In situ high-temperature Raman and FTIR spectroscopy of the phase transformation of lizardite

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ABSTRACT

A temperature-dependent in situ micro-FTIR and micro-Raman spectroscopic investigation was performed on powdered (FTIR, Raman) and single-crystal (Raman) lizardite-1T samples between room temperature and 819 °C. Between room temperature and 665 °C, the OH stretching bands shift to lower wavenumbers, demonstrating a weak expansion of the O3-H3…O2 interlayer distance. Band deconvolution of FTIR and Raman spectra at room temperature show differences in the number of bands in the OH stretching region with respect to group theory: four (FTIR) and six (Raman) OH stretching bands, respectively. This number reduces to four (Raman) at non-ambient temperatures either caused by LO-TO splitting, the presence of non-structural OH species or the presence of different lizardite polytypes and/or serpentine polymorphs as well as defects. During dehydroxylation, the evolution of the integrated intensity of the OH bands suggests a transport of hydrogen and oxygen as individual ions/molecule or OH⁻. A significant change in Raman spectra occurs between 639 and 665 °C with most lizardite peaks disappearing contemporaneously with the appearance of forsterite-related features and new, non-forsterite bands at 183, 350, and 670 cm⁻¹. A further band appears at 1000 cm⁻¹ at 690 °C. The long stability of Si-O-related bands indicates a delayed decomposition of the tetrahedral sheet with respect to the dehydroxylation of the octahedral sheet. Moreover, evidence for a small change in the ditrigonal distortion angle during heating is given. In general, all appearing non-forsterite-related frequencies are similar to Raman data of talc and this indicates the presence of a talc-like intermediate.

Keywords: Lizardite, dehydroxylation, Raman, and FTIR spectroscopy, talc-like intermediate

INTRODUCTION

Lizardite Mg3Si2O5(OH)4 is a member of the serpentine group. The 1:1 phyllosilicate consists of brucite-like octahedral sheets connected to SiO₄ tetrahedral sheets. The most common structures are the one-layer 1T and the two-layer 2Hₐ polytypes, although other polytypes are known. At ambient pressure and temperatures above 500 °C, lizardite breaks down to anhydrous phases, i.e., forsterite and enstatite. A probable mechanism for the liberation of the hydroxyl groups is a reaction of two adjacent hydroxyl groups that presumably recombine to a water molecule (e.g., Guggenheim et al. 1987). Theoretical studies have shown that the two symmetrically distinguishable hydroxyl groups (Fig. 1) in each lizardite polytype can be recognized by their infrared (IR) and Raman active bands in the frequency range between 3500 and 3800 cm⁻¹ (Balan et al. 2002; Auzende et al. 2004). Temperature-dependent FTIR and Raman studies are therefore suitable to study the structural change of serpentine minerals, which is difficult to understand by X-ray techniques (XRD) alone (e.g., Guggenheim and Zhan 1998). In general, temperature-dependent peak shifts or intensity/absorbance changes are caused by changes in the bond length, symmetry changes, or short- to large-scaled structural rearrangements (e.g., Zhang et al. 2006; Gillet 1996). Intensity changes of the OH Raman bands and changes in absorbance of the OH IR bands can serve as a (semi-)quantitative information source to estimate the amount of OH or H₂O within the crystal structure. Previous Raman and FTIR studies on serpentine minerals were mainly used as tools to distinguish serpentine polymorphs (e.g., Rinaudo et al. 2003) or to determine the structural behavior at high-pressure/high-temperature (HP/HT) conditions within the stability region of

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FIGURE 1. Crystal structure of lizardite-1T (Mellini 1982) projected along the b axis showing the positions of the inner and inner-surface hydroxyl groups. Some authors, e.g., Auzende et al. (2004), designate the “inner-surface” OH also as “outer” OH.
EXPERIMENTAL METHODS

Sample material

The investigated lizardite sample, for which no origin is given, was taken from the ETH Zurich mineral collection. The electron microprobe-based determination of the stoichiometry yields $(\text{Mg}_{2.84}\text{Fe}_{0.04}\text{Al}_{0.03})\text{(Si}_{2.02}\text{)}\text{O}_{5\text{(OH)}}_{4.09}$ and a total water content of 13.3 wt%, which was estimated by thermogravimetry. The sample mainly consists of the 1:1 polytype, although there are intergrowths of curled 1:1-layers and regions with a high concentration of stacking faults based on transmission electron microscopy (TEM) analyses (Trittschack and Grobéty 2012).

