



Article High-Pressure Raman Spectroscopy and X-ray Diffraction Study on Scottyite, BaCu₂Si₂O₇

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Abstract: In situ high-pressure synchrotron X-ray diffraction and Raman spectroscopic experiments of scottyite, BaCu₂Si₂O₇, were carried out in a diamond anvil cell up to 21 GPa at room temperature. X-ray diffraction patterns reveal four new peaks near 3.5, 3.1, 2.6 and 2.2 Å above 8 GPa, while some peaks of the original phase disappear above 10 GPa. In the Raman experiment, we observed two discontinuities in $d\nu/dP$, the slopes of Raman wavenumber (ν) of some vibration modes versus pressure (P), at approximately 8 and 12 GPa, indicating that the Si-O symmetrical and asymmetrical vibration modes change with pressure. Fitting the compression data to Birch–Murnaghan equation yields a bulk modulus of 102 ± 5 GPa for scottyite, assuming K_0' is four. Scottyite shows anisotropic compressibility along three crystallographic axes, among which *c*-axis was the most compressible axis, *b*-axis was the last and *a*-axis was similar to the *c*-axis on the compression. Both X-ray and Raman spectroscopic data provide evidences that scottyite undergoes a reversible phase transformation at 8 GPa.

Keywords: scottyite; high pressure; phase transformation

1. Introduction

Rock-forming minerals that occur on the Earth's crust may change structures and physical properties in the mantle environment. Numerous high-pressure experiments in Earth sciences were focused on silicate minerals that are directly relevant to the Earth's interior. Some orthosilicate minerals, such as olivine, show that the SiO₄ tetrahedron is relatively rigid; not until pressures as high as 30 GPa do SiO₄ tetrahedron start to show signs of distortion. The distortion of SiO₄ tetrahedron is interpreted as due to the distortion of Si-O-Si linkage, which was recognized by the kink in the $d\nu/dP$ (the slopes of Raman wavenumber (ν) versus pressure (P)) plot [1]. In contrast, before the distortion of SiO₄ tetrahedron, the most easily distorted crystal structure took place on the bond connection between cation and SiO₄ tetrahedron. Therefore, it would be interesting to look into sorosilicates that have two tetrahedrons shared and linked by one oxygen atom, forming a basic structural unit of Si₂O₇. The SiO₄ tetrahedron is considered to be a rigid body and not easy to be compressed.

Scottyite, named after Michael M. Scott, is a Ba-Cu sorosilicate, BaCu₂Si₂O₇, [2]. As a newly found mineral, scottyite is a very rare mineral, first reported as an unnamed Ba-Cu silicate from Eifel, Germany [3–5]. It was then identified as an orthorhombic system with space group *Pbnm* and unit-cell parameters a = 6.866 Å, b = 13.901 Å, c = 6.902 Å, and V = 625.70 Å³ [6]. It was later found in many other localities, including the Wessels mine, Kalahari Manganese Fields, Northern Cape Province, Republic of South Africa, from which the unit-cell parameters are determined as a = 6.8556 Å, b = 13.1725 Å, c = 6.8901 Å, and



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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). V = 622.21 Å³ on the basis of a single-crystal XRD study [2]. The structure of scottyite is based on a tetrahedral framework consisting of SiO₄ and CuO₄ tetrahedrons. The flattened CuO₄ tetrahedrons share corners with one another to form chains parallel to the c-axis. These chains are interlinked by Si₂O₇ tetrahedral dimers and Ba²⁺ cations, which are bonded to seven O atoms of irregular coordination numbers. Within the CuO₄ tetrahedral chain, the Cu-O-Cu angle is 124.49°, which is responsible for the antiferromagnetic coupling in BaCu₂Si₂O₇ [2,7–9].

