

Hydrogen positions and thermal expansion in lizardite-1T from Elba: A low-temperature study using Rietveld refinement of neutron diffraction data

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ABSTRACT

The structure of lizardite-1T from Monte Fico, Elba, was refined in space group $P31m$ using neutron diffraction data, measured at 8, 150, and 294 K, and full-profile Rietveld refinement techniques. The lattice parameters at 8 K [$a = 5.3267(2)$, $c = 7.2539(6)$ Å], 150 K [$a = 5.3260(2)$, $c = 7.2574(6)$ Å], and 294 K [$a = 5.3332(2)$, $c = 7.2718(6)$ Å] show nonlinear expansion, with nearly all volume change above 150 K. H positions were precisely refined at 8 K. The inner H4 atom deviates from the idealized 0,0,z positions and is disordered over three symmetry-related positions 0.24 Å away from the ternary axis. The outer H3 atom location is consistent with the previous single-crystal X-ray structure refinement. On the basis of the present thermal expansion data and previous compressibility measurements, the following equation of state for lizardite-1T is proposed: $V_{P,T} = V_0[1 + 32.8 \times 10^{-6}(T - 294) - 15.5 \times 10^{-4}(P - 0.001)]$. Accordingly, the constant volume condition requires geothermal gradients on the order of 15 °C/km.

INTRODUCTION

Because of its simplicity, the structure of lizardite, ideally $\text{Mg}_3\text{Si}_2\text{O}_5(\text{OH})_4$, is the archetype model for Mg-rich serpentines (Mellini and Viti 1994). The structure was described with two modules: the 1:1 layer and the interlayer region. The latter involves a sheet of closest-packed octahedral OH groups, which faces the tetrahedral bases of the adjacent 1:1 layer at a distance that allows hydrogen bonds to be formed. Previous refinements, using X-ray data obtained from seven single crystals, showed that the internal dimensions of the 1:1 layer do not change with chemical substitution or increasing pressure. Instead, the interlayer thickness varied significantly, suggesting that the thickness depends on the hydrogen bonds linking adjacent layers (Mellini 1982; Mellini and Zanazzi 1987, 1989).

The accurate determination of H positions is difficult with X-ray data. Although a neutron single-crystal data set is optimum, lizardite crystals of sufficient size are unknown. Thus, we conducted a powder neutron diffraction study by full-profile Rietveld refinement, using a polycrystalline sample of almost pure lizardite-1T from Monte Fico, Elba. This material (Mellini and Viti 1994) occurs in large masses that exhibit excellent crystallinity, as seen from X-ray and electron diffraction. To our knowledge, no other occurrence gives material of similar quality.

EXPERIMENTAL METHODS

Sample preparation

The Monte Fico lizardite is compositionally homogeneous (Viti and Mellini, in preparation). Average values

(Mellini and Viti 1994) are as follows: $\text{MgO} = 37.97$, $\text{SiO}_2 = 39.65$, $\text{Al}_2\text{O}_3 = 2.86$, $\text{FeO} = 3.97$ wt%.

Lizardite crystals, 0.1–0.5 mm in diameter, and grains made up of several crystals were selected under the optical microscope. The whole was then gently crushed and analyzed by X-ray powder diffraction to determine the mineralogic purity and crystallinity. Care was taken to avoid chrysotile impurities, which occasionally fill interstices between lizardite crystals, and deformation of lizardite crystals during grinding. The X-ray diffractograms showed only one small extra peak resulting from chrysotile. Lizardite peaks did not indicate stacking faults. For example, the 100 reflection did not show a tail toward higher 2θ values.

Data collection

Neutron diffraction data were collected using the multi-detector powder diffractometer (TAS3) at the steady-state DR3 reactor at Risø National Laboratory in Denmark. The instrument is a two-axis neutron diffractometer situated at a thermal beam. It has a Debye-Scherrer geometry with a vertical axis and a bank of 20 He-3 detectors spanning about 106° in 2θ (Als-Nielsen et al. 1988). Each detector covers approximately 5.3°, so step scans of $100 \times 0.053^\circ$ give diffraction patterns of 2000 data points. A vertically focusing Ge(711) monochromator set at $2\theta = 85.1^\circ$ provided a beam of incident neutrons with wavelength 1.06966(2) Å, as determined from refinement of a diffraction pattern from a standard sample ($\alpha\text{-Al}_2\text{O}_3$, $R\bar{3}c$, 5.12984 Å, 55.28333°). The divergence of the incident (8 mm wide $\times$ 30 mm high) and diffracted beams was defined to 10 min by one Soler collimator placed before