As reference, a lizardite-1 single crystal from the Monte Fico quarries, Elba Island/Italy, was used (private collection, Marcello Mellini). Comprehensive mineralogical and chemical data of this sample are provided by Mellini and Viti (1994) and Fuchs et al. (1998).

Powdered lizardite was gently pressed in a diamond-anvil cell (DAC) without gasket until a transparent film was generated. Single transparent pieces of this film were deposited in a sapphire crucible and used for all FTIR in situ dehydroxylation experiments. Raman experiments were performed on powdered samples and a Monte Fico single crystal, placed into a sapphire crucible.

FTIR and Raman spectroscopy

FTIR measurements were performed with a Bruker VERTEX 80v spectrometer equipped with an Olympus BX41 microscope (20× objective) in backscattering configuration at the GFZ Potsdam. All examined spectra were measured with an argon laser (488 nm, 300 mW). The spectrometer was calibrated against the emission bands of a neon lamp. Spectra acquisition time was 10 s and 4 spectra were cumulated for each temperature step. In general, spectra were measured in the frequency ranges of 100–1200 and 3500–3800 cm$^{-1}$.

Non-isothermal runs were monitored to study the dehydroxylation of lizardite. The Raman measurements were obtained between 21 and 819 °C. An initial heating rate of 130 °C/min was used to heat the sample from 21 to 253 °C, afterward the heating rate was fixed to 100 °C/min for 50 °C increments (to 459 °C) and 50 °C/min for 25 °C increments, starting with the 484 °C spectrum and ending with the 819 °C spectrum. Single measurements after each temperature increment required 5 min owing to the Raman setup conditions. The FTIR data were collected between 21 and 665 °C. Initially, a heating rate of 100 °C/min was used for the following temperature intervals: 21–98, 98–201, 201–304, 304–356, 356–407, 407–433, and 433–459 °C. FTIR spectra were recorded after each temperature interval. A linear heating rate of 1 °C/min and a continuous measurement was chosen from 459 to 665 °C.

Fitting procedure

Peak deconvolution and integration were performed with the PeakFit software package. The low-frequency range (100–1200 cm$^{-1}$) and high-frequency range (3500–3800 cm$^{-1}$) were treated independently. The Savitzky-Golay algorithm was used to smooth single spectra. After baseline subtraction, all spectra were deconvoluted using a combined Gaussian-Lorentzian Amp function for which the width at the base and the full-width at half maximum (FWHM) are refinable parameters. A minimum number of peaks was chosen to guarantee a best fit to allow the comparison between individual time and temperature resolved spectra as shown by Auzende et al. (2004).

RESULTS

Raman and FTIR spectra at ambient conditions

The low-frequency range of the Raman spectra obtained at RT is characterized by five well-resolved strong bands that are accompanied by six weak bands (Fig. 2). Additional bands (Table 1) observable in the deconvoluted lizardite-17 spectrum of the Monte Fico sample are very weak or not present in spectra obtained from the powdered sample (Fig. 2). Precise peak positions and band assignments of both lizardite samples are summarized in Table 1. In general, the positions of the main peaks in the low-frequency range are identical with data summarized by Rinaudo et al. (2003). However, some bands were assigned to chrysotile by the latter (Table 1). We cannot unambiguously distinguish between purely lizardite-related or chrysotile containing, because these bands occur in the Monte Fico lizardite sample also, which contains up to 25 wt% chrysotile (Viti 2010). The 510 cm$^{-1}$ band assigned to deformation vibrations of SiO$_4$-AlO$_4$ tetrahedra is visible in the spectrum of the Monte Fico crystal.
but is missing in the powdered sample due to the low-AI content of our sample. The antisymmetric Si-O stretching frequencies centered at around 1072 and 1099 cm\(^{-1}\) are present but an accurate fitting in the in situ recorded spectra is difficult owing to their low intensity and large FWHM (Fig. 2).

