RSC Advances



PAPER

View Article Online
View Journal | View Issue

and Zheshuai Lin*c



Cite this: RSC Adv., 2018, 8, 9946

structure variation of chalcopyrite type LiGaTe₂†

V. V. Atuchin,*ab Fei Liang, S. Grazhdannikov, L. I. Isaenko, Atuchin, S. Molokeev, P. G. Krinitsin, M. S. Molokeev, P. Prosvirin, Xingxing Jiang^c

Negative thermal expansion and electronic

The LiGaTe $_2$ crystals up to 5 mm in size were grown by the modified Bridgman–Stockbarger technique and the cell parameter dependence on temperature in the range of 303–563 K was evaluated by the X-ray diffraction analysis. The thermal behavior of LiGaTe $_2$ is evidently anisotropic and a negative thermal expansion is found along crystallographic direction c with coefficient -8.6×10^{-6} . However, the normal thermal expansion in two a directions with coefficient 19.1×10^{-6} is dominant providing unit cell volume increase on heating. The atomic mechanism is proposed to describe this pronounced anisotropic expansion effect. The electronic structure of LiGaTe $_2$ is measured by X-ray photoelectron spectroscopy and the band structure is obtained by DFT calculations. The pressure response from 0 to 5 GPa was calculated and a normal crystal compression is found. This work indicates that LiGaTe $_2$ is promising as an IR NLO or window material for many practical applications because the thermal expansion coefficients of this telluride are not big. We believe that these results would be beneficial for the discovery and exploration of new IR optoelectronic polyfunctional metal tellurides.

Received 3rd February 2018 Accepted 5th March 2018

DOI: 10.1039/c8ra01079j

rsc.li/rsc-advances

Introduction

Telluride crystalline materials are of great importance in modern microelectronics and photonics.^{1–12} Because of specific features of metal-tellurium chemical bonds, tellurides possess narrow-bandgap semiconductor or metallic properties.^{1,2,6–8,11,12} Due to their low-wavenumber phonon spectrum, semiconductor tellurides are transparent in the mid-IR range and they can be used as optical window materials. However, the number of known noncentrosymmetric tellurides is not big

and, only for several crystals, the linear and nonlinear properties are obtained. $^{13-22}$ The high potential of LiGaTe $_2$ for the optical frequency conversion in the mid-IR spectral range has been demonstrated in several studies, and the structural and optical properties of this material were measured. 13,16,17,20,23

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c8ra01079j

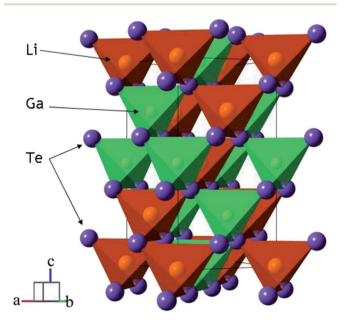


Fig. 1 The crystal structure of $LiGaTe_2$ chalcopyrite. The unit cell is outlined. Lone lithium and gallium atoms are omitted for clarity.

^aLaboratory of Optical Materials and Structures, Institute of Semiconductor Physics, SB RAS, Novosibirsk 630090, Russia. E-mail: atuchin@isp.nsc.ru

^bLaboratory of Semiconductor and Dielectric Materials, Novosibirsk State University, Novosibirsk 630090, Russia

^{&#}x27;Key Laboratory of Functional Crystals and Laser Technology, Technical Institute of Physics and Chemistry, Chinese Academy of Sciences, Beijing 100190, China. E-mail: zlin@mail.ipc.ac.cn

⁴Laboratory of Crystal Growth, Institute of Geology and Mineralogy, SB RAS, Novosibirsk 630090, Russia

^eLaboratory of Functional Materials, Novosibirsk State University, Novosibirsk 630090, Russia

Laboratory of Crystal Physics, Kirensky Institute of Physics, Federal Research Center KSC SB RAS. Krasnovarsk 660036. Russia

^{*}Department of Physics, Far Eastern State Transport University, Khabarovsk 680021, Russia

^hSurface Science Laboratory, Boreskov Institute of Catalysis, SB RAS, Novosibirsk 630090, Russia

Paper

 ${\rm Li_2Te}$ to the starting stoichiometric composition at the level up to 16 at%. To avoid the material contamination by lithium oxide and/or nitride, the sample handling was produced in the Ar atmosphere in a specialized dry box. Upon the completion of the active synthesis stage, the melt was maintained at the melting

temperature for 24 h.