the sample and twenty 10 min Soller collimators placed after the sample in front of each He-3 detector. The resulting instrumental resolution was about 18 min FWHM at $2\theta = 45^\circ$. About 4.7 g of powdered lizardite sample were introduced in a sealed cylindrical vanadium container (9 mm inner diameter $\times$ 40 mm height and 0.1 mm wall thickness). Before sealing, the vanadium container was evacuated and filled with He exchange gas (~ 300 mbar) to ensure thermalization of the powder grains.

For diffraction experiments at room temperature (294 K), the vanadium container was placed directly on the diffractometer sample table, where a special device provided for sample rotation around the ω axis of the diffractometer during data collection. The low-temperature diffraction patterns at 8(1) and 150(1) K were obtained using a Displex cooling device with a cylindrical aluminum vacuum jacket with an overall wall thickness of 2 mm in the diffraction plane. This device was fixed onto the diffractometer sample table, and no sample rotation was provided. The vanadium container was screwed directly onto the low-temperature copper block of the cooling device. This copper block can be heated above base temperature (approximately 8 K) and is also in direct contact with a calibrated germanium thermometer and a 100 Ω platinum resistance thermometer. The germanium thermometer was used for measuring temperatures up to 80 K, and the platinum resistor for measuring temperatures from 50 to 300 K. Temperatures were monitored at 10 ms intervals and recorded simultaneously with the step-scan data.

The data collection was, in all cases, divided into several experiments, each defined by its monitor count measured in the primary beam between entrance collimator and sample. Monitor counts on the order of 10^4 – 10^5 were used, corresponding to about 1–10 h. Thus, difference patterns between successive experiments could be calculated to check for structural changes during the measurement, as might be expected for the low-temperature experiments if the sample were not in complete thermal equilibrium. For the latter experiments, the raw observed intensities were corrected by subtracting the contribution of the empty cryostat. Because these corrections were small in comparison with the resulting intensities, no additional treatment of errors was made, and estimated standard deviations obtained from counting statistics were used throughout. The background level and fluctuations of these patterns were, within 1.9–2.5 times the estimated standard deviation from counting statistics, the same as those obtained at room temperature. No visible effect of graininess, which might arise from the lack of sample rotation in the cryostat, was observed.

The last detector had a greater noise fluctuation in all experiments, and an intensity drift was occasionally observed for the first detector. After the early stages of refinement, these data were therefore excluded from least-squares calculations. Other regions were eventually excluded to eliminate impurity peaks as noted below.

In Figure 1, the observed neutron powder diffraction patterns at 8, 150, and 294 K are shown. Note that the peak definition is greatly improved in low-temperature experiments, particularly in the high- 2θ region.

Structure refinement

The GSAS program system (Larson and Von Dreele 1994) was used for Rietveld refinement. The space group $P31m$ and atomic parameters were taken from single-crystal X-ray diffraction results (Mellini and Viti 1994) for the same specimen used in the present study. The octahedral site occupancy (hereafter referred to as "Mg site" to be consistent with previous papers) was set at 0.947 Mg + 0.053 Fe to correspond to the chemical analysis (average of ~ 50 crystals). A check for the constancy of these values during the final stages of refinement showed only negligible changes, confirming the microprobe results. No account was made for the low Al content (see above), considering that the neutron-scattering length of Al (3.45 fm) is close to that of the elements for which it might substitute, i.e., Si (4.15 fm) and Mg (5.38 fm).