For a deconvolution of the overlapping OH stretching frequencies at RT, four peaks were used for the FTIR spectra (Fig. 3a) and six for the related Raman spectra (Fig. 3b). Six bands were required also to fit OH bands in the Raman spectrum obtained from the Monte Fico single crystal (Table 1). This amount of bands is not compatible with theoretical data of the serpentine polymorph chrysotile by Kloprogge et al. (1999), who indicated the presence of just four Raman active OH stretching modes of the inner OH groups (3688 cm\(^{-1}\)) and the stretching modes of the inner surface OH groups (3688 cm\(^{-1}\)) and the stretching modes of the inner OH groups (3703 cm\(^{-1}\)). The fourth deconvoluted band at 3669 cm\(^{-1}\) (RT) is not indicated among the 51 transverse optical modes of the inner-surface OH groups and the stretching vibrations of the inner OH groups show a linear temperature relationship, whereas two linear segments are necessary to describe the

### Temperature-dependent Raman- and FTIR-spectra in the OH stretching region

At temperatures higher than RT, the number of peaks necessary to deconvolute the OH bands adequately is four, both for the Raman (Fig. 7) and IR spectra (Fig. 8). There are no additional OH-related bands visible during heating. The frequency shifts of the four bands attributed to OH are illustrated in Figure 9 (FTIR data) and Figure 10 (Raman data) as a function of temperature. Following Balan et al. (2002), three bands (Fig. 3) are attributed to the out-of-phase stretching modes of the inner-surface OH groups (3652 cm\(^{-1}\)), the in-phase stretching modes of the inner-surface OH groups (3688 cm\(^{-1}\)) and the stretching modes of the inner OH groups (3703 cm\(^{-1}\)). The fourth deconvoluted band at 3669 cm\(^{-1}\) (RT) is not indicated among the 51 transverse optical modes indicated by Balan et al. (2002). The in-phase stretching modes of the inner-surface OH groups and the stretching vibrations of the inner OH groups show a linear temperature relationship, whereas two linear segments are necessary to describe the

### Table 1. Lizardite-17 related Raman and FTIR bands and their assignments at room temperature

<table>
<thead>
<tr>
<th>Band assignment(^{\dagger})</th>
<th>Lizardite-17 Raman bands (cm(^{-1}))</th>
<th>Lizardite-17 Raman bands (cm(^{-1}))</th>
<th>Lizardite-17 FTIR bands (cm(^{-1}))</th>
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**Notes:** as = antisymmetric, s = symmetric, chr = attributed to chrysotile vibrations by Rinaudo et al. (2003); (1) Indicates the five major Raman bands of lizardite in the low-frequency range.

\(^{\dagger}\) Rinaudo et al. (2003).

\(^{\dagger\dagger}\) Farmer (1974).

\(^{\ddagger}\) Sample described by Mellini and Viti (1994), but measured individually for this study.
thermal frequency evolution of the out-of-phase vibration of the inner-surface OH groups. The temperature at the intersection of the segments is indicated by an arrow in Figure 9a. This temperature coincides approximately with the temperature of an accelerated dehydroxylation behavior (Fig. 11a). The unassigned band shows a similar non-linear frequency evolution (Fig. 9b).
Libowitzky (1999) showed that the OH stretching frequencies can be used to calculate O…O and H…O distances in hydrous minerals, i.e., the lizardite interlayer distances (for comparison see Guggenheim and Zhan 1998) are determined from the OH stretching frequencies. Using this method, the interlayer distance (O2-O3 distance) derived from the FTIR data increases from 2.89 ± 0.06 Å at RT to 2.90 ± 0.06 Å at 582 °C. These distances correspond to an O2–H3 distance of 2.23 ± 0.01 Å at RT to 2.27 ± 0.01 Å at 582 °C.