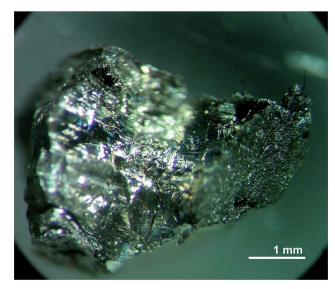
LiGaTe2 is a positive uniaxial crystal and it has a chalcopyrite type structure ($I\bar{4}3d$ with the closest packing).²⁴ The structure of LiGaTe₂ is shown in Fig. 1.16,25 Each Li, Ga and Te atom occupies only one crystallographic position, the Li-Te bond distance is 2.736 Å and, for the Ga-Te bond, the value is 2.611 Å. For comparison, in the LiGaSe₂ structure (wurtzite type, Pna2₁) there are two positions of Se. The Li-Se bond distance changes from 2.493 to 2.565 Å, while the Ga-Se bonds range from 2.389 to 2.405 Å. The nonlinear crystal LiGaTe2 is characterized by the biggest band-gap energy of $E_{\rm g}=2.41$ eV among tellurides, which is essentially bigger than those of LiInTe₂ (1.5 eV), AgGaTe2 (1.32 eV) and AgInTe2 (1.03 eV). A sufficiently large birefringence (0.094) for LiGaTe₂ allows one to obtain its roomtemperature phase-matching in the whole transparency range (1.66–21 μ m). Its nonlinear coefficient d_{36} was estimated by the phase-matched second harmonic generation to be 43 pm V⁻¹, which is about 7 times higher than that of the related orthorhombic LiGaS2 and about 4.5 times relative to LiGaSe2. The conversion efficiency in LiGaTe2 for SHG 10.6 µm was found to be higher than that of AgGaSe2.16 Besides, this telluride is considered as a promising crystal for neutron detection.26,27 However, in several studies, it was reported on that crystal growth of LiGaTe2 is not trivial because of its high tellurium volatility and pronounced chemical activity of lithium at high temperatures. 20,28-30 This may result in the defect generation and stoichiometry deviations, and these effects should be accounted to reach the optical quality crystals.

The second stage was the melt homogenization in a sealed quartz ampoule with an inner pyrolytic carbon coating. The homogenization temperature did not exceed 50 K above the melting point of LiGaTe2. This level was selected because the incongruent evaporation of the melt appeared above this temperature.29,30 The LiGaTe2 crystals up to 5 mm in size were grown from the homogenized melt by the Bridgman-Stockbarger technique in the two-zone vertical resistance furnace with the diaphragm in a sealed pyrocarbon inner-coated quartz ampoule. The ampoule was first evacuated to the residual pressure of 10⁻³ atm and then filled with a high purity Ar (P = 0.2 - 0.5 atm). The temperature at the start of crystallization was a few degrees below the LiGaTe2 melting point. The maximum gradient near the crystallization front was 2 K mm^{-1} and the ampoule movement rate to the cold zone was 2.5 mm per day. The crystal growth was performed for 20 days. The photo image of the grown crystal is shown in Fig. 2.

For a practical NLO crystal, the crystal growth and optical response are equally important. Consequently, the thermal and chemical properties, including electronic structure analysis, play the most crucial role. However, only a few theoretical studies focus on these issues.31-36 To date, there are no reports about the dependence of structural parameters of LiGaTe₂ on temperature and experimental measurements of electronic structure. Thus, the present study is aimed at the LiGaTe2 crystals growth and observation of the electronic structure in parallel by the X-ray photoelectron spectroscopy (XPS) and theoretical methods, where XPS is extremely sensitive to chemical state of the crystal surface and theoretical model is a power tool to consider the relations between crystal structure and physical properties. Besides, the dependence of structural parameters of LiGaTe2 on temperature is explored combining experimental and theoretical methods.