During the first stage of Rietveld refinement, all parameters were fixed, except for the scale factor, background coefficients (a three-term cosine Fourier series), and unit-cell parameters. The instrument zero point was fixed at 0.034° (2θ), as determined by the standard α - Al_2O_3 refinement. Subsequently, the peak shape was refined using a pseudo-Voigt profile function, which contained the Gaussian U , V , W parameters (Caglioti et al. 1958), a Lorentzian $\gamma = X/(\cos \theta)$ parameter to account for particle size, and a cut-off of the wings at 0.1% of the peak maximum. The U , V , and W Gaussian parameters were fixed at values obtained from the standard α - Al_2O_3 refinement, and only the particle-size parameter was refined. The Lorentzian X values obtained from the three patterns ranged from 11.4 (8 K) to 12.2 (294 K). A March-Dollase preferred-orientation correction (Dollase 1986) was introduced, with [001] as the preferred direction, and the corresponding parameter converged to the same value of 1.17(1) for all refinements.

Special care was taken in background modeling to avoid effects from small chrysotile impurities. A tiny quantity of chrysotile impurity was isolated and mounted in a Debye-Scherrer camera equipped with CHI-70, a Gandolfi device (Gregorkiewicz 1994; Serrano-Santos and Gregorkiewicz 1995). The resulting film was converted to a digitized diffraction pattern with intensity vs. scattering angle (Nguyen and Jeanloz 1993). Its comparison with the observed and calculated X-ray patterns of lizardite indicated that a small amount of chrysotile might account for many discrepancies, such as a few weak extra peaks, shoulders, and an anomalous broadening at the base of some lizardite peaks. Because chrysotile exhibits much lower crystallinity, the discrepancies appear as broad, nonrandom fluctuations of the background and could easily be recognized also in the neutron patterns of our lizardite sample. Seven chrysotile contributions were then removed from the observed neutron patterns by subtrac-

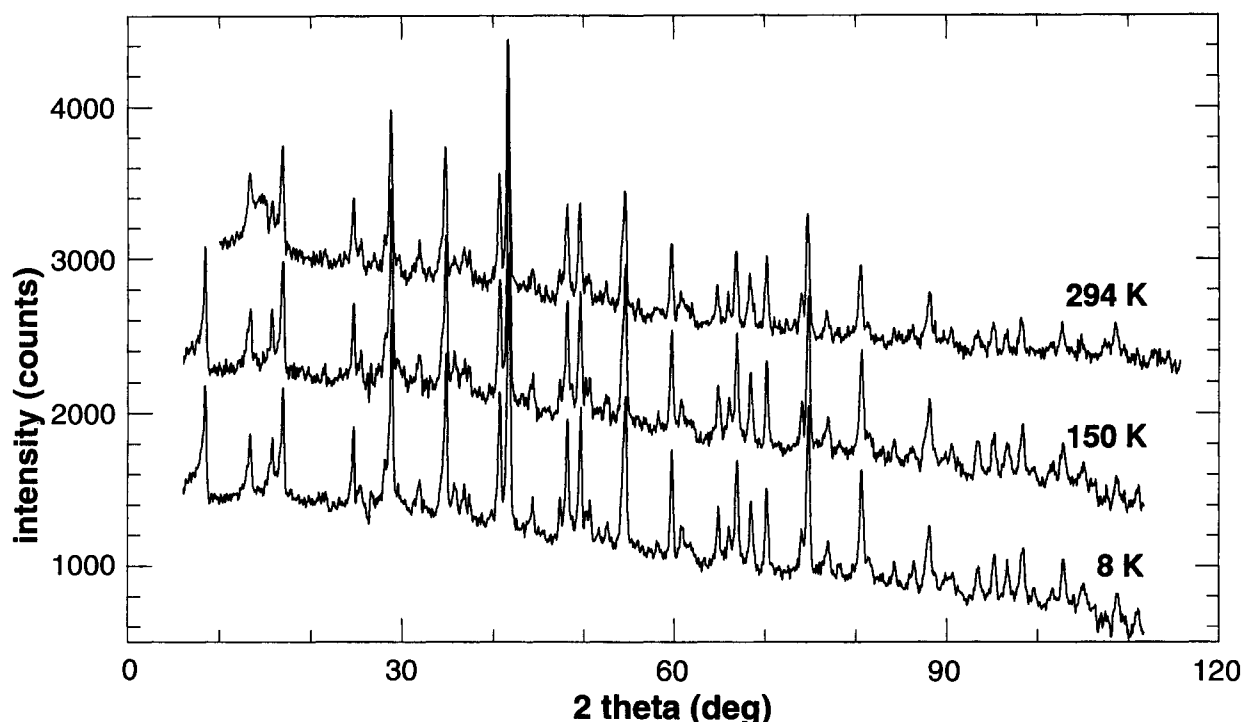


FIGURE 1. Observed neutron powder diffraction patterns for lizardite-1T at 8, 150, and 294 K. The 150 and 294 K patterns are displaced for comparison.

tion as fixed background points. After such corrections, the refinement converged to χ^2 values of 1.57, 1.37, and 1.06 and R_{Bragg} values of 0.19, 0.17, and 0.27, respectively, for the 8, 150, and 294 K refinements.

