \text{Al}_{\text{Si}}^{\dagger} - \text{H}_{\text{i}}^{\dagger} \leftrightarrow \text{Al}_{\text{Si}}^{\dagger} + \text{H}_{\text{i}}^{\dagger} \]  

(2)

A similar reaction should be possible for Li:

\[ \text{Al}_{\text{Si}}^{\dagger} - \text{Li}_{\text{i}}^{\dagger} \leftrightarrow \text{Al}_{\text{Si}}^{\dagger} + \text{Li}_{\text{i}}^{\dagger} \]  

(3)

These reactions are equivalent to, for example, eqs. 1b and 1a, respectively, in Campone et al. (1995) and eq. 1 of Jain and
Nowick (1982). That the H and Li are both given subscript $i$ should not be taken to suggest that they occupy the same interstitial site: $\{\text{Al' Si} - \text{H}^\dagger_i\} \times$ can also be written (more correctly) as $\{\text{Al' Si} - \text{OH}^\dagger_1\} \times$, for example, and the Li probably occupies the large open channel parallel to the $c$ axis.

For simplicity, we assume that in a simple H–Li exchange, it should be possible to combine the two reactions, upon which the Al$_{\text{Al}}$ cancel out. In order to distinguish between the H or Li in ‘interstitial’ sites associated with Al (immobile), and those in a dissociated, mobile configuration, we use ‘$i2$’ and ‘$i1$’ for the former and latter, respectively. Herein, ‘$i1$’ always refers to an interstitial site where the occupying ion is free to move. Therefore, combining equations 2 and 3, and adding this notation, we have:

$$\{\text{Al' Si} - \text{H}^\dagger_{i2}\} \times + \text{Li}^\dagger_{i1} \leftrightarrow \{\text{Al' Si} - \text{Li}^\dagger_{i1}\} \times + \text{H}^\dagger_{i2}$$

which describes the dissociation of [AlLi] and [AlH], forming two mobile ions that swap places forming two new immobile defects.

---

**Fig. 5.** Profiles of [AlH] and [AlLi] from experiments (a) HQHP-pH1 and (b) HQHP-pH5, fitted. The measured Li content is corrected to account for the [LiOH] defect. Model parameters: total time = 3600 s, $\Delta x = 25 \mu m$. Initial and boundary conditions: (a) $\sum i_1$ (mobile) $+ \sum i_2$ (immobile, Al-associated) = 24.4 $/10^6$Si; $\sum i_1$ (initial) = 17.7 $/10^6$Si; $\sum i_1$ (boundary) = 0 $/10^6$Si. (b) $\sum i_1$ (mobile) $+ \sum i_2$ (immobile, Al-associated) = 24.9 $/10^6$Si; $\sum i_1$ (initial) = 17.9 $/10^6$Si; $\sum i_1$ (boundary) = 0 $/10^6$Si. All other fit parameters ($D$, $K$, $\gamma_2$, $\epsilon$) are provided in Table 3. Fits to data from other experiments are provided in the Supplementary Appendix.

**Fig. 6.** The profile from experiment HQHP-TIB2, resolved, with the [AlH], [LiOH], [HOH] (assumed to be represented by a band at 3475 cm$^{-1}$) and $\sum$ Li modelled simultaneously. This fit was done manually, i.e. no non-linear least-squares regression was attempted, so is not guaranteed to be the best fit. Model parameters: $\Delta x = 25 \mu m$, total time = 3600 s, $\log_{10} D$ ($m^2 s^{-1}$) = 9; $\sum i_1$ (mobile) $+ \sum i_2$ (immobile, Al-associated) = 105 $/10^6$Si; $\sum i_3$ (immobile, O-associated) = 8 $/10^6$Si; $K_1$ = 0.1; $K_2$ = 0.2.