The powder diffraction data of LiGaTe₂ were collected at room temperature with a Bruker D8 ADVANCE powder diffractometer (Cu-K α radiation) and linear VANTEC detector. The step size of 2θ was 0.016° and the counting time was 1.5 s per step. The 2θ range of $5-70^{\circ}$ was measured with the 0.6 mm divergence slit, but the 2θ range of $70-140^{\circ}$ was measured with the 2 mm divergence slit. The intensities and obtained esd's were further normalized: $I_{i \text{ norm}} = I_{i} \times 0.6/\text{(slit width)},$ $\sigma_{\text{norm}}(I_{i}) = \sigma(I_{i}) \times 0.6/\text{(slit width)},$ taking into account the actual divergence slit width value which was used to measure each particular intensity I_{i} , and saved in the xye-type file. So, the transformed powder pattern had a usual view over the whole 2θ range of $5-140^{\circ}$, but all high-angle points had small esd values.

2. Experimental methods



LiGaTe₂ crystals were obtained in three stages.³⁷ At the first stage, we obtained the charge by fusing the elementary components Li, Ga, and Te in a glass carbon crucible inside the sealed quartz ampoule. The ampoule was evacuated to the residual pressure of 10^{-3} atm. The synthesis temperature was 1250 K, the initial components purity was Li 99.99%, for Ga and Te – 99.9999%. All the metal reagents were supplied by Alfa Aesar. The accuracy of reagent mass measurements was ± 4 mg. The pyrosynthesis reaction proceeds with the liberation of a large amount of heat. At the conditions, Li reacts with the crucible walls and it is partially consumed. To compensate this effect, we introduced an excess of

Fig. 2 Photo image of the LiGaTe₂ crystal.

RSC Advances Paper

Additionally, to see the structural parameters dependence on temperature, fourteen X-ray diffraction (XRD) patterns in the range of $2\theta = 5-120^{\circ}$ were collected at fourteen different temperatures in the range from 303 K to 563 K spending 35 min for each pattern. The measurements were carried out using the temperature set TTK450 (Anton Paar). The recorded XRD patterns are shown in Fig. 3. It was found that the sample decomposes at a low rate in the temperature range of 303-443 K, but, after that, the rate increases and the impurity peaks prevent a good fitting. Therefore, the XRD patterns from 443 K to 563 K were recorded from a new LiGaTe2 powder sample. However, the decomposition at 583 K also leads to a large amount of Te impurity, and a further heating leads to the total sample decomposition and we stop the experiment.

The XPS spectra were recorded using a SPECS (Germany) photoelectron spectrometer equipped with a hemispherical PHOIBOS-150-MCD-9 analyzer and FOCUS-500 (Al Ka radiation, $h\nu = 1486.74$ eV, 200 W) monochromator. Just before the measurements, the binding energy (BE) scale was calibrated using the positions of the peaks of Au $4f_{7/2}$ (BE = 84.0 eV) and Cu $2p_{3/2}$ (BE = 932.67 eV) core levels. The LiGaTe₂ powder sample was prepared for the XPS experiment. To decrease a possible contamination by air agents, the LiGaTe2 crystal was ground in agate mortar into the glove box filled with high-purity argon. In this way, however, the chemical reaction between LiGaTe₂ and agate species can not be excluded. The LiGaTe₂ powder sample was loaded onto a conducting double-sided copper scotch and transfered into the input vacuum chamber without a contact with the air environment. The details of the sample handling and BE energy scale calibration methods can be found elsewhere. 38-40 The base pressure of a sublimation ionpumped chamber of the system was less than 5×10^{-10} mbar during the present experiments. Besides the survey photoelectron spectra, the narrow spectral regions Li 1s, Te 3d, Ga 2p, Ga 3d, Te 4d and the valence band were recorded. The survey spectra were taken at the analyzer pass energy of 50 eV and the detailed spectra were registered at 20 eV. The concentration ratios of the elements on the sample surface were calculated from the integral photoelectron peak intensities which were

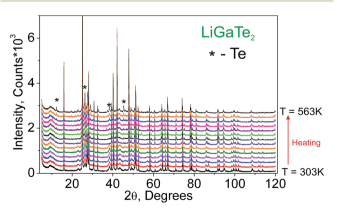


Fig. 3 The XRD patterns recorded from the LiGaTe₂ sample in the range from 303 K to 563 K. The impurity peaks related to the Te component are marked by the asterisk.

corrected with the theoretical sensitivity factors based on Scofield photoionization cross sections.41 For the peak fitting procedure, a mixture of Lorentzian and Gaussian functions was used together with the Shirley background subtraction method.

