Quantification of water in hydrous ringwoodite


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Ringwoodite, $\gamma-(\text{Mg,Fe})_2\text{SiO}_4$, in the lower 150 km of Earth's mantle transition zone (410–660 km depth) can incorporate up to 1.5–2 wt% H$_2$O as hydroxyl defects. We present a mineral-specific IR calibration for the absolute water content in hydrous ringwoodite by combining results from Raman spectroscopy, secondary ion mass spectrometry (SIMS) and proton-proton (pp)-scattering on a suite of synthetic Mg- and Fe-bearing hydrous ringwoodites. H$_2$O concentrations in the crystals studied here range from 0.46 to 1.7 wt% H$_2$O (absolute methods), with the maximum H$_2$O in the same sample giving 2.5 wt% by SIMS calibration. Anchoring our spectroscopic results to absolute H-atom concentrations from pp-scattering measurements, we report frequency-dependent integrated IR-absorption coefficients for water in ringwoodite ranging from 78,180 to 158,880 Lmol$^{-1}$cm$^{-2}$, depending upon frequency of the OH absorption. We further report a linear wavenumber IR calibration for H$_2$O quantification in hydrous ringwoodite across the Mg$_2$SiO$_4$-Fe$_2$SiO$_4$ solid solution, which will lead to more accurate estimations of the water content in both laboratory-grown and naturally occurring ringwoodites. Re-evaluation of the IR spectrum for a natural hydrous ringwoodite inclusion in diamond from the study of Pearson et al. (2014) indicates the crystal contains 1.43 ± 0.27 wt% H$_2$O, thus confirming near-maximum amounts of H$_2$O for this sample from the transition zone.

Keywords: IR spectroscopy, water in nominally anhydrous minerals, transition zone, mineral-specific absorption coefficient, SIMS, Raman spectroscopy, proton-proton scattering, ringwoodite

INTRODUCTION

Ringwoodite was first described in the Tenham meteorite found in Queensland, Australia, by Binns et al. (1969) and named after the Australian mineral physicist Ted Ringwood. The first terrestrial occurrence of ringwoodite was recently reported by Pearson et al. (2014), who discovered a hydrous ringwoodite inclusion in ultra-deep diamonds from Juina, Brazil. Ringwoodite is one of the major components of the Earth’s mantle transition zone (e.g., Bernal, 1936; Akaogi and Akimoto, 1977; Anderson and Bass, 1986; Bina and Wood, 1986; Irifune, 1987; Ringwood and Major, 1967) and although nominally anhydrous, ringwoodite is well-known to incorporate up to 1.5–2 wt% of water as (OH)$^-$ point defects in both laboratory-grown and naturally occurring samples (e.g., Kohlstedt et al., 1996; Bolfan-Casanova et al., 2000; Smyth et al., 2003; Pearson et al., 2014).

Water in nominally anhydrous minerals (NAMs) influences phase relations (e.g., Hirschmann, 2006), melting temperature (e.g., Hirth and Kohlstedt, 1996; Hirschmann et al., 2009; Tenner et al., 2012), and physical properties including electrical conductivity (e.g., Yoshino et al., 2009), thermal conductivity (Thomas et al., 2012), elasticity (e.g., Jacobsen, 2006), and rheology (e.g., Kavner, 2003). Several OH defect mechanisms have been studied in ringwoodite, mainly involving charge-compensating vacancies at the octahedral Mg-Fe sites, the tetrahedral Si site, and normally vacant interstitial sites (e.g., Kudoh et al., 2000; Smyth et al., 2003, 2004; Chamorro Pérez et al., 2006; Blanchard et al., 2009; Mrosko et al., 2013; Panero et al., 2013; Purejav et al., 2014; Yang et al., 2014).