At this stage, atomic parameters were allowed to refine. Isotropic displacement factors were refined for four groups of atoms, namely Si-Mg-Fe, O1-O3-O4, H3-H4, and O2. To improve convergence, 35 soft constraints, corresponding to the Si-O, Mg-O, O-O, Si-Si, and O-H distances, were imposed. The ideal values for these parameters were taken from lizardite single-crystal X-ray data (Mellini and Viti 1994), except for the O-H distance, which was set to 0.97(5) Å (Chiari and Ferraris 1982). The weight of the soft constraints was progressively reduced during successive refinement cycles. Atomic coordinates were refined, beginning with Si, Mg, and O1, O3, O4 and followed by the H atoms and O2.

Least-squares refinements using 8, 150, and 294 K data converged to χ^2 values of 1.35, 1.10, and 0.61 and to R_{Bragg} values of 0.12, 0.14, and 0.14, respectively (Table 1). In Figure 2, the observed and calculated patterns, corresponding to the Rietveld refinement using 8 K data, are given along with the difference curve.

RESULTS

Unit-cell parameters

Unit-cell parameters are shown in Table 1. The room-temperature cell [$a = 5.3332(2)$, $c = 7.2718(6)$ Å] matches values obtained by X-ray diffraction of two selected crys-

tals [$a = 5.330(4)$ and $5.338(4)$, $c = 7.257(6)$ and $7.269(6)$ Å; Mellini and Viti 1994]; the current values represent the average of about 10^7 single crystals. The unit-cell parameters decrease upon cooling, with the c parameter being more sensitive to temperature than the a parameter (Fig. 3). Maximum variation occurs between 150 and 294 K, and a nonlinear dependence of the a and c unit-cell parameters of lizardite with respect to temperature is evident. These results emphasize the need for caution in assumptions regarding extrapolations of unit-cell parameters, especially for thermodynamic calculations. The value calculated between 150 and 294 K is accepted as a lower estimate of the expansion coefficient of lizardite. Nonlinear behavior was noted by Mellini and Zanazzi (1989) in their determination of lizardite compressibility.

TABLE 1. Rietveld refinement parameters and unit-cell data of lizardite-1T at 8, 150, and 294 K

T (K)	8	150	294
R_p	0.027	0.028	0.024
R_{wp}	0.035	0.036	0.031
$R(F^2)$	0.124	0.137	0.143
No. obs	274	271	250
No. p	24	23	23
χ^2	1.349	1.103	0.612
a (Å)	5.3267(2)	5.3260(2)	5.3332(2)
c (Å)	7.2539(6)	7.2574(6)	7.2718(6)
V (Å ³)	178.25(1)	178.28(1)	179.12(1)

Note: R_p = profile residual, R_{wp} = weighted profile residual, $R(F^2) = R_{\text{Bragg}}$, No. obs = number of reflections $F^2(hkl)$ extracted, No. p = number of variables.