3. Ab initio calculation methods

The electronic band structures calculations for LiGaTe2 were performed by a CASTEP package.42 The generalized gradient approximation (GGA-PBE) functionals⁴³ were selected to be the exchange-correlation (XC) functional. The ion-electron interactions were described by optimized norm-conserving pseudopotentials.44 The orbital electrons of Li 2s¹, Ga 3d¹⁰4s²4p¹, Te 5s²5p⁴ were treated as valence electrons and the correlation energy (U = 5.0 eV) was used to describe the Hubbard U of Ga 3d electrons. The kinetic energy cutoff of 880 eV and Monkhorst-Pack k-point meshes⁴⁵ spanning less than 0.04 \mathring{A}^{-3} (4 × 4 × 2) in the Brillouin zone were chosen to ensure the sufficient accuracy of current methods. The hydrostatic pressure (from 0 to 5 GPa; step 0.5 GPa) was applied on the LiGaTe₂ crystal and geometry optimization calculations were performed until the energy change, the maximum force and maximum displacement were less than 5.0×10^{-6} eV per atom, 0.01 eV Å⁻¹, and 5.0×10^{-4} Å, respectively. The convergence tests revealed that the above computational parameters are accurate enough for the purposes of this study. The phonon dispersion spectra of LGT were calculated by the linear response method.46 Notably, the primitive cell constants under different temperatures were fixed in phonon calculations.

4. Results and discussion

4.1. Temperature effect

The crystal structure refinement was performed by the package DDM which accounts the esd values of each point by a special weight scheme.47 The powder pattern has a complex background curve due to the existence of an amorphous phase in the sample, and this was the reason for applying the DDM program instead of commonly used Rietveld programs. All diffraction peaks were indexed by the tetragonal cell $(I\bar{4}2d)$ with parameters close to those of LiGaTe2.16 Therefore, the crystal structure shown in Fig. 1 was taken as a starting model for the refinement. The thermal parameter of Li was not refined and other

Table 1 Main parameters of processing and refinement of the LiGaTe₂ sample

Compound	${ m LiGaTe}_2$
Space group	$Iar{4}2d$
a, Å	6.33757 (2)
c, Å	11.70095 (5)
V , \mathring{A}^3	469.966 (4)
Z	4
2θ -interval, $°$	5-140
R_{DDM} , %	9.48
$R_{\rm exp}$, %	6.98
R_{exp} , % χ^2	1.36
R_{B} , %	4.74

Paper

0.3 LiGaTe₂ Intensity, Counts*10 4 0.2 0.1 60 80 100 120 20 40 60 80 100 120 140 2θ, Degrees

Fig. 4 Difference Rietveld plot of LiGaTe₂.

ions were refined with isotropic thermal parameters. The refinement was stable and it gave low *R*-factors, as evident from Table 1 and Fig. 4. The atom coordinates and main bond lengths are summarized in Tables S1 and S2,† respectively.

The additional fourteen powder patterns recorded at the temperatures from 303 K to 563 K were also treated by the DDM program and this gave the thermal dependencies of unit cell parameters and cell volume with a good reliability (Fig. 5, Table S3 \dagger). With the temperature increase, one can see a strong increase of a cell parameter, but cell parameter c noticeably decreases. This means that the thermal behavior of LiGaTe₂ is evidently anisotropic and the negative thermal expansion is found in crystallographic direction c. However, the normal thermal expansion in two a directions is dominant; thereby, the cell volume still increases on heating, as shown in Fig. 5c. The thermal expansion coefficients obtained for LiGaTe₂ are summarized in Table 2. Thus, this telluride should be classified as an anisotropic positive thermal expansion (APTE) material.