Because of its likely abundance (>50% of a pyrolite model) in the lower 150 km of the Earth’s mantle transition zone (410–660 km depth) and for its ability to incorporate significant amounts of H$_2$O, we have undertaken this study on determining the absolute water content in ringwoodite. Infrared (IR) spectroscopy is one of the most common tools for analysis of water in minerals, for which mineral-specific absorption coefficients are required. The absolute water content in minerals was conventionally determined by quantitative dehydration weight analyses (e.g., Aines and Rossman, 1984), but more suitable for the typically very small amounts (μg) of material from high-pressure and high-temperature syntheses, two additional methods have been developed: elastic recoil detection analysis (e.g., Aubaud et al., 2009; Bureau et al., 2009; Withers et al., 2012) and proton-proton (pp) scattering (e.g., Gose et al., 2008; Thomas et al., 2008; Reichart and Dollinger, 2009).
In addition to theoretically based IR calibrations for water in ringwoodite (Balan et al., 2008; Blanchard et al., 2009), experimentally determined IR absorption coefficients have been estimated for ringwoodite either by using general calibrations (Paterson, 1982; Libowitzky and Rossman, 1997) or from secondary ion mass spectrometry (SIMS) measurements of water contents, which we note are usually calibrated against the water content in silicate glasses or minerals measured by IR. Absorption coefficients for water quantification have been determined for ringwoodites with fayalite component (Fa100, Fe2SiO4) through Fa40 and for pure forsterite (Fo100, Mg2SiO4) composition (Koch-Müller and Rhede, 2010) but not for compositions spanning the likely Fe-content of the mantle transition zone from Fo75 to Fo95.

In this study we calibrate the water concentrations in synthetic hydrous ringwoodite from IR spectroscopy against independently determined values from Raman spectroscopy, SIMS and the absolute H-concentration measured by pp-scattering microscopy. Broad-beam pp-scattering has previously been applied to mm-sized samples (Thomas et al., 2008). However, until recently this quantification technique was not feasible for smaller samples synthesized under very high pressure-temperature conditions in the multi-anvil press. Here, we employ pp-scattering experiments on synthetic ringwoodite crystals less than 200 μm in largest dimension and quantify 3D distributions of atomic hydrogen at μm spatial resolution. We also present hydrogen depth-profiles and 2D hydrogen maps and H2O concentrations in samples used for the spectroscopic calibration. We calculate absorption coefficients for a suite of hydrous ringwoodite with varying Fe- and water concentrations, test their wavenumber-dependence, and compare results with literature data. The IR calibration can thus be used to determine the absolute water content in hydrous ringwoodite, which we also apply to the natural hydrous ringwoodite inclusion in diamond reported by Pearson et al. (2014).

### MATERIALS AND METHODS

#### SAMPLES

Samples used in this study were synthesized at high P-T conditions in multi-anvil presses and cover a range of H2O contents and compositions (Table 1). Chemical analyses and other characterizations of the single crystals used in this study have been published elsewhere: for samples MA120, MA62, MA56, and MA75, see Koch-Müller and Rhede (2010), Taran et al. (2009), and Koch-Müller et al. (2009); Koch-Müller et al. (2011); for sample SZ0820, see Ye et al. (2012); for samples SZ9901 and SZ0104, see Smyth et al. (2003); Smyth et al. (2004), Thomas et al. (2012), and Jacobsen et al. (2004); for sample SZ0570, see Mao et al. (2011). High-quality, crack- and inclusion-free isotropic single crystals ranging in longest dimension from 50 to 500 μm were selected for analyses. The color of the ringwoodites used here varies from colorless to dark blue to almost opaque depending on Fe concentration.