The evolution of the integrated absorbance with increasing temperature of the four OH bands extracted from the in situ FTIR experiments results in a sigmoidal shaped dehydroxylation curve. Initial dehydroxylation starts at 450 ± 20 °C. The dehydroxylation maximum lies at 540 °C (Fig. 11a). The absorbance of the individual inner-surface and inner OH stretching modes also starts to decrease at around 450 °C (Fig. 11b). The average intensity of the unassigned band is picking up with beginning of the dehydroxylation reaction.

**FIGURE 5.** (a–f) Low-frequency range of temperature-dependent Raman measurements representing the phase transformation of lizardite-17 to forsterite; (f) sample was heated progressively to 819 °C and afterward quenched and re-measured at RT (21 °C).
The sum curve of the four Raman OH stretching band intensities (including the unassigned band, for comparison see Figs. 3, 7, and 8), which occur at temperatures higher than RT, gently decreases between RT (Fig. 12a) and 450 °C. The minimum of the derivative, functionally related to the maximum in the OH release, occurs at around 640 °C. Above 665 °C the bands disappear completely. The inner-surface OH stretching bands start to decrease in intensity at around 550 to 600 °C (Fig. 12b). Similar to the behavior observed in the FTIR spectra, the unassigned band has on average a higher intensity during the dehydroxylation than at lower temperatures.

**DISCUSSION**

**OH bands and their assignments**

 Whereas RT low-frequency Raman spectra of lizardite are compatible with previous data (e.g., Rinaudo et al. 2003), OH stretching frequencies in both the Raman and FTIR data show unusual band multiplication compared to group theory considerations by Balan et al. (2002) and Prencipe et al. (2009). Additional bands to fit the spectrum for the OH stretching modes of lizardite at RT has been recognized previously (e.g., Auzende et al. 2004) and additional OH bands are also known from spectra of 2:1 phyllosilicates (Zhang et al. 2005). Three different causes are most likely responsible for the appearance of the extra OH band: (1) longitudinal optic-transverse optic (LO-TO) splitting, (2) additional OH groups such as non-structural OH groups, also related to disorder, and (3) mixture of lizardite polytypes and/or chrysotile-like layers. According to hybrid Hartree-Fock/Density functional theory (HF/DFT) calculations of the vibrational spectrum of lizardite, Prencipe et al. (2009) interpreted additional bands observed by Auzende et al. (2004) as originating from LO-TO splitting of the three OH vibrational modes. Additional OH
bands in Raman spectra of kaolinite are also attributed to LO-TO splitting (Farmer 1998). Prencipe et al. (2009) and Farmer (1998) comprehensively presented the physical background of such additional band appearances related to LO-TO splittings. The key parameter for LO-TO splitting is the size of individual crystals compared to the Raman laser wavelength (Farmer 1998). IR data contain mostly TO modes, except for powdered samples with crystallite sizes greater than the IR wavelength, whereas Raman data are frequently affected by LO-TO splitting (Farmer 1998). Hence, a wavelength depending behavior, i.e., band multiplication owing to sample size related parameters with respect to the IR and Raman laser wavelength, may result in additional bands in our RT FTIR and Raman data sets, with four observable IR bands, but six Raman bands. Thus, the transparent film used probably shows a less complex OH stretching band pattern (dominated by TO modes) than the resulting patterns of the single crystal and the loose powder used in the Raman experiments.