**Fig. 7.** Models (solid lines) representing the $^2$H-in/H-out experiment, together with data (circles) from experiment HQHP-H$^2$H. The model parameters are provided in Table 4. Model parameters: $\Delta x = 25 \mu m$; total time = 3600 s. Initial and boundary conditions (for other model parameters, see Table 4): $\sum i_1$ (mobile) $+ \sum i_2$ (immobile, Al-associated) = 18.5 $/10^6$Si; $\sum i_3$ (mobile) = 16 $/10^6$Si; $\sum i_4$ (boundary) = 16 $/10^6$Si.
For the purposes of modelling, this is again simplified by removing the Al to give equation 5:

\[
H^*_{i2} + Li^*_{i1} \leftrightarrow Li^*_{i2} + H^*_{i1}
\]  

The method for modelling this process is described briefly here, with further details, including fitting routines, estimations of uncertainties, model limitations, etc., in the Supplementary Appendix. The method involves dividing the total model time into a series of small steps, i.e. the explicit finite difference method. Each time step comprises a diffusion step (\(\Delta x = 2.5 \mu m\), \(\Delta t\) adjusted to maintain numerical stability), where the mobile H and Li diffuse (diffusion only on the \(i_1\) site, i.e. only \(Li^*_{i1}\) and \(H^*_{i1}\)), assuming that a single diffusion coefficient can describe both H and Li, then a reaction step where the reaction in equation 5 is modelled. This is done using the equilibrium expression (herein \(K_1\) for the reaction in equation 5) specified, using square brackets for concentration (now using units of atoms per \(10^6\) Si):

\[
K_1 = \frac{[H^*_{i1}][Li^*_{i1}]}{[Li^*_{i2}][H^*_{i2}]}
\]

and with various constraints of mass balance and site balance imposed:

\[
\sum i_1 = [Li^*_{i1}] + [H^*_{i1}]
\]

\[
\sum i_2 = [Li^*_{i2}] + [H^*_{i2}]
\]

\[
\sum H = [H^*_{i1}] + [H^*_{i2}]
\]

\[
\sum Li = [Li^*_{i1}] + [Li^*_{i2}]
\]

Solving these equations yields the concentrations of \(Li^*_{i1}\), \(H^*_{i1}\), \(Li^*_{i2}\) and \(H^*_{i2}\), which is possible if four of \(K_1\), \(\sum i_1\), \(\sum i_2\), \(\sum H\) and \(\sum Li\) are known (further information in the Supplementary Appendix). After the reaction step, diffusion occurs again, then reaction, and so on, until the total model time is reached. The final model output is a series of curves representing the concentration of each defect as a function of distance from the boundary, which are then fitted to the data using non-linear least-squares regression. Uncertainties (95% confidence intervals) for the fitted

### Table 4. Results from fitting the profiles described from the \(^2\)H-in/H-out experiment.

| Fixed variables | Fitted variables |
|-----------------|------------------|
| \(\sum i_2 + \sum i_1\) | \(\sum i_1 = 0.1\) |
| \(\sum H\) (initial) | \(\log_{10}D\) (m² s⁻¹) | \(\chi^2\) |
| \(\sum H\) (boundary) | \(-9.28 (0.03)\) | 0.89 |
| Log₁₀\(K_3\) | \(-10.28 (0.03)\) | 0.89 |

See Table 3 for notation details. \(K_3\) = equilibrium constant from equation 14. In this case \(K_3\) was set to 1. Again, fits were done with both \(\sum i_1 = 0.1\) and 1.
model parameters, $K_1$ and $D$, were determined using the constant chi-squared boundary method (Press et al., 2007). The initial and boundary concentrations of $\Sigma i_1$, $\Sigma i_1$ and $\Sigma i_2$ were set manually for each experiment. The best fit parameters and uncertainties are presented in Table 3 for the five H-in/Li-out experiments modelled in this way. The $\Sigma i_1$ parameter is problematic, as we have no independent constraints on its value, so, for internal consistency only, this was set to either 0.1 or 1 atoms/$10^6$ Si. This is discussed further below with regards to model caveats.

Model fits for profiles from two H-in/Li-out experiments are presented in Fig. 5, showing that the model successfully reproduces the forms and lengths of the profiles.

**[AlH] and [LiOH] defects in H-in/Li-out experiments**

A limitation of the model development above is that it did not consider the [LiOH] defects that are visible in the FTIR spectra of most experiments as a band at $\sim 3481$ cm$^{-1}$ (e.g. Baron et al., 2015; Frigo et al., 2016). Adding this defect adds considerable complexity and is only treated briefly here (a detailed description is provided in the Supplementary Appendix), and non-linear least-squares regressions were not attempted – all fitting was done visually only.