It is interesting to consider the atomic mechanism of this pronounced anisotropic expansion effect. The dependences of d(Ga-Te) and d(Li-Te) bond lengths on temperature are shown in Fig. 6. Accordingly, as the temperature increase from 303 to 563 K, the Ga-Te bond length fluctuation is only 0.2% (Fig. 6a). This implies that the Ga-Te bond length is almost persistent, manifesting the rigidity of GaTe₄ tetrahedra when being heated. In comparison, the Li-Te bond length exhibits a higher flexibility and it increases from 2.751 to 2.775 Å (by 0.9%, almost five times than that of the Ga-Te bond, Fig. 6b). Coupled with the Li-Te bond elongation, the neighboring rigid GaTe₄ tetrahedra rotate with each other, leading to the enlargement of the Ga-Ga-Ga angle from 94.57° to 95.06° (Fig. 6c). This effect directly results in the increase of the projection along the a-axis and the contraction of that along c-axis in the Ga-Ga distance with the increasing temperature, giving rise to the positive and negative thermal expansions of these two dimensions, respectively. Therefore, the APTE behavior of LiGaTe₂ is mainly ascribed to the synergetic effect between the configuration modification of "rigid" GeTe4 and "flexible" LiTe4 tetrahedra, as depicted in Fig. 7.

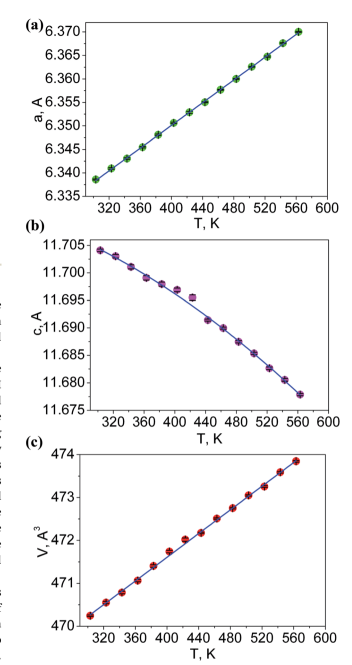


Fig. 5 Temperature dependences of cell parameters: (a) a; (b) c; (c) cell volume V.

Table 2 Thermal expansion coefficients in LiGaTe₂

Crystallographic parameter	Thermal expansion coefficient, K ⁻¹	
a c V	$19.1 \times 10^{-6} \\ -8.6 \times 10^{-6} \\ 29.4 \times 10^{-6}$	

As it is known, the thermal expansion response is strongly related to intrinsic phonon vibrational modes. In order to elucidate the mechanism of negative thermal expansion along

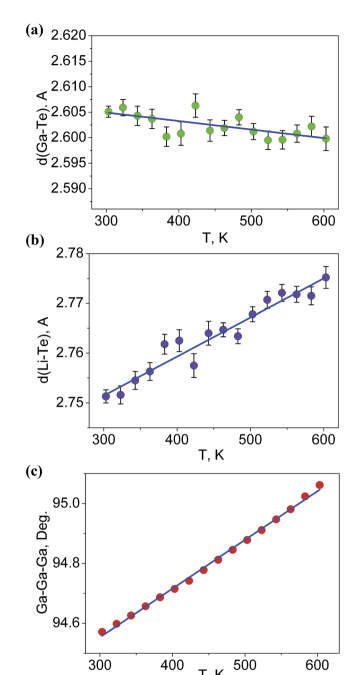


Fig. 6 The dependences of (a) d(Ga-Te), (b) d(Li-Te) bond lengths and (c) Ga-Ga-Ga angle on temperature.

the c axis, we performed the phonon calculations of LGT at different temperatures (303, 403, 503 and 603 K) (Table 3). The lattice parameters are fixed and the atomic positions are fully relaxed in calculations. The relationship between the Grüneisen parameter and thermal expansion coefficient is given as:

$$\alpha = \frac{C_{\rm V}\rho}{K_{\rm T}}\gamma$$

where α is the thermal expansion coefficient, $C_{\rm v}$ is the principle heat capacity, ρ is density, $K_{\rm T}$ is isothermal bulk moduli, γ is the macroscopic Grüneisen parameter. The macroscopic Grüneisen

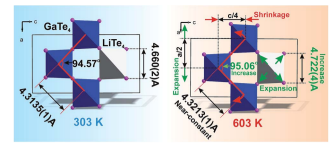


Fig. 7 The model which explained the increase of a cell parameter and the decrease of c cell parameter in LiGaTe $_2$ on heating. The bond length d(Li-Te) increases, but d(Ga-Te) lengths stay almost unchangeable under heating. This leads to the asymmetrical deformation of the $(\text{GaTe}_4)_3(\text{LiTe}_4)$ ring and the increase of Ga-Ga-Ga angle which is responsible for the expansion of parameter a and the shrinkage of the parameter c.