#### 3D HYDROGEN MICROSCOPY BY PROTON-PROTON SCATTERING

Proton-proton scattering at proton energies up to 25 MeV has been used in a broad-beam configuration to perform hydrogen depth microscopy in mm-sized mineral platelets, as described...
in Thomas et al. (2008) or Gose et al. (2008). A new detector setup has been installed at the nuclear microprobe SNAKE at the Munich tandem accelerator lab that enables 3D hydrogen microscopy of μm-sized objects by scanning a μm-focused beam. A detailed description can be found in Reichart and Dollinger (2009). The protons scattered from an elastic scattering reaction of an incident proton and a hydrogen atom are detected in coincidence by two pairs of Si-detectors with 16 strips on the front and back side each. The strips and a narrow timing window allow filtering of hydrogen events from all other scattering events with a detection limit below 1 at-ppm (1 atom out of 10^6 other atoms). The energy information of both protons is used to gain depth information with about μm depth resolution. As both coincident protons from the pp-scattering reaction have to be detected at the back of the sample, the scattering geometry requires unsupported samples. With a proton energy of 22 MeV as used in our analyses, thin but stable samples can be prepared. Our samples were ground and polished until they had a final thickness <100 μm (SZ9901 39 μm, SZ0570 71 μm, SZ0820 50 μm). Crystal edges were glued to copper grids with mesh sizes ranging from 150 to 400 μm. From the scattering probability and the proton beam current, the number of coincident events from a certain thickness gives the atomic hydrogen in H-atoms/cm^2. We convert this to H atoms per total atoms using the atomic density of the samples determined by other methods such as X-ray crystallography.

**IR SPECTROSCOPY**

In preparation for IR measurements single crystals were doubly polished using diamond spray. Unpolarized IR spectra of randomly oriented synthetic ringwoodite were recorded from 1850 to 4000 cm^−1 at ambient conditions using a Bruker Tensor 37 FTIR spectrometer with a KBr beamsplitter, globar source, and Hyperion microscope with MCT detector. Up to 1024 scans were taken with a resolution of 2 cm^−1 through a 100 × 100 μm aperture. The sample thickness was determined using the eyepiece reticle and stage micrometer scale of the optical microscope. The final thicknesses varied from 13 to 84 μm depending on the specimen (Table 1). A linear baseline correction in the integration limits between 2000 and 3900 cm^−1 was applied for all our spectra. Peak positions, integrated absorbances, band maxima, and area-weighted average (Libowitzky and Rossman, 1996, 1997) of band positions were determined using PeakFit (Systat Software, Inc.). We applied a Gaussian/Lorentzian distribution function to all component bands. Polarized measurements are not needed for isotropic minerals, where the absorbance is linearly dependent on the species concentration and thickness of the sample. Here we use one-directional absorbance values derived from unpolarized spectra of cubic ringwoodite and multiply them by three for subsequent calculations.

**RAMAN SPECTROSCOPY**

The same single crystals used in the IR study were characterized with confocal microRaman spectroscopy with no additional sample preparation required. All analyses were performed in backscattering geometry using an Andor Shamrock 0.3 m spectrograph (1800 grooves/mm grating) coupled to an Andor Newton DU970 EM-CCD camera, an Olympus optical microscope and a long working distance Mitutoyo 100× objective (LWD VIS, NA = 0.70, WD = 6.0 mm). The 458-nm line of a solid-state Melles Griot laser source with ∼250 mW output power was used for sample excitation. Laser intensity at the sample was ∼18 mW. For all measurements a confocal aperture of 100 × 100 μm was used and spectra were acquired from 50 to 1300 cm^−1. Counting times ranged from 10 to 100 s with 3 accumulations. Counting times for spectra in the frequency range from 2800 to 4000 cm^−1 were 100 s at 5 accumulations. After a linear baseline correction was performed, spectra were integrated between 2000 and 3900 cm^−1. The KOG glass (Thomas et al., 2008) was used as reference material. A molar volume correction factor of 0.79 for differences between sample and reference material was used for quantification (see Mrosko et al., 2011).

**SECONDARY-ION MASS SPECTROMETRY**

SIMS measurements of hydrogen in ringwoodite were carried out on the Cameca NanoSIMS 50L scanning ion microscope at the Carnegie Institution of Washington. Polished single crystals were mounted in indium. To flatten the surface and fill cavities around the crystals, the whole assembly was pressed carefully with 4.9 t in a hydraulic hand press. The ^16OH^- signal was standardized using natural and synthetic minerals described in Koga et al. (2003) and Hauri et al. (2006): olivines SynFo100, SynFo68, GRR1012, KLV-23; orthopyroxenes KMB-1, India, Kenya, A288; garnets MON-9, ROM263-9, ROM263-25, ROM263-52; clinopyroxene PMR-53. We also used additional olivines and pyroxenes from the Monastery kimberlite that were studied by Bell et al. (2004): olivines ROM177, ROM250-13, ROM250-2; clinopyroxenes ROM271-10, ROM271-16, ROM271-21; orthopyroxene ROM273. Detailed analytical setup and methodology information is described in Hauri et al. (2006) and Koga et al. (2003).