As with the reactions presented above, we assume that [LiOH] can dissociate, forming a mobile H or a mobile Li. If we assume that there exists also a [HOH] defect, then we have:

$$[\text{H}_3^+ - \text{O}^- - \text{H}_i^+] + \text{Li}_i^+ = [\text{Li}_3^+ - \text{O}^- - \text{H}_i^+] + \text{H}_i^+$$  \hspace{1cm} (11)

now denoting one of the interstitial sites as $i_3$, and keeping the mobile interstitial site as $i_1$. As another simplifying assumption, the other $i$ site is assumed to not be involved in the reaction, and is left as $'i'$. It is of course possible that there also exists a reaction involving a defect with Li$_2$O stoichiometry, however this possibility is not considered. Now, adding the reaction in equation 11 into the reaction step in the diffusion–reaction model (the equilibrium constant for this reaction is herein denoted $K_2$):

$$K_2 = \frac{[\text{H}_i^+][\text{Li}_i^+]}{[\text{Li}_i^+][\text{H}_i^+]^2}$$  \hspace{1cm} (12)

and assuming that the free $\text{H}_i^+$ and $\text{Li}_i^+$ can move between defects associated with O$^-$ and Al$_{i_3}$, profiles for [LiOH], [HOH], [AlH] and [AlLi] are generated simultaneously.

An example model output is shown in Fig. 6 along with the data from experiment HQHP-TIB2. Modelling results show that the two Li-bearing defects [LiOH] and [AlLi] show out-diffusion, where [HOH] and [AlH] show in-diffusion profiles. One shortcoming of the model is that it requires a [HOH] defect, however such a defect is not definitively known in the FTIR spectra of quartz – this is discussed further below with other model caveats.

**$^2$H-in/H-out experiment**

The profiles from this experiment appear to show error-function forms (Fig. 3i), which might be considered as consistent with simple concentration-dependent diffusion, i.e. equation 1. However, if one assumes that H–Li exchange requires a diffusion–reaction model then there is no obvious reason that this should not also be the case for H–$^2$H exchange. Therefore, a diffusion–reaction model was constructed based on equation 13, i.e. the $^2$H equivalent of equation 4:

$$[\text{Al}_{i_3}^+ - ^2\text{H}_i^+] + \text{H}_i^+ \leftrightarrow [\text{Al}_{i_3} - \text{H}_i^+] + ^2\text{H}_i^+$$  \hspace{1cm} (13)

where, as before, ‘i1’ represents a mobile position, and ‘i2’ is immobile. We assume that the equilibrium constant for this reaction (herein K3) can be approximated as 1, at least within the
Table 5. Results from approximate visual fitting of two $^2$H-in/H+Li out experiments.

|                  | $\Sigma_1$ | $\Sigma_2$ | $\Sigma_{Li}$ (initial) | $\Sigma_{Li}$ (boundary) | $\Sigma_H$ (initial) | $\Sigma_H$ (boundary) | $\log_{10} K_1$ | $\log_{10} K_3$ |
|------------------|-------------|-------------|--------------------------|--------------------------|-----------------------|-----------------------|----------------|----------------|
| DQHP-TIB2        | 1           | 134         | 54.9                     | 2                        | 80                    | 4                     | -9.1           | -1.05          |
| *DQHP-BRA3       | 1           | 29          | 17.9                     | 0.5                      | 12                    | 0.1                   | -5.7           | -0.92          |

See Table 3 for notation details. $K_1$ and $K_3$ are the equilibrium constants from equations 6 and 14, respectively. These fits were done only with $\Sigma_1 = 1$.

limitations of our experimental and analytical methods:

$$K_3 = \frac{[H^+_1][^2H^+_1]}{[H^+_1][H^-_1]} \approx 1$$  \hspace{1cm} (14)

The profile from this experiment was then modelled and fitted, as above, with H contents (as H$_2$O) determined using $\varepsilon = 89,000$ L cm$^{-2}$ mol$^{-1}$ (Thomas et al., 2009), and $^2$H contents using $\varepsilon = 79,700$ L cm$^{-2}$ mol$^{-1}$. The derivation of the $\varepsilon$ value for H$_2$O is described in full in the Supplementary Appendix, however, simply $\varepsilon$ for H$_2$O is adjusted to minimise the variance of $^2$H + H (atomic basis) for all points along the whole profile, whilst keeping $\varepsilon$ for H$_2$O constant. The assumption is that $^2$H + H should be constant along the profile in the $^2$H-in/H-out experiment, i.e. every $^2$H gained is compensated by one H being lost.