Table 3 The detailed atomic vibrations (cm⁻¹) for the optical modes by the first-principles calculations

Mode	303 K	403 K	503 K	603 K	
E	54.099	54.249	54.379	54.346	
E	54.099	54.249	54.379	54.346	
\mathbf{B}_2	75.147	75.198	75.308	75.321	
E	79.036	79.157	79.263	79.381	
E	79.036	79.157	79.263	79.381	
B_1	84.277	84.360	84.508	84.643	
A_2	94.289	93.849	93.444	93.083	
A_2	117.890	117.494	117.150	116.817	
A_1	139.510	139.153	138.849	138.488	
\mathbf{B}_2	186.916	186.231	185.674	185.160	
B_1	192.383	191.624	191.000	190.352	
E	199.834	199.356	198.992	198.668	
E	199.834	199.356	198.992	198.668	
E	208.451	208.116	207.867	207.655	
E	208.451	208.116	207.867	207.655	
B_2	300.283	297.476	294.839	292.215	
E	306.310	303.520	300.971	298.416	
E	306.310	303.520	300.971	298.416	
E	311.169	308.399	305.858	303.364	
E	311.169	308.399	305.858	303.364	
B_1	312.970	310.324	307.852	305.353	

parameter γ is the sum of the microscopic Grüneisen parameters (γ_i) of the phonon model by weight. And then, the microscopic Grüneisen parameter of an individual vibrational mode with number i can be defined as (the negative of) the logarithmic derivative of the corresponding frequency ω_i :

$$\gamma_i = -rac{V}{\omega_i}rac{\partial \omega_i}{\partial V}$$

and *V* is the volume. So, if the frequency of a vibrational mode increases with the increasing volume (increasing temperature), this phonon mode would contribute to the negative thermal expansion (NTE).

For a chalcopyrite structure with space group $I\bar{4}2d$, symmetry allows the optical vibrations to be classified into 15 normal modes: $\Gamma = A_1 + 2A_2 + 3B_1 + 3B_2 + 6E$, where B_2 and E are polar Raman active modes having transverse optical (TO) and

longitudinal optical (LO) modes, A_1 and B_1 are nonpolar Raman active modes, and A_2 is an optically inactive mode. As seen in Table 3, six vibrational modes (low frequency region) exhibit an anomalous wavenumber increase on the increasing temperature. So, we can assigned the negative thermal expansion of LGT to these vibration modes. As shown in Fig. S1,† the B_2 modes located at 75.14 cm $^{-1}$ (300 K) mainly contribute to the rotation of (GaTe₄) tetrahedra, coupling the Li⁺ slide migration along the c axis. Therefore, the negative thermal expansion of LGT along the c axis can be attributed to the tetrahedra rotation, combining the experimental measurements and theoretical calculations.⁴⁸

4.2. Comparison between XPS spectra and band structure

The survey photoemission spectrum recorded from the LiGaTe₂ sample is shown in Fig. S2.† All detected spectral features were successfully attributed to constituent element core levels and Auger lines, except for the weak C 1s and O 1s bands shown in Fig. S3 and S4,† respectively. A comparatively narrow dominant component of the C 1s line (284.8 eV) was related to adventitious hydrocarbons adsorbed at the surface from the air, and the low-intensity additive component at 289.7 eV was attributed to carbonate species. In the O 1s line, two components can be revealed. The lower binding energy component at \sim 530.4 could be attributed to the surface oxide, and a higher binding energy component appeared at \sim 531.5 eV could be related to the initial stages of surface hydration. Thus, it can be concluded that LiGaTe2 is not chemically inert and it actively interacts with atmospheric agents. From the technological point of view, this should be accounted for at crystal cutting and polishing stages.