Non-structural OH species have been suggested as an alternative explanation for additional OH band(s) in vibrational spectra of phyllosilicates (Wang et al. 2002; Zhang et al. 2006, 2010). However, most of these bands appear during the heat treatment and before significant dehydroxylation (e.g., Zhang et al. 2010). These bands are often situated at wavenumbers significantly different from that of initial OH bands. In the case of talc, such extra bands occur within the range of the primary OH bands (Zhang et al. 2006). In our experiments, the band at 3669 cm\(^{-1}\) (FTIR) and 3670 cm\(^{-1}\) (Raman), respectively, labeled as the unassigned band in Figure 3, can be interpreted as resulting from some non-structural OH species already present at RT, but increasing during dehydroxylation. A third possible cause for the additional bands may be the presence of different polytypes, turbostratic stacking faults or curved, chrysotile-like layers within the lizardite sample (Trittschack and Grobéty 2012). Defects have been suggested as an explanation for additional hydroxyl bands in amphiboles (Burns and Strens 1966), micas (Beran 2002), and kaolinite and dickite (Brindley et al. 1986). For samples containing more than one polymorph of a phase, individual frequency shift rates (\(\delta v/\delta T\)) for specific modes of individual polymorphs may change the degree of overlap. This changes the number of bands necessary for an accurate fit of the measured spectra. Brindley et al. (1986) document such temperature-dependent OH band specific shifts in kaolinite and dickite. However, taking the possibility of a LO-TO splitting into account, such a solution is simplistic.

**Figure 7.** (a–d) OH stretching frequency range of temperature-dependent Raman measurements representing significant intensity reduction between RT and 665 °C, the highest temperature where OH signals are still present.
The lizardite dehydroxylation I: Comparison between FTIR, Raman, and TGA data

The temperature at which the absorbance of the OH bands starts to decrease in the FTIR spectra is comparable with the dehydroxylation onset observed in TGA analysis of the same sample (Trittschack and Grobéty 2012). The rate of the intensity decrease (Fig. 11) is, however, more rapid than the TGA derived dehydroxylation rates in experiments with comparable heating rate (540 °C compared to ~650 °C \( \leq T_{\text{max}} \leq ~720 °C \)) reported by Trittschack and Grobéty (2012). The explanation for the discrepancy may be that the two techniques monitor different steps of the dehydroxylation process. Whereas FTIR/Raman spectroscopy is sensitive to the primary hydrogen and hydroxyl liberation steps, TGA monitors the exit of the dehydroxylation products from the sample e.g., the weight loss after the products leaves the reactant grain and the powder. Thus, the TGA signal is initially delayed relative to the spectroscopic signal. The differences in dehydroxylation maxima observed in FTIR and Raman data are related to the different samples used. Also, the preparation of the samples used for the FTIR and Raman experiments were different: whereas for the Raman spectroscopy a gently pressed powder and a single crystal were used, the FTIR spectroscopy was obtained on a thin transparent film pressed in a DAC. The partial pressure (or activity) of the product species (water, hydroxyl, hydrogen, and/or oxygen molecules) along the diffusion path decreases more rapidly in a thin film than in a loose powder and single crystal, respectively. This increases the absolute dehydroxylation reaction rate in the thin film, which may explain the accelerated intensity loss observed in the FTIR measurements. A similar explanation has been suggested for the differences in IR spectra between annealed pyrophyllite samples of a varying thickness (Wang et al. 2002).

The lizardite dehydroxylation II: Kinetic aspects and their spectroscopic indications

The activation energy \( E_A \) as function of the reaction progress estimated by TGA based dehydroxylation studies of lizardite shows an initial increase from 200 to over 350 kJ/mol followed by a plateau (Trittschack and Grobéty 2012). For heating rates similar to the one used here, the activation energy plateau is reached at about 50 °C below the temperature at which the OH bands have lost almost 60% (at 639 °C), respectively, 98% (665 °C)