**RESULTS**

**IR SPECTROSCOPY**

Unpolarized IR spectra of ringwoodite with forsterite numbers Fo100-Fo83 are illustrated in Figure 1. Spectra are offset for clarity. The IR spectra show broad absorption bands of OH stretching vibrations with maxima of the main band ranging from ∼3130 to 3174 cm^−1 and additional bands at ∼3656 to 3675 cm^−1, 3531 to 3568 cm^−1, and ∼2500 cm^−1 (see Table 1 and Figure 1). Band intensities vary and band maxima of the main band shift to higher wavenumbers with increasing iron content.

**RAMAN SPECTROSCOPY**

Unpolarized Raman spectra for ringwoodites used in this study are shown in Figure 2. The spectrum of pure Mg-ringwoodite (Figure 2A) displays expected Raman modes at ∼322, 444, 665, 794, and 829 cm^−1, which are due to antisymmetric (T2g) and symmetric (A1g) stretching vibrations of the isolated SiO4 tetrahedra (Chopelas et al., 1994), whereas an association of the Raman bands at ∼300 cm^−1 and 400 cm^−1 with octahedral cation vibrations is under discussion (McMillan and Akaogi, 1987; Chopelas et al., 1994). Representative spectra for more
iron-rich samples show similar patterns (Figure 2B), with the strongest Raman peaks at ∼794 and 835 cm⁻¹ dominating. In addition, low-frequency bands with slightly varying intensities and positions at ∼200 cm⁻¹ are present in the Fe-bearing samples (Figure 2B). These signals might be associated with localized modes generated by Fe substitution (Fe²⁺, Fe³⁺) (Kleppe et al., 2002; Kleppe and Jephcoat, 2006), which explains their absence in pure Mg-ringwoodite (Figures 2A,B). Additional weak modes at >700 cm⁻¹ might be associated with vibrations due to Si₂O₇ (Kleppe and Jephcoat, 2006). The high-frequency range (Figures 2A,D) of all samples studied here shows broad OH band features due to OH stretching vibrations in accordance with observations from IR spectroscopy, which allow water quantification by Raman spectroscopy. Figure 2C shows the Raman spectrum of the reference silica glass used in this study containing 332 wt ppm H₂O (Thomas et al., 2008). Note, that Figure 2B is meant for illustrative purposes only; spectra have not been normalized and intensities are given in arbitrary units representing measurements taken on different days with different count rates, laser intensities and accumulations, allowing only qualitative comparison.

FIGURE 1 | Unpolarized IR spectra (black) including band deconvolution (red) of Fe-free (Fo100) and Fe-bearing hydrous ringwoodites at room pressure. Spectra show broad absorption features with band maxima ranging from ∼3130 to 3174 cm⁻¹, from ∼3531 to 3588 cm⁻¹, and from ∼3656 to 3675 cm⁻¹. The feature at ∼2500 cm⁻¹ has been assigned to overtones by e.g., Hofmeister and Mao (2001) and was excluded from further quantification calculations. Spectra are offset for clarity.

FIGURE 2 | Raman spectra of Mg-ringwoodite (A) and Fe-bearing ringwoodite (B) showing characteristic lattice vibrations in the low-frequency range and OH bands in the high-frequency range (D). The spectrum of the reference glass KOG with 332 wt ppm is shown in (C). Intensities are given in arbitrary units.