2. Experimental Method

2.1. Raman Spectroscopy

Small chips of natural scottyite crystal (RRUFF Project, deposition no. R120077, from the Wessels mine, Republic of South Africa) with a dimension of 80 μm were used as a starting material in the Raman spectroscopic experiment. In this high pressure work, a disk of T301 stainless steel with a thickness of 250 µm was used as a gasket. The gasket was first compressed between the diamond anvils to make an indentation on which a hole of 150 µm was then drilled for the sample chamber. The sample was loaded in the chamber using a 4:1 methanol:ethanol mixture as pressure-transmitting medium. It was then gradually compressed to high pressure and then decompressed to room pressure. The ruby fluorescence method was used for pressure measurement [10] after each pressure increment. We measured more than two ruby grains to make sure that the pressure of the compressed sample remained hydrostatic. A Horiba iHR550 unit installed at the Department of Earth Sciences, National Cheng Kung University, was used in the Raman spectroscopic study. A semiconductor laser beam of 458 nm was used as an excitation source. The laser light of 100 mW was focused on the sample in the diamond cell, and Raman signals were collected in a back-scattered way by the CCD detector. The recording time of each Raman spectrum was 60 s (averaged by 3 times of acquisition). During each measurement, pressure was recorded before and after each Raman spectrum was collected. Both compression (loading) and decompression (unloading) processes were conducted.

2.2. X-ray Diffraction

Synchrotron X-ray diffraction method was applied to the study of the behaviors of natural scottyite (RRUFF Project, deposition no. R120077, from the Wessels mine, Republic of South Africa) at high pressure. The powder of scottyite was then loaded into the hole (200 μ m in diameter) in a T301 stainless steel gasket, prepared in the same way as the high pressure Raman measurement. After the chamber was loaded with the sample, it was then filled with silicon oil as a pressure-transmitting medium, and then gradually compressed to high pressure and decompressed to room pressure. Pressure determination was achieved by observing the shift of the ruby R1 emission line. More than two ruby grains were measured, to make sure that the pressure remained hydrostatic.

The X-ray diffraction (XRD) experiment was carried out at PLS-II 5A beamline at the Pohang Accelerator Laboratory, South Korea. The size of the X-ray beam was confined to 100 μ m and the position of the beam was adjusted to fall on the center of the sample chamber in each run. Pressure was gradually increased up to about 17 GPa and then decreased relatively rapidly back to the ambient conditions. The scattering data were collected by MAR345 detector and integrated using the FIT2D v10.132 (ESRF) [11]. Lattice parameters were calculated by a least-square method developed by Novak and Colville [12].

3. Results

3.1. Raman Spectroscopy

The Raman spectra of scottyite are arbitrarily divided into four regions. Region 1 (ν_1 and ν_3 modes), between 800 and 1100 cm⁻¹, contains bands attributed to the Si-O symmetric and antisymmetric stretching vibrations within the SiO₄ tetrahedrons. Region 2, between 660 and 700 cm⁻¹, includes bands resulting from the Si-O_{br}-Si (O_{br}: bridging O atom) bending vibrations within the Si₂O₇ tetrahedral dimers. Region 3 (ν_2 and ν_4 modes)

ranges from 420 to 660 cm⁻¹, and contains modes related to the O-Si-O symmetric and anti-symmetric bending vibrations within the SiO₄ tetrahedrons. The bands in Region 4, below 420 cm⁻¹, are mainly related to Cu–O interactions and lattice vibrational modes, as well as the rotational and translational modes of SiO₄ tetrahedrons [2].

Table 1 shows the comparison of our results of Raman data of scottyite with the previous investigation at the room pressure. Some of the representative Raman spectra of scottyite were taken during the loading and unloading processes of the experiment in this study (Figure 1). The v_1 mode (895 cm⁻¹) and v_3 mode (1016, 958 and 863 cm⁻¹) in Region 1, Si-O_{br}-Si bending mode (675 cm⁻¹) in Region 2, and ν_2 mode (458 cm⁻¹) and ν_4 mode $(608, 579 \text{ and } 560 \text{ cm}^{-1})$ in Region 3 are observed at the ambient conditions in this study. From Figures 1 and 2a,b, the frequencies of most Raman modes are found to increase with increasing pressure, except peak R_2 , ν_4 , ν_2 , R_{4c} , R_{4f} , R_{4g} and R_{4k} between 8 and 12 GPa. The intensity of most of the peaks becomes weaker with increasing pressure while some Raman bands (such as R_2 , v_4 and v_2 mode) remain quite sharp at all pressures. It is found that the v_1 mode of scottyite becomes broader, and one distinct peak (new peak) is found on the right shoulder when the pressure exceeds 8 GPa, and an additional new peak was found above 12 GPa. The intensity of the new peak becomes stronger and shifts to a lower wavenumber with increasing pressure. The R_2 mode of scottyite remains quite sharp at all pressures, but shifts to lower wavenumber with increasing pressure between 8 and 12 GPa. The ν_{3a} mode of scottyite becomes weaker with increasing pressure between 8 and 12 GPa, and disappears above 12 GPa. We also found some new peaks on v_1 , R_{4d} modes between 8 and 12 GPa and ν_1 , ν_4 , R_{4a} , R_{4c} , R_{4i} modes above 12 GPa.