FIGURE 8. (a–d) OH stretching frequency range of temperature-dependent FTIR measurements representing a significant reduction in absorbance between RT and 562 °C.
of their intensity in the Raman spectrum. DFT calculations and almost all previous works (e.g., Gualtieri and Ferrari 2006; Molina-Montes et al. 2008a, 2008b; Sperinck et al. 2011) propose a recombination of neighboring proton and hydroxyl ions to H₂O molecules as the primary reaction step, followed by diffusion of the products (H₂O molecules) along the interlayer to the surface. Several distinct recombination variants are possible, which differ by the distance between the reacting hydroxyls. Trittschack and Grobéty (2012) interpreted the increasing $E_a$ as a result of these parallel recombination steps, whereby the close distance, low $E_a$ recombination reactions occur at the beginning of dehydroxylation. With increasing reaction progress, longer distance recombination variants with higher $E_a$ become more frequent. The plateau $E_a$ corresponds to the recombination variant with the highest $E_a$ and the slight decrease indicates the increasing influence of transport in certain parts of the sample (large grains). This interpretation is compatible with DFT calculations (Molina-Montes et al. 2008a, 2008b; Sperinck et al. 2011), but not with experimental work by Gualtieri and Ferrari (2006) and Gualtieri et al. (2012), who indicate one-dimensional diffusion of H₂O molecules through the six-membered silicate rings. In the latter case, H₂O-related modes should be visible in in situ Raman and FTIR spectra of partially dehydrated samples. Martin (1977) and Datta (1991) described a broad, infrared active band between 3200 and 3500 cm⁻¹ in thermally pretreated chrysotile samples, which they assign to H₂O modes. Such bands were not present in our spectra. Also Zhang et al. (2010) did not find any H₂O-related bands in their FTIR spectra of partially dehydrated sericite, muscovite, pyrophyllite, and talc. Zhang et al. (2005) prefer a hydrogen/oxygen/hydroxyl hopping mechanism over H₂O molecule diffusion as the transport mechanism for the dehydroxylation products. The transport by hopping requires similar bond breaking and formation steps as the initial liberation reactions. The transport mechanism would have similar $E_a$ as the primary reactions, and could, therefore, also become rate-limiting. Thus, transport could also contribute to the $E_a$ plateau.

The increased intensity of the unassigned OH mode during dehydroxylation in both the FTIR and Raman spectra may be related to such transient hydroxyl groups formed during transport. This interpretation is tentative, because the unassigned hydroxyl band is the result of the deconvolution of strongly overlapping bands.

**Dehydroxylation products: A talc-like intermediate and forsterite**

To approximately 450 °C, the spectra are only affected by a constant frequency shift of the Raman and IR bands, which relate to thermal expansion of the structure. The intensity of all bands remains nearly constant to this temperature (Figs. 4 and 11a). Above this temperature, bands assigned to atom groups belonging to the octahedral sheet, i.e., $A_{1g}$ mode of lizardite (Fig. 6b) and OH stretching vibrations (Fig. 9a), show a strongly discontinuous behavior (Figs. 6b and 6f). This is interpreted as resulting from
changes in the atomic bonding environment. These changes occur at about the same temperature where the absorbances/intensities of the OH bands start to decrease. Thus, the onset of dehydroxylation affects only the octahedral sheet. The different behavior of the in-phase (Fig. 9a) and out-of-phase OH stretching mode frequencies (Fig. 9c) as function of temperature, however, is surprising. The in-phase stretching mode shows a break in the slope of $\delta \nu/\delta T$, whereas the slope of the out-of-phase mode is characterized by a linear slope. Modes originating from the same atomic group should show the same frequency behavior. A possible cause may be the deconvolution procedure.