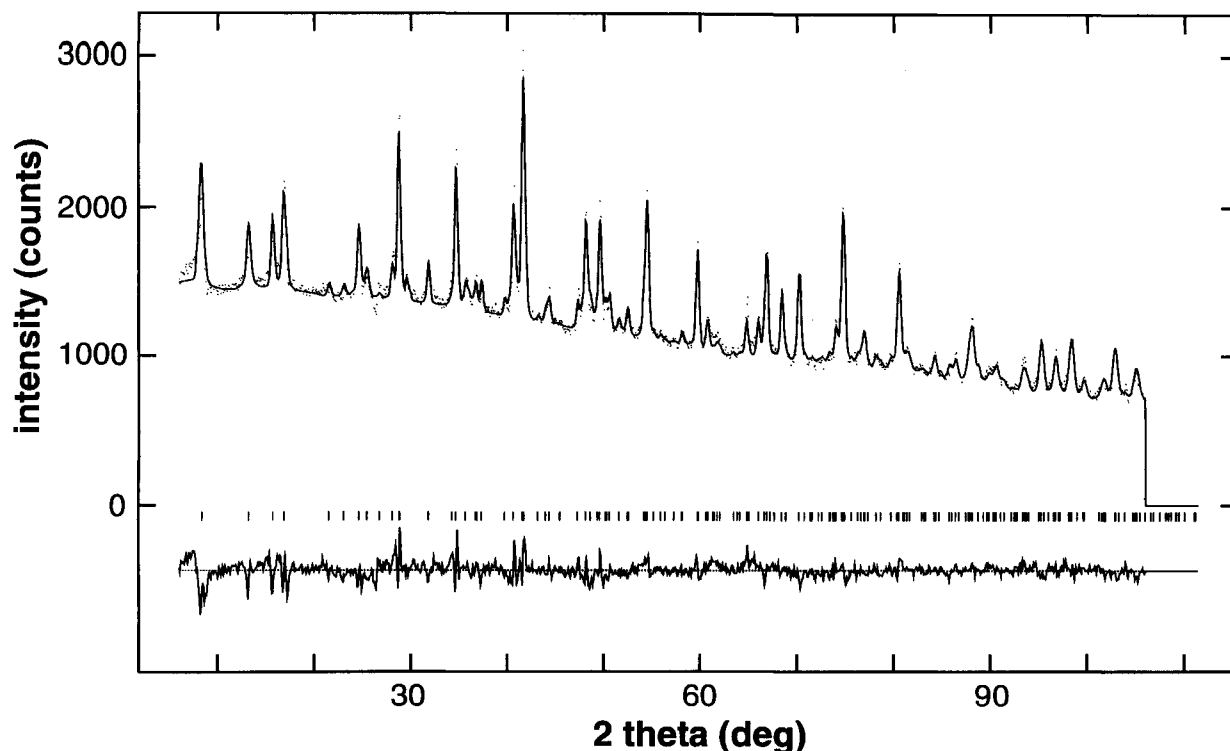


FIGURE 2. Observed (points), calculated (solid line), and difference pattern corresponding to the Rietveld refinement using the 8 K data. Tick marks indicate positions of allowed Bragg reflections.

Structural results

The Rietveld refinement based on powder neutron diffraction closely matches the results from single-crystal X-ray diffraction (Table 2, cf. Table 2 in Mellini and Viti

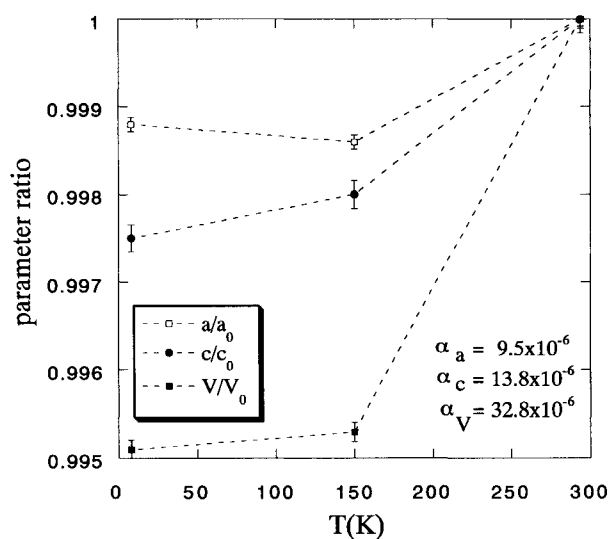


FIGURE 3. Normalized temperature dependence of the unit-cell parameters of lizardite-1T. The α values are the expansion coefficients measured between 150 and 294 K. Error bars correspond to ± 1 estimated standard deviation.

1994). Only the atomic coordinates obtained from the 8 K refinement are reported here because they represent the most significant structural result. As in the previous study, negative ditrigonalization characterizes the 1T polytype, as predicted by Mellini (1982). The deviation from hexagonal configuration is $\alpha = -2.6^\circ$, which compares to the -2.7 and -2.6° obtained by X-ray analysis at room temperature.