The discontinuities of the linear trend on the dv/dP (Raman modes shift as function of pressure) are also shown in Tables 2 and 3. All the modes of scottyite show discontinuity in the dv/dP slope at ~8 GPa, and some modes show additional discontinuity at 12 GPa. During the unloading process, all vibrational modes persist until 8 GPa where all modes shift back to the position as the loading process. Below 8 GPa, the decompression data are similar to those of the compression ones. There are seven peaks that show negative slop between 8 and 12 GPa (R_2 , v_4 , v_2 , R_{4c} , R_{4g} and R_{4k}).

The discontinuity behaviors of each region are as the following: for Region $1(v_1 \text{ and } v_3 \text{ modes})$, between 800 and 1100 cm⁻¹, the slope of wavenumber vs. pressure (dv/dP) is from 3.51–4.29 cm⁻¹/GPa to 2.99–5.59 cm⁻¹/GPa. Some new peaks show the slope (dv/dP) from 2.68 to 5.24 cm⁻¹/GPa. For Region 2, between 660 and 700 cm⁻¹, the slope of wavenumber vs. pressure (dv/dP) is from 2.68 cm⁻¹/GPa to 1.32 cm⁻¹/GPa, but shows a negative trend between 8 and 12 GPa. For Region 3 (v_2 and v_4 modes), between 420 to 660 cm⁻¹, the slope of wavenumber vs. pressure (dv/dP) is from 1.9–2.26 cm⁻¹/GPa to 0.78–1.89 cm⁻¹/GPa, and two peaks show a negative slope between 8 and 12 GPa. For Region 4, below 420 cm⁻¹, the slope of wavenumber vs. pressure (dv/dP) is from 0.13–3.14 cm⁻¹/GPa to -0.93–2.44 cm⁻¹/GPa.

Table 1. Major Raman bands for scottyite on a previous experiment study and on this Raman spectroscopic study.

Major Raman Bands	ν ₁ Strong, Sharp; SiO4 Symmetric Stretching	ν ₂ Very Strong, Sharp; SiO ₄ Symmetric Bending	$ u_3$ Relatively Weak; SiO ₄ Anti-symmetric Stretching	ν ₄ Relatively Strong, Sharp; SiO ₄ Anti-Symmetric Bending	R ₂ ¹ Strong, Sharp; Si-O-Si Bending Mode	${R_4}^1$ Strong to Weak; SiO ₄ Rotational and Translational Modes
Yang et al. [2]	896	459	1019, 958, 866	612, 578, 560	675	<420
This study	895	458	1016, 958, 863	608, 579, 560	675	358, 315, 303, 243, 220, 170, 159, 141, 120, 86, 75

 1 R_{2} , R_{4} means second and fourth region on the Raman spectra of scottyite.



Figure 1. A series of Raman spectra of scottyite showing the effect of pressure on the Raman modes. The v_1 and v_4 mode is the more intense Raman modes in scottyite. The distinct peak (new peak) is marked " \uparrow ". Arrows (\leftarrow , \rightarrow) indicate the direction of peak movement.



Figure 2. The variations in the wavenumber for each Raman mode of scottyite with pressure. (**a**) Wavenumber range: 420–1120 cm⁻¹; (**b**) wavenumber range: 50–420 cm⁻¹. Data for loading process are shown as solid symbols (two runs: red triangle and blue round shape) and those for unloading process as open symbols. The dotted line means the discontinuity of the slope on each Raman mode. There are some new peaks (green/pink square shape) during the loading process which are on v_1 , R_{4d} modes between 8 and 12 GPa and R_{4a} , R_{4c} , R_{4j} modes above 12 GPa. The solid lines are the eyeball linear fit of the data.