The small temperature-dependent shift of the O-Si-O bending mode located at 129 cm⁻¹ (RT) to lower wavenumbers is related to the change in ditrigonal distortion of the tetrahedral sheet as demonstrated by X-ray diffraction (Guggenheim and Zhan 1998). The shifting rate $\delta \nu/\delta T$ is one order of magnitude smaller than for all other bands (~0.439 cm⁻¹/100 °C). The first intensity changes in Si-O-related Raman modes occur above 600 °C (Fig. 4) where the hydroxyl modes have lost more than half of their initial intensity (Fig. 12a). The tetrahedral sheet is thus not affected by the initial changes in the octahedral sheet. The large jump in frequency of the $A_{1g}$ mode of lizardite down to 183 cm⁻¹ and the jump in frequency of the $v_3$ Si-O-Si band is assigned to product phase(s) of the dehydroxylation. Under ambient, dry conditions the first recognizable dehydroxylation product is mainly amorphous, but crystallizes upon further heating to forsterite, followed by enstatite (Brindley and Hayami 1963). The stable products of the dehydroxylation reaction of pure magnesian lizardite under hydrothermal conditions are talc, forsterite, and water (O’Hanley et al. 1989). Various authors (e.g., McKelvy et al. 2004; Brindley and Zussman 1957) reported low-angle peaks in X-ray diffractograms of thermally treated serpentine phases, indicating that amorphization is not complete and that dehydroxylation produces intermediate partially ordered structures. Although differing in detail, early studies by Ball and Taylor (1963) and Brindley and Hayami (1965) proposed reaction sequences with disordered dehydroxylated intermediates. They infer the presence of a partially ordered structure from the topotactic relationship between forsterite and serpentine (Brindley and Hayami 1963). Based on a peak corresponding to a $d$-spacing of 14 Å, McKelvy et al. (2006) proposed the formation of a lamellar oxyhydroxide meta-serpentine characterized by a doubled cell dimension along the $c$ axis and a heterogeneous OH distribution. Their DFT calculations support the formation of such a 14 Å structure, which is transformed to an amorphous phase with progressive dehydroxylation. Two different partially ordered, anhydrous intermediates, dehydroxylate I and II, are proposed by MacKenzie and Meinhold (1994) based on $^{25}$Mg and $^{29}$Si shifts in NMR spectra of heated chrysotile samples. According to these authors, dehydroxylate I has a structure similar to the parent phase, whereas the $^{29}$Si NMR position of dehydroxylate II is similar to talc. Viti (2010) suggests the formation of intermedi-
ate, partially ordered dehydroxylation products on the bases of DTG/DTA analyses of lizardite. Their weight-loss rate curves show two maxima, a feature also observable in DTG experiments of Trittschack and Grobéty (2012). The first maximum is attributed to the partial dehydroxylation of the initial lizardite leading to the intermediate phase. The dehydroxylation of the latter would be responsible for the second maximum. Three new features at 183, 350, and 670 cm⁻¹ in the spectrum at 665 °C, which cannot be assigned to either lizardite or forsterite, are interpreted as originating from one or more intermediate phases. Supported by a band appearing at 1000 cm⁻¹ (690 °C), it is possible to assign them as Raman bands related to talc as reported by Fumagalli et al. (2001). The bands located at 670 and 1000 cm⁻¹ are also observable in the spectrum of the quenched dehydroxylated sample, where they are shifted to 686 and 1021 cm⁻¹, respectively. Referring to Fumagalli et al. (2001) these bands can be assigned to a Mg(OH)₂ mode (183 cm⁻¹), a symmetric SiO₄ mode (350 cm⁻¹), and one O-Si-O-Si mode in case of the 670 cm⁻¹ (686 cm⁻¹ at RT) feature (Fig. 5d) and one Si-O stretching mode represented by the 1000 cm⁻¹ feature at 819 °C and 1021 cm⁻¹, respectively, in the spectra of the quenched sample. The strongest of these three weak single bands in the OH stretching mode region (Fig. 7d) is also consistent with the spectrum of talc (Fumagalli et al. 2001). This interpretation supports NMR observations by MacKenzie and Meinhold (1994) who proposed a talc-like structure for their dehydroxylate II phase. During dehydroxylation, the partial pressure of the volatile products may be sufficiently high in certain regions of the sample to nucleate talc, a stable product of the lizardite dehydroxylation in the hydrous system (O’Hanley et al. 1989). The band at 183 cm⁻¹ appears also in the dehydroxylation of chrysotile (Trittschack and Grobéty, unpublished manuscript), where it is present for a much broader temperature range.

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FIGURE 12. (a) Decrease of the integrated intensity of all Raman OH stretching bands as function of temperature and corresponding derivative curve and (b) for the individual inner-surface OH stretching bands and the unassigned band. All symbols have the identical assignment as in Figure 10.

REFERENCES CITED