WATER CONTENTS AND MINERAL-SPECIFIC ABSORPTION COEFFICIENTS

Ringwoodite water contents determined by confocal micro-Raman spectroscopy, SIMS, and pp-scattering (Table 1) are in good agreement. Based on SIMS analysis, the H₂O concentration of the five samples studied here ranges from 0.36 wt% H₂O to 2.5 wt% H₂O (see Table 1). Figure 3 shows hydrogen maps and depth profiles from selected areas derived from pp-scattering analyses for μm-sized crystals of Fo100, 90, and 83 super-glued on copper grids. Depth profiles allow separation of surface water and internal bulk hydrogen (or glue contamination, see Figure 3A), measured as H atoms per square cm. Assuming density and composition, the depth scale and thickness is calculated from energy loss of the protons, giving H atoms per total atoms in the selected areas. Background-corrected values for Fo100, 90, and 83 were calculated as 1.77 ± 0.35 wt% H₂O, 1.13 ± 0.23 wt% H₂O, and 0.91 ± 0.18 wt% H₂O, respectively, assuming all detected H is incorporated as hydroxyl groups.

From independently determined water concentrations (cH₂O), peak-fitted integrated areas (Aᵢ) from IR spectroscopy, sample
FIGURE 3 | Sample images, hydrogen maps, and depth profiles of ringwoodites SZ0820 (Fo100, A), SZ0570 (Fo83, B), and SZ9901 (Fo90, C) from pp-scattering. The total scan area is marked in the sample images. From the hydrogen maps we are able to select areas of interest for quantitative depth profiles (B, C in log scale for better background visibility). Outside areas may be covered by the supporting mesh grid or contaminated by glue residue, as clearly observed in the depth profile from (Continued)
thickness (d) and sample density (ρ), mineral-specific IR absorption coefficients (Table 2) were calculated for Fo60, 83, 87, 90, and 100 compositions using the Beer-Lambert law, ε = (A_t × 1.8)/(d × ρ × c_H2O), where d is the sample thickness in cm, ρ is the sample density in g/cm³, and c_H2O is the H2O content in wt%. Note that we use the integrated absorbance from one crystallographic direction multiplied by three for our calculations. Based on Raman spectroscopy, absorption coefficients for ringwoodite of Fo60 to Fo100 composition range from 78,180 ± 23,460 Lmol^-1 cm^-2 to 158,880 ± 47,660 Lmol^-1 cm^-2 (Table 2). The relative error of absorption coefficients is mainly determined by the error of the water concentration from all analyses (7–22% relative uncertainty from SIMS, 20% relative uncertainty from pp-scattering and 30% relative uncertainty from Raman spectroscopy), the surface quality of the samples, and spectral fits.

### DISCUSSION

IR spectroscopy allows investigation of water incorporation mechanisms in NAMs (e.g., Libowitzky and Beran, 2006). For ringwoodite, mechanisms include protonation of oxygen sites (hydroxyl, OH⁻) associated with the vacant and partially vacant 16c and 16d octahedral sites, [V_Mg(OH)_2]^x, Mg²⁺ substitution for Si⁴⁺ on the tetrahedral site, [Mg₆(OH)_2]^{12}x, tetrahedral silicon vacancies with a hydrogarnet type defect, [V_Si(OH)_4]^{12}x, and defects on tetrahedral edges, [Mg/Fe²⁺/Fe³⁺(OH)_2]^{12}x (e.g., Kudoh et al., 2000; Smyth et al., 2003; Blanchard et al., 2009; Mrosko et al., 2013; Panero et al., 2013; Purevjav et al., 2014; Yang et al., 2014). IR spectra of synthetic ringwoodite reported are similar to previous studies (e.g., Bolfan-Casanova et al., 2000; Kudoh et al., 2000; Smyth et al., 2003; Chamorro Pérez et al., 2006; Mrosko et al., 2013).