		1	'1		v_2 v_3			ν ₃ ν ₄				R_2
Kaman Mode	ν_1	New	New	v_{1a}	ν_2	v_{2a}	ν_3	v_{3a}	ν_4	New	v_{4a}	R_2
<8 GPa slope	4.27	-	-	4.12	1	.90	4.28	3.51	2.26	-	2.09	2.68
8–12 GPa slope	5.59	5.24	-	4.09	3.57	-1.33	4.29	-	-4.68	-	2.44	-3.35
>12 GPa slope	3.62	3.97	2.68	3.35	1.56	0.78	-	-	1.25	1.12	1.89	1.32
					-: no da	ta.						

Table 2. The variations in the slope $(d\nu/dP, \text{cm}^{-1}/\text{GPa})$ of each Raman mode in scottyite.

Table 3. The variations in the slope $(d\nu/dP, cm^{-1}/GPa)$ of each Raman mode in scottyite.

										R_4						
Raman Mode	New a	~	a b	с	N		N		f g					j	,	
		и		<i>c</i> 1	<i>c</i> 2	- New	а	New	e	f1	<i>f</i> 2	g1 g2	— n	1	$\begin{array}{c c} j \\ \hline j1 & j2 \\ \hline 0.96 & (0) \\ 1.79 & - \\ 2.35 & 0.95 & (0) \\ \end{array}$	ĸ
<8 GPa slope	-	3.14	2.44	2	.31	-	1.66	-	1.62		1.95	1.48	0.13	0.58	0.96	0.57
8–12 GPa slope	-	1.94	-	5.78	-4.27	-	-	1	1.43	5.54	-5.71	-3.72	1.69	3.91	1.79	-1.35
>12 GPa slope	1.18	1.53	-	2.44	-	0.92	-	1.75	1.07	1.76	0.19	-0.93 0.96	-	-	2.35 0.95	0.12
1.																

-: no data.

3.2. X-ray Diffraction Studies

A series of X-Ray Diffraction patterns are overlain to show the structural changes with pressure in scottyite in Figure 3. Several new peaks were observed above 8 GPa, which indicates the transformation of scottyite to high-pressure phase (Figure 3). The variations in d-spacings of the diffraction peaks as a function of pressure for scottyite are plotted in Figure 4. The calculated cell parameters, *a*, *b* and *c*, with the change in pressure are plotted in Figure 5, and tabulated in Table 4.



Figure 3. A series of X-Ray Diffraction patterns showing the sequences of phase transitions in scottyite during the loading and unloading processes. Scottyite exhibits four new peaks above 7–8 GPa (indicated by " \uparrow " and "*") suggesting the onset of phase transition. In addition, some peaks disappeared or merged (indicated by " \downarrow " and "#") when the pressure is over 10 GPa. The scottyite phase was recovered during the unloading process.



Figure 4. The variations in d-spacing for each diffraction peak in various phase of scottyite. (**a**) 2.6–6.8; (**b**) d-spacing range: 1.8–2.6. (*: Intensity weak, the trend is for reference only; solid circles: loading; open circles: unloading).

(a)

(b)



Figure 5. The compressibility of the lattice parameters (*a*, *b* and *c*) of scottyite with pressure up to 17 GPa (solid circles: loading; open circles: unloading).

Pressure (GPa)	а	b	С	V	a/a ₀	<i>b/b</i> 0	c/c_0	V/V_0
0.01	6.876	13.173	6.864	621.72	1	1	1	1
1.00	6.849	13.138	6.822	613.86	0.996	0.997	0.994	0.987
2.00	6.81	13.140	6.817	610.01	0.990	0.997	0.993	0.981
2.92	6.793	13.099	6.775	602.85	0.988	0.994	0.987	0.970
4.01	6.778	13.089	6.761	599.82	0.986	0.994	0.985	0.965
5.02	6.766	13.063	6.74	595.71	0.984	0.992	0.982	0.958
5.52	6.765	13.047	6.715	592.69	0.984	0.990	0.978	0.953
6.10	6.754	13.047	6.709	591.19	0.982	0.990	0.977	0.951
7.20	6.730	13.028	6.685	586.13	0.979	0.989	0.974	0.943
8.22	6.698	12.997	6.656	579.43	0.974	0.987	0.970	0.932
9.06	6.735	12.946	6.667	581.30	0.979	0.983	0.971	0.935
10.08	6.703	12.897	6.723	581.19	0.975	0.979	0.979	0.935
11.19	6.714	12.887	6.729	582.22	0.976	0.978	0.980	0.936
12.29	6.741	12.868	6.602	572.68	0.980	0.977	0.962	0.921
13.14	6.723	12.835	6.583	568.05	0.978	0.974	0.959	0.914
14.26	6.725	12.807	6.584	567.06	0.978	0.972	0.959	0.912
15.03	6.659	12.813	6.649	567.30	0.968	0.973	0.969	0.912
16.06	6.66	12.859	6.653	569.77	0.969	0.976	0.969	0.916
16.40	6.498	12.883	6.688	559.88	0.945	0.978	0.974	0.901
			Unle	oading				
12.29	6.752	12.935	6.612	577.47	0.982	0.982	0.963	0.929
8.05	6.737	13.002	6.684	585.48	0.980	0.987	0.974	0.942
6.12	6.739	13.014	6.693	586.99	0.980	0.988	0.975	0.944
4.02	6.763	13.075	6.758	597.58	0.984	0.993	0.985	0.961
1.08	6.833	13.170	6.813	613.11	0.994	1.000	0.993	0.986
0.01	6.878	13.188	6.864	622.61	1.000	1.001	1.000	1.001