H positions

Two H sites (Fig. 4) occur in lizardite. H3 is the outer H atom, which is bonded to O3 and links the octahedral

TABLE 2. Atomic positions and isotropic mean-square displacements (U_{iso}) of lizardite-1T obtained from Rietveld refinement of the 8 K pattern

	x	y	z	U_{iso} (8 K)	U_{iso} (150 K)	U_{iso} (294 K)
Si	$\frac{1}{3}$	$\frac{2}{3}$	0.070(1)	20(8)	33(9)	42(8)
Mg	0.324(2)	0	0.447(2)	20(8)	33(9)	42(8)
O1	$\frac{1}{3}$	$\frac{2}{3}$	0.292*	2(5)	5(5)	71(7)
O2	0.507(2)	0	-0.014(2)	88(12)	112(13)	157(13)
O3	0.665(2)	0	0.587(1)	2(5)	5(5)	71(7)
O4	0	0	0.291(3)	2(5)	5(5)	71(7)
H3	0.646(3)	0	0.715(2)	78(23)	157(19)	321(22)
H4	0.046(6)	0.046(6)	0.172(4)	78(23)	157(19)	321(22)

Note: The displacement parameters corresponding to the 150 and 294 K refinements are reported for comparisons. U_{iso} reported as squared picometers.

* Not refined in order to fix unit-cell origin.

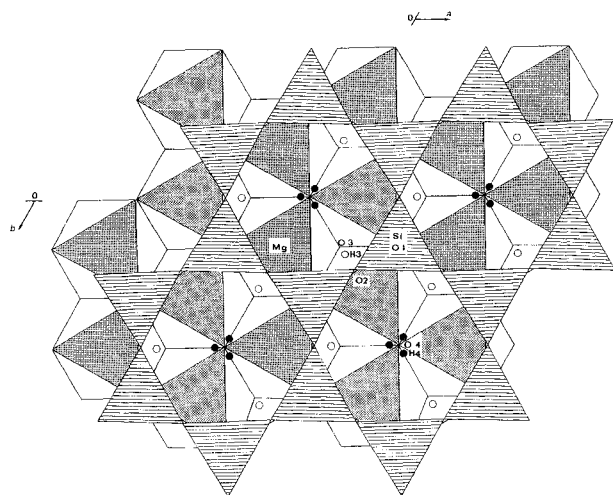


FIGURE 4. Projection of the crystal structure of lizardite-17 from the 8 K refinement. Open circles = H3 = outer H atom, solid circles = H4 = inner H atom. Note that the structure is viewed from the bottom with c pointing away from the observer.

sheet of a given 1:1 layer to the tetrahedral sheet of the adjacent 1:1 layer. H4 is the inner H atom, bonded to O4, and ideally located on the ternary rotation axis perpendicular to the ditrigonal ring with coordinates 0,0, z .

H3 was located, although with greater accuracy, near the position obtained by the previous X-ray determination (0.646,0,0.715 vs. 0.660,0,0.710). The important point is that this study confirms previous conclusions. In particular, from Figure 4, we note a correlated atomic displacement. In projection, H3 is slightly displaced from its idealized position on the perpendicular over O3 and toward O2. In concert with this, the O2 atom in the adjacent 1:1 layer is displaced toward H3, giving a slightly ditrigonal distortion of the six-membered ring of silicate tetrahedra. Thus, the interlayer hydrogen bond strengthens to give an O2-H3 distance of 2.10 Å.

For H4, a short O-H distance (O4-H4 $\approx$ 0.8 Å) suggests that the 0,0, z coordinate is an average position, with H4 disordered over three or six symmetry-related positions off the ternary axis. Among various alternatives tested in the last stages of the 8 K refinement, two cases are considered here. In the first case, H4 is displaced along [100], with subsequent decrease of the H4-O4-Mg angle, whereas in the second case H4 is displaced along [110], with subsequent increase of the H4-O4-Mg angle. The latter refinement, corresponding to H4 at 0.046, 0.046, 0.172 (i.e., at 0.24 Å off the ternary axis), gave the best agreement with observed data according to the test by Hamilton (1965). This result is explained as a combined effect of tetrahedral ring ditrigonalization and octahedral substitution. In particular, H4, in the observed off-axis position, points toward one of the three O2 atoms that, because of the ditrigonalization of the ideally hexagonal six-membered tetrahedral ring, moved toward the center of the ring (Fig. 4). Thus, the displacement short-