Following the band assignments of Mrosko et al. (2013), based on heating experiments, and Hofmeister and Mao (2001), the main OH band at ∼3130 cm⁻¹ is most likely composed of various bands and associated with protonation of octahedral vacancies; the bands at 3656 to 3675 cm⁻¹ and 3531 to 3568 cm⁻¹ are associated with tetrahedral defects of either the hydrogarnet substitution or Mg replacing Si atoms, and the features at ∼2500 cm⁻¹, present in anhydrous and hydrous ringwoodite are due to overtones of Si-O vibrations. Contingent upon the increasing Fe-concentration in our samples we observe the expected shift of the main OH band to higher wavenumbers (e.g., Libowitzky, 1999; Mrosko et al., 2013), from ∼3130 cm⁻¹ in Fo100 to 3174 cm⁻¹ in Fo83. Mrosko et al. (2013) report that an increasing iron concentration also affects the high-frequency band at ∼3675 cm⁻¹, which in this case causes a shift to lower wavenumbers. Here, we do not observe a significant negative shift of this band. This might be due to the relatively low iron content and the low Fe-contrast between our samples (from Fo100 to Fo83), hence such a shift might be resolvable only when comparing samples with more extreme differences in total Fe-concentration. We also do not observe a direct correlation between sample Fe-content and water concentrations for our synthetic ringwoodites. The latter are consistent with prior reported values, which typically range up to 2.5 wt% H₂O and 1.48 wt% in synthetic and natural samples, respectively (Kohlstedt et al., 1996; Smyth et al., 2003; Pearson et al., 2014; Yang et al., 2014). In agreement with Koch-Müller and Rhede (2010), we observe a general increase in the integrated molar absorption coefficient with decreasing OH-stretching frequency of the band maximum (Table 2 and Figure 5).

A comparison of ringwoodite-specific IR absorption coefficients derived from water quantification using a variety of analytical techniques is shown in Figure 5. The two solid lines give the general trends of absorption coefficients for water in hydrous minerals and glasses vs. wavenumber of the corresponding OH bands (Paterson, 1982; Libowitzky and Rossmann, 1997). Since more than one peakfit solution exists for the broad OH band pattern in ringwoodite, we chose to plot band maxima vs. absorption coefficient instead of weighted mean wavenumbers, as suggested in Libowitzky and Rossmann (1997). The good correlation between absorption coefficient and band maxima reflects the dominance of the main OH feature in the IR spectra, i.e., the prevalence of the associated incorporation mechanism. For ringwoodite we observe a frequency-dependence of the absorption coefficient, which shows an increase with decreasing wavenumber due to interrelations of OH dipole strength, O-H…O geometry and OH band intensity (for details see Koch-Müller and Rhede, 2010), similar to what has been communicated by Paterson (1982) and Libowitzky and Rossmann (1997) (Figure 5).

Our observations support conclusions of Koch-Müller and Rhede (2010) that the use of a single absorption coefficient for water quantification in ringwoodite is not recommended. Instead, a frequency-dependent calibration specific to ringwoodite is needed, which reflects a range of OH-O distances connected to variance in sample composition, i.e., element concentrations, densities, and molar volumes. From Raman spectroscopy, we determine absorption coefficients ranging from 78,180 ± 23,460 Lmol⁻¹ cm⁻² to 158,880 ± 47,660 Lmol⁻¹ cm⁻² for ringwoodite samples of Fo60 to Fo100 composition (see Table 2 and Figure 4). All values (except the Raman value for Fo87) for our ringwoodite plot below the calibration curves of Paterson (1982) and Libowitzky and Rossmann (1997), which is in good agreement with findings by Balan et al. (2008), Blanchard et al. (2009), and Koch-Müller and Rhede (2010). Koch-Müller and Rhede (2010) report wavenumber-dependent absorption coefficients for synthetic Fe-Mg-ringwoodites with Fa100 to Fo60 compositions. Based on their experimentally determined wavenumber dependence and density/molar volume dependence of the calibration constant they estimate absorption coefficients of 101,613 Lmol⁻¹ cm⁻² and 105,877 Lmol⁻¹ cm⁻².
### Table 2 | Comparison of absorption coefficients for ringwoodite from this study and literature.