Table 4. The unit cell parameters, molar volume and their normalized ratios of scottyite in the loading and unloading processes.

At first glance, the diffraction patterns of scottyite shown in Figure 3 do not seem to show a very significant change with pressure. However, closer examination of these patterns reveals some minor changes during the compression process. With an increase in pressure, we observed that four new diffraction peaks show up near 8 GPa. Figures 3 and 4a,b display four new peaks in the intervals between (031) and (210); (040) and (102); near (051) and (250) diffractive planes, the d-spacing of which is near 3.5, 3.1, 2.6 and 2.2 Å, respectively, above 8 GPa. Some peaks either disappear or were merged with other peaks when the pressure was over 10 GPa, such as (031), (210), (201), (022), (151) and (321) diffractive planes. The appearance of more diffraction peaks implies that the high-pressure phase of scottyite has a lower symmetry.

The variations in a/a_0 , b/b_0 , c/c_0 and molar volume with pressure for scottyite are listed in Table 4. The compression data are fitted to the Birch–Murnaghan equation (Equation (1)).

$$P = \frac{3}{2} Ko \left[(Vo/V)^{7/3} - (Vo/V)^{5/3} \right] \left\{ 1 - \frac{3}{4} (4 - Ko') \left[(Vo/V)^{2/3} - 1 \right] \right\}$$
(1)

We obtain a bulk modulus (K_o) value of scottyite as 102 ± 5 GPa (assuming that K_o' is 4) (Figure 6).



Figure 6. The compression data as fitted to the Birch–Murnaghan equation, to give a bulk modulus, K_o , of 102 GPa, assuming $K_o' = 4$. Above 8 GPa, (assuming that the structure remains unchanged) it is clearly seen that all the data deviate from the compression curve.

4. Discussion

4.1. Anisotropic Compressibility

We found that the *c*-axis was the most compressible axis in scottyite, the *b*-axis was the least, and the *a*-axis was similar to the *c*-axis on the compression (Figure 5 and Table 4). The scottyite structure is based on a tetrahedral framework consisting of SiO₄ and CuO₄ tetrahedrons. The CuO₄ tetrahedrons are considerably flattened, and they share corners to form chains parallel to the *c*-axis. The chains are interlinked by the Si₂O₇ dimers oriented parallel to the *b*-axis. [2,6,7] The largest bond distance of M-O was Ba-O (2.825 Å) where Ba²⁺ cations are in the framework channels. The SiO₄ is a rigid body and is linked to the Si₂O₇ dimer, so it is hard to compress. However, CuO₄ tetrahedron layers are parallel to the *a*-axis and *c*-axis relatively easy to compress.

4.2. The Pressure of the Phase Transition

The four new peaks in the diffraction patterns and the discontinuities in the slopes of dspacing versus pressure plot all occurred at approximately 7–8 GPa, which is quite distinct for scottyite. Figure 5 shows the variations in the *a*-, *b*- and *c*-parameters of scottyite with pressure, assuming that the scottyite structure remains during the loading and unloading processes. From Figure 5, we observed a kink in their compression curves at about 8 GPa in all three crystallographic axes. Above 9 GPa, the lattice parameters behave in a less compressible way, and show larger values than those extrapolated from the low pressures, which is not reasonable. The offset of the lattice parameters on the *a*-axes and *c*-axes direction is similar. All the lattice parameters are reversible under the unloading process. The deviation from the normal compression behavior in scottyite is more clearly seen in Figure 6. Below 8 GPa, the compression data can be fitted to a bulk modulus of 102 GPa with $K_o' = 4$ by the Birch–Murnaghan equation (Equation (1)), but the data show a large deviation from the normal compression curve above 8 GPa. Some XRD peaks disappeared or merged with other peaks when the pressure exceeded 10 GPa.