When K is set to 1, the modelled profiles approach error-function shapes, as we observe in the $^2$H-in/H-out experiment. It appears to be possible to generate error-function forms that fit the data with any positive value of $\Sigma_1$ (total mobile H + $^2$H) up to the total value of H + $^2$H. As $\Sigma_1$ decreases, D increases, and where $\Sigma_1 = H + ^2H \approx 18.5 / 10^6$ Si (the total H + $^2$H is set to 18.5 atoms per $10^6$ Si, see Fig. 7, Table 4), D equals that determined by fitting the data to equation 1.

However, the starting material for the $^2$H-in/H-out experiment prior to the initial equilibration step was the same as that used for the ‘pH series’ (HQHP-pH1, -pH3, -pH5 and -pH7, crystal BRA3). The stepped profiles from the latter series could not be well fitted with $\Sigma_1$ over ~3–5 / 10$^6$ Si – further information is provided in the Supplementary Appendix. If we then consider the results of internally consistent fitting of both the ‘pH series’ and this HQHP-$^2$H experiment, the diffusion coefficient associated with $^2$H–H exchange is around one order of magnitude slower than that of H–Li exchange. This is not intuitively reasonable if we assume that diffusivity should increase with decreasing mass and ionic radius of the diffusant, but has been observed before – for example, the monovalent cation with the highest measured diffusivity in diffusion experiments to date is Na (Frischat, 1970). However, we note that Frischat (1970) used very different experimental methods (little information is provided, however assuming the methods are similar to Frischat (1968)) – radioactive $^{22}$Na was diffused into quartz from a film of $^{22}$NaCl at atmospheric pressure, then $^{22}$Na vs. depth profiles were measured using serial grinding and by measuring the residual activity.

Overall, we can therefore state that whilst the results of the $^2$H-in/H-out experiment do not require a diffusion–reaction model, a diffusion–reaction model can reproduce the approximately error function profile shapes.

$^2$H-in/Li + H out experiments

Modelling these experiments is considerably more challenging than the H-in/Li-out or $^2$H-in/H-out experiments, given that there are now (at least) six unknowns to be solved for in the reaction step, being the concentrations of [AlH], [Al$^2$H], [AlLi] plus the mobile Li, H and $^2$H (i.e. those in the ‘i1’ positions). This is already an incomplete description – ideally we should have more unknowns, i.e. the above plus [LiOH], [HOH] and [H$^2$OH] (and potentially [Li$_2$O]). Taking the simplified system, i.e. ignoring the O-associated defects, there is no analytical solution to the simultaneous equations in the reaction step, meaning that the equations have to be solved numerically. For this reason, as with the model presented above including the Al- and O-associated defects, the measured profiles from these experiments were not fitted by non-linear least-squares regression. Instead, the profiles of [AlH], [Al$^2$H] and [AlLi] from two experiments were visually fitted only by adjusting the initial and boundary concentrations of Li and H, $\Sigma_1$, $\Sigma_2$ and D.

Models are presented in Fig. 8, taking $K_1 = 0.1$ (H–Li exchange, equation 4) and $K_3 = 1$ (H–$^2$H exchange, equation 13) which are broadly consistent with the model outputs presented above. These models generate profiles with several interesting features. Firstly, the [AlH] profiles show maxima at ~300 µm from the crystal edge, where we would normally expect the maxima to be in the crystal cores. Secondly, the near-interface out-diffusion section of the [AlH] profile is concave-up, which is broadly the opposite of the error-function profile form. Thirdly, the [Al$^2$H] profile is concave-down near the interface, again the opposite of the error-function form. All of these model outputs are consistent with the data.

Of particular note are the local maxima in [AlH] profiles. Such features would generally be described as ‘uphill-diffusion’, i.e. diffusion of an element against its own concentration gradient, which is not uncommon in multi-component diffusion systems (Zhang, 2010). In this case, all modelled diffusion is downhill – the maxima are due to the presence of a reaction step in the model rather than uphill diffusion. Even though non-linear least-squares regression was not used (the profiles were matched to models by visual estimations only), these results provide clear support to the use of the diffusion–reaction model. They also support the use of such a model for the $^2$H-in/H-out experiments presented above, despite the model not being required for an adequate fit.