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<th>Experiments</th>
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<th>Absorption coefficient (L/molH₂Ocm²) (Raman this study)</th>
<th>Absorption coefficient (L/molH₂Ocm²) (SIMS this study)</th>
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For Mg-ringwoodite, respectively, with an OH band maximum of 3120 cm⁻¹. These values are in good agreement with our experimentally determined average absorption coefficient for Mg-ringwoodite from SIMS, Raman, and pp-scattering of 98,280 Lmol⁻¹cm⁻² (Table 2) with a slightly increased band maximum of 3130 cm⁻¹.

From our absorption coefficients based on OH band maxima (Figure 5) and data points obtained by Koch-Müller and Rhede (2010) we calculate a regression line, which can be used to determine IR absorption coefficients for ringwoodites: \( y = -207.35x + 761,228 \), where \( y \) is the absorption coefficient in Lmol⁻¹cm⁻² and \( x \) is the band maximum of the strongest OH band in cm⁻¹. Based on this linear regression the value for Mg-ringwoodite with a band maximum of 3130 cm⁻¹ calculates as 112,223 Lmol⁻¹cm⁻² (Table 2) and for a band maximum of 3120 cm⁻¹ as 114,296 Lmol⁻¹cm⁻². Koch-Müller and Rhede (2010) conclude that wavenumber-independent absorption coefficients, such as those for olivine and quartz (Thomas et al., 2009), are valid for minerals of the same composition with comparable atomic distances. Absorption coefficients, however, strongly depend on structure changes. This has been shown for SiO₂ polymorphs, where coefficients increase from quartz to coesite and stishovite, as the densities of the structures and thus the mean O-O distances decrease (Thomas et al., 2009; Koch-Müller and Rhede, 2010). Koch-Müller and Rhede (2010) explain the frequency dependence of the absorption coefficient in ringwoodite by strongly differing mineral compositions (Fe/Mg substitution), which may result in a continuous decrease of the O-H—O distances with increasing Mg. They find that within the polymorphic series of \( (\text{Mg,Fe})_2\text{SiO}_4 \) the absorption coefficient increases with decreasing density and molar volume, which we do not observe for the synthetic ringwoodites of Fo100 to Fo83 composition (Figure 4).

Our work illustrates that using general IR calibration trends reported by Paterson (1982) and Libowitzky and Rossman (1997) would underestimate water concentrations in ringwoodite as previously discussed by Koch-Müller and Rhede (2010). Our absorption coefficients give up to 34% higher H₂O contents than the general calibration trends would estimate. We have produced a linear wavenumber-dependent IR calibration for water quantification in a ringwoodite series representing Fₐ to Fₒ compositions, which will result in more accurate estimations of the water content in natural and synthetic ringwoodites.

Finally, using this new calibration for water in ringwoodite we re-evaluated the OH-absorption spectrum of the natural hydrous ringwoodite diamond inclusion from Pearson et al. (2014). Note that Pearson et al. (2014) utilized absorption coefficient and density estimates for pure Mg-ringwoodite (3.9 g/cm³). This leads to a large uncertainty, which was estimated to be ±50% by Pearson et al. (2014). The combination of IR spectra (OH band maxima) and density data from a suite of synthetic ringwoodite samples studied here (Table 1) allows a better density estimate of ∼3.65 g/cm³ for their ringwoodite inclusion and indicates a forsterite number of ∼90. Usage of the new density and a more suitable absorption coefficient (106,002 Lmol⁻¹cm⁻²) for the OH band maximum at 3160 cm⁻¹ results in a water content of 1.36–1.50 wt% H₂O in their natural sample. While the
new wavenumber-dependent calibration confirms originally projected H₂O concentrations (1.35–1.49 wt% H₂O), it reduces the uncertainty by more than half, to ~20%. Our new estimate for the uncertainty in the water content of the natural hydrous ringwoodite (Pearson et al., 2014) includes a Gaussian error propagation of relative errors from peak-fitted integrated areas (5%), sample density (5%), sample thickness (15%), and absorption coefficient (10%). With an absolute H₂O content of 1.43(±0.27) wt%, we thus confirm that the natural inclusion from the transition zone contains near-maximum amounts of H₂O.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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