In addition, we also found the discontinuities of each Raman mode in the $d\nu/dP$ plot at near 8 GPa, and some Raman modes showed negative slop of $d\nu/dP$ between 8 and 12 GPa. Most of the trend with the negative slop of $d\nu/dP$ belonged to R_2 (Si-O-Si bending mode), ν_4 (SiO₄ anti-symmetric bending) and ν_2 (SiO₄ symmetric bending) which deserves further investigation in order to understand the mechanism of the phase transition. We conclude that scottyite has encountered a structural phase transition at 8 GPa and the transition is reversible.

4.3. Post-Scottyite Structure Analysis

Preliminary high-pressure Raman and X-ray investigation provided evidence that a reversible phase transition takes place at 8 GPa in scottyite. We have tried to analyze the post-scottyite structure with GSAS-II program [13], and the most suitable candidate for the high pressure phase is maybe the monoclinic system of the Bravais lattice C2/m, with unit-cell parameters of a = 5.67 Å, b = 5.12 Å, c = 18.58 Å, $\beta = 91.64^{\circ}$ and molar volume of V = 539.3 Å³ at 11.19 GPa. The fitting pattern is shown in Figure 7. However, since there are still many peaks that cannot be matched, the structure that we proposed above may need further experimental data and analysis to justify it.



Figure 7. Comparison of the XRD pattern at 11.19 GPa with the diffraction peaks of candidate high-pressure phase obtained by GSAS-II. The blue lines are the XRD peak positions at 11.19 GPa and the orange lines are the diffraction peaks obtained by GSAS-II. The inserted figure is the entire diffraction pattern.

A comparison of crystallographic data of $BaM_2Si_2O_7$ -type (M = Be, Mg, Mn, Fe, Co, Zn, and Cu) minerals [2,14–18] shows that the structure with the monoclinic system is M = Mg, Mn, Fe, Co, and Zn. There is a strong resemblance among them in the structure (Table 5). The radius of Cu²⁺ is similar to Mg²⁺, Co²⁺ and Zn²⁺, so we speculate that the post-scottyite phase may be similar among these compounds. However, this still needs to be verified by a single-crystal X-ray diffraction experiment.

Table 5. The crystal structure of some BaM₂Si₂O₇-type minerals with the monoclinic system.

Charry i and Formurale	Space		Unit-Cell	Parameters	5	Si-O-Si (°)
Chemical Formula	Group	а	b	С	β	
BaFe ₂ Si ₂ O ₇ [14]	$P2_{1}/c$	7.488	13.785	7.085	118.23	127.2
BaCo ₂ Si ₂ O ₇ [15]	C2/c	7.213	12.781	13.762	90.299	124.5
BaMg ₂ Si ₂ O ₇ [16]	C2/c	7.246	12.714	13.748	90.211	125.2
BaMn ₂ Si ₂ O 7 [17]	C2/c	7.295	12.963	14.032	90.248	no data
BaZn ₂ Si ₂ O ₇ -25 °C [18]	C2/c	7.278	12.801	13.687	90.093	124.8

From the Table 5, we know that the angle of Si-O-Si in the monoclinic $BaM_2Si_2O_7$ -type structure is between 124.5° and 127.2°, which is smaller than in the scottyite structure

(134.3° [2]). In other words, the compression deformation of the scottyite crystal maybe mainly occurs in Si-O-Si linkage formation, and thus results in CuO_4 tetrahedra layer were compressed and adjusted. This then makes the *a*-axis and *c*-axis directions the most to be compressed.

The compression behavior of scottyite may be applied to silicates of similar structure, which is of significant meaning in the understanding of bonding characteristics of sorosilicates. Despite the fact that the structure of the post-scottyite phase has not been solved in this study, it is feasible that detailed single crystal XRD experiments in the future will help to resolve it.

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