The valence band spectrum and the nearest constituent element core levels are shown in Fig. 8. The mixed-states bands with several less pronounced components are found over the binding energy range of 0–17 eV. At \sim 19.5 and \sim 40 eV, the bands related to Ga 3d and Te 4d are observed, respectively. In Fig. 9–11, the representative element core levels Li 1s, Te 3d and Ga 2p are shown. The Li : Ga : Te ratio was estimated by Li 1s,

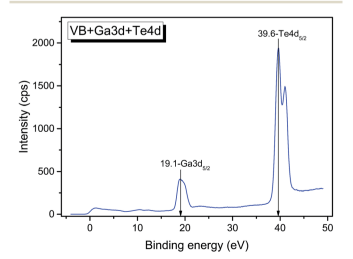


Fig. 8 Detailed XPS spectrum of the Ga 3d and Te 4d doublets from $LiGaTe_2$.

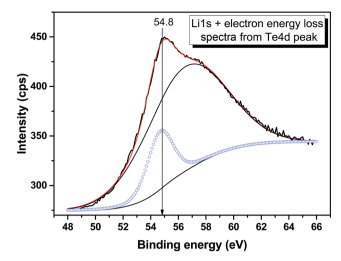


Fig. 9 Detailed XPS spectrum of the Li 1s core level from LiGaTe₂.

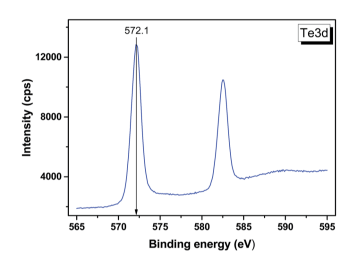


Fig. 10 Detailed XPS spectrum of the Te 3d doublet from LiGaTe₂.

Te 4d and Ga 3d peak areas and tabulated atomic sensitivity factors. The relative element ratio for the powder sample is Li: Ga: Te = 0.38:0.22:0.40 and that indicates a noticeable enrichment of the LiGaTe₂ surface by Li. Previously, a similar effect was observed for closely related compound LiGaS₂. Thus, the Li segregation to the crystal surface may be a general feature of compounds LiGaX₂ (X = S, Se, Te). The ratio Ga: Te = 0.55 is in reasonable consistence with nominal composition Ga: Te = 0.50. The calculations were made without carbon signal accounting. The set of the element core levels measured for LiGaTe₂ is shown in Table 4. Comparatively, the BE values obtained for Li 1s, Te 3d and Ga 2p core levels in LiGaTe₂ are in a good relation with the BE values obtained earlier in several other representative compounds.

The calculated electronic total and partial densities of states (DOS and PDOS) projected on the constituent elements in LiGaTe₂ are displayed in Fig. 12. LiGaTe₂ is non-magnetic compound, so it spin-up and spin-down band structure (and DOS) are degenerated. Therefore, we only plotted one spin tunnel component for clarity. It is noted that the Te 4d orbitals

RSC Advances

8x10³ - (80) 7x10³ - (5x10³ - (5x10³

Fig. 11 Detailed XPS spectrum of the Ga 2p doublet from LiGaTe₂.

cannot be simulated by DFT because they are inner-shell electron levels and there are no suitable pseudopotentials that take them into account. As displayed in Fig. 12a, the XPS spectrum

and electron density of states are reasonably matched in the energy region from -20 to 0 eV. This confirms the validity of the plane-wave pseudopotential method in its application to LiGaTe₂. Some bond characteristics can be deduced from the PDOS: (i) the Ga 3d orbitals are strongly localized at -19.1 eV, which is consistent with the calculated strong DOS peak at -18.16 eV. It hybridizes little with the orbitals of other elements, suggesting that Ga 3d electrons are not bonded with Te orbitals. Notably, the Hubbard U values mainly changes the position of Ga 3d orbitals (from -15.2 eV at U = 0 eV to -18.16 eV at U = 5 eV); (ii) the Te 5s orbitals are relatively localized at the inner energy level around -10 eV, which is slightly hybridized with Ga 4s and 4p electrons; (iii) from 10 to 0 eV, the electron states are mainly composed of