Model caveats and shortcomings

Whilst the model presented above is successful at fitting the results of different experiments with general internal consistency, there are several caveats, limitations and simplifying assumptions that warrant further discussion.

First, in equation 5, and then throughout, it was implicitly assumed that two dissociation and association reactions occur simultaneously. This is not physically realistic – the two reactions would ideally be modelled separately.
Second, and probably the most problematic shortcoming of this model is that it requires free H$_i$ and Li$_i$ to be invoked, the former not being associated with any known band in FTIR spectra of quartz. This means that the model needs to assume some concentration of H$_i$ + Li$_i$, i.e. the sum concentration of the mobile cations at any given time ($\sum i$, equation 7). The power of FTIR spectroscopy in being able to distinguish different point defects is somewhat inconvenient here – we would probably not be considering this caveat if only the total H content, rather than full FTIR spectra had been measured. In this study, all profiles were fitted with $\sum i$ set to both 0.1 and 1 atoms per 10$^6$ Si for internal consistency. Some profiles could be well fitted with much lower or higher values of $\sum i$ however 0.1 and 1 atoms per 10$^6$ Si allowed all profiles to be fitted – further information is provided in the Supplementary Appendix. In lieu of any obvious better alternative, we also made the assumption that these mobile cations (H$_i$ and Li$_i$) move into their immobile states ([AlH] and [AlLi]) at the end of the experiment, i.e. they move from ‘i1’ to ‘i2’ sites on quench.

Third, in all models, we assume that a single diffusion coefficient is sufficient along the ‘i1’ pathway. There is, however, no reason to assume that (for example) the diffusion coefficient of Li is equal to that of H. By making this assumption, we are effectively applying a H–Li inter-diffusion coefficient, although it is not stated as such (see eq. 5 of Stalder (2021) and the associated text for a discussion of this concept). Again, this is done for simplification – if we considered this as inter-diffusion, we would need two ‘tracer’ diffusion coefficients, one for Li and one for H (three for the $^2$H-in/H + Li out experiments), and D would be concentration-dependent, i.e. different at every point along the profile. At this stage of model development, the aim is to minimise the number of variables. It is likely, however, that the uncertainties related to D due to this model limitation are minor relative to those related to the unconstrained nature of $\sum i$ (second point, above).

Fourth, in modelling the HQHP-TIB2 profile, a [HOH] defect was assumed – this was represented by a band at ~3475 cm$^{-1}$, which has not been assigned to this defect. The main reason was that the absorbance of this band appears to be inversely proportional to that of the 3481 cm$^{-1}$ [LOH] band (Fig. 9). Some arguments for and against this attribution are presented in the following paragraph.

The 3475 cm$^{-1}$ band was present, although barely resolvable, in experiments HQHP-pH1, -pH3, -pH5 and -pH7. Additionally, polarised FTIR spectra show that the total absorbance of this band is entirely dominated by its ELc contribution (see the Supplementary Appendix), as with most bands in the OH stretching region of quartz – this means that if the band represents a defect with H$_2$O stoichiometry, it almost certainly includes structurally-bound OH. Jollands et al. (2020b) predicted that a defect with H$_2$O stoichiometry should have bands at 3411 and 3498 cm$^{-1}$, with the latter showing most absorbance with ELc, which would be broadly consistent with our suggestion, though there is no evidence for the former band (although it would probably be subsumed by the [AlH] bands, if it were present). Additionally, Kats (1962) suggested that a band at 3470 cm$^{-1}$ was not associated with any cation other than H$^+$ (summarised by Kronenberg et al., 1986). However, it was not observed in quartz crystals grown in the pure SiO$_2$-H$_2$O system (Stalder and Konzett, 2012).

Fifth, we have assumed that the equilibrium constants K1, K2 and K3 can be assumed to be constant along the profiles. This is unlikely to be exactly correct, however the ability to fit different profile forms using single equilibrium constants suggests that this effect is minor relative to the other caveats.

**The effect of pH**

No effect of experimental pH was observed, in terms of either interface H and Li contents, or diffusion profile length/geometry. The original motivation for conducting experiments at different pH conditions was to determine whether a different activity of protons in the fluid would change the interface H/Li ratio, which may then affect the profile length and shape. That no effect of pH was observed does not mean that no effect is possible, however it could be explained by the fact that (1) the difference in pH is considerably lower at experimental conditions than at room temperature or (2) the total amount of Li and H that can be incorporated into the crystals is so low, given the low Al content, that any small variations in Li and H concentration might not be resolvable. Experiments conducted at conditions where the relative pH differences are larger (i.e. at higher pressure) could be enlightening, as could experiments using quartz crystals with higher Al contents.