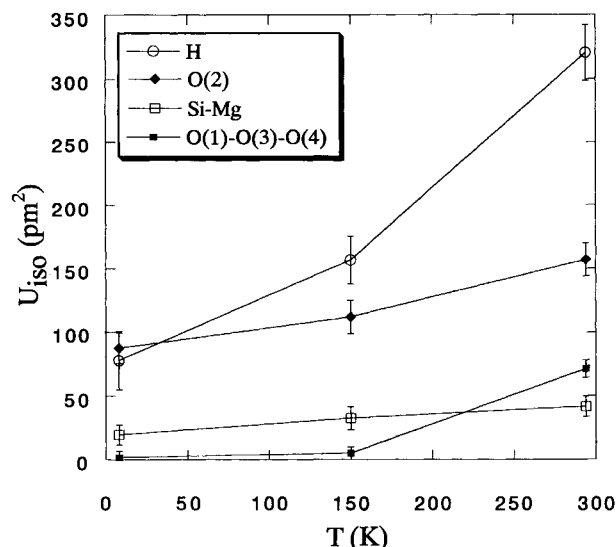


FIGURE 5. Temperature dependence of the isotropic Debye-Waller factor in lizardite-17. Error bars correspond to ± 1 estimated standard deviation.

ens some O2-H4 distances. Moreover, the observed increase of the H4-O4-Mg angles is expected for the substitution of Mg with ions of higher charge, as seen in dioctahedral micas or trioctahedral micas with different elements in the octahedral positions (see, e.g., Serratos and Bradley 1958; Giese 1979).

Debye-Waller parameters

Mellini and Viti (1994) observed anomalously large Debye-Waller factors for O2, in particular for the U_{11} component. They tentatively attributed this anomaly to static or dynamic positional disorder arising from negative-to-positive ditrigonalization of the six-membered rings in the tetrahedral sheets. In Rietveld powder diffraction refinements, it is difficult to obtain meaningful anisotropic Debye-Waller parameters, and it is known that, even in the isotropic case (U_{iso}), values may greatly differ from those obtained by single-crystal diffraction of the same material. The present result can, however, be used for comparison of the data obtained at different temperatures. In fact, Figure 5 shows a general decrease of Debye-Waller factors with temperature, which is almost already complete at 150 K. However, whereas mean-square displacement amplitudes of all other O atoms approach zero, O2 (as well as the H atoms) maintains values well above zero. This means that, for those atoms, the Debye-Waller factors cannot entirely be explained by thermal motion, in support of the above-mentioned idea of positional disorder, which might result from occasional positive ditrigonalization of the tetrahedral sheet.

DISCUSSION AND CONCLUSION

We derive an approximate equation of state of lizardite, using the present thermal-expansion data (but great-

er expansion coefficients are expected if measured at higher temperatures) for the Elba lizardite, together with compressibility coefficients calculated from the data reported for the near-isochemical Coli lizardite (Mellini and Zanazzi 1989). The approximate relations are as follows:

$$a_{P,T} = a_0[1 + 9.5 \times 10^{-6}(T - 294) - 2.2 \times 10^{-4}(P - 0.001)]$$

$$c_{P,T} = c_0[1 + 13.8 \times 10^{-6}(T - 294) - 11.2 \times 10^{-4}(P - 0.001)]$$

$$V_{P,T} = V_0[1 + 32.8 \times 10^{-6}(T - 294) - 15.5 \times 10^{-4}(P - 0.001)].$$

Clearly P (here given in kilobars) and T are not analogous in lizardite. Although linear expansion along [100] is roughly similar to that along [001], the compressibility along [001] is considerably larger than that along [100]. Balancing effects of P and T may be estimated by considering volume expansion and compressibility coefficients. From the above equations, we find that a 50 °C increase is offset by a 1 kbar increase. Assuming 3 is a reliable specific gravity for serpentinites, the 1 kbar increase relates to a depth of 3.3 km. Therefore, no important volume change would occur for serpentinites emplaced under a geothermal gradient of 15 °C/km. Of course, if the gradient is higher, a volume increase would occur; conversely, if the gradient is lower, contraction would occur.

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