**Reconciling observations from H-in and H-out experiments?**

One peculiar observation is that our previous experiments considering H–Li exchange through Li-in/H-out diffusion (Jollands et al., 2020a) gave results that appear to be inconsistent with those presented in this study. Where the H-in/Li-out experiments in this study generally yielded H concentration profiles with complex shapes, those in the Li-in/H-out experiments showed shapes that were, within uncertainty, consistent with error-function forms (Jollands et al., 2020a).

In fact, if the flux in the diffusion–reaction model is reversed, i.e. such that the boundary conditions of Li and H lead to Li in-diffusion, but all other parameters (K, D, number of sites) are kept constant, profile forms close to error functions are the result. To illustrate this further, the Li and [AlH] profiles from the 807°C, Li-in/H-out experiments of Jollands et al. (2020a) (their Q1L1, Q2L1, Q3L1, Q4L1 and Q5L1) were re-fitted using the diffusion–reaction model. With $\sum i$ = 1/10$^6$Si, we obtain log$_{10}$D (m$^2$ s$^{-1}$) ≈ −8.5 to −9.5, and log$_{10}$K1 = −0.4 to 0.1 (Table 3), and the profiles from the previous study were fitted satisfactorily (Fig. 10, and Supplementary Appendix). Fitting the same profiles to a standard error-function solution to Fick’s second law gives log$_{10}$D (m$^2$ s$^{-1}$) between −11.1 to −11.8 (see Supplementary dataset 1 of Jollands et al., 2020a for original fit parameters), i.e. 2–3 orders of magnitude lower than that obtained by using the diffusion–reaction model. Another observation of the Jollands et al. (2020a) investigation was a relatively large spread in measured D at any given temperature – approximately ±0.3 orders of magnitude around the mean. It is likely that this was due to the use of a variety of different starting materials, however there was no clear systematics observed. Alternatively, this spread in D might relate to the application of a simple diffusion model to a system where a more complex diffusion–reaction model is warranted. However, attempts to re-fit all data from the previous study using the new model were unsuccessful in this regard, the difference between D becomes larger – this almost certainly relates to the fact that we have little or no constraint on the elusive $\sum i$ parameter, either in terms of absolute value or relative values between different starting materials.
The original objective of the Jollands et al. (2020a) investigation was to provide diffusion coefficients that could be applied to Li-in/H-out profiles observed previously in some volcanic quartz crystals (Charlier et al., 2012; Tollan et al., 2019). Whilst it now appears that the simple diffusion model applied by Jollands et al. (2020a) was not strictly appropriate (we previously used equation 1), it is reasonable to assume that their phenomenological diffusion coefficients are appropriate for use as a diffusion chronometer, at least within their quoted uncertainties. This is because the compositional range of their starting materials (in terms of H and Li concentrations) straddled the composition of their studied volcanic quartz, from the Bishop Tuff. Regardless of our above discussion and the considerable complexities in this system, we would still propose that simple solutions or diffusion coefficients are appropriate for purposes of H diffusion chronometry using volcanic quartz in situations where H is lost and Li gained, at least within any reasonable uncertainty.

Conclusion

On the basis of a series of diffusion experiments and in situ analyses, this study demonstrates that H, $^3$H and Li diffusion in quartz involves a process where H and other monovalent cations move by dissociating from a charge-compensating species, generally Al$^{3+}$, then moving rapidly to another charge-balancing site, where they become immobilised. Models invoking this process can create different profile forms consistent with experimental observations: step shapes in H-in/Li-out experiments, localised humps in $^3$H-in/Li $^+$ out experiments, and error-function forms in Li-in/H-out and $^3$H-in/H-out experiments. Models describing this complex diffusion mechanism of H in quartz have been proposed previously (e.g. Kronenberg and Kirby, 1987) – this study provides independent support.

Supplementary material. All FTIR spectra and LA-ICP-MS data are available at https://doi.org/10.17632/v3p5fw65bv.1. To view the Supplementary Appendix for this article, please visit https://doi.org/10.1180/mgm.2021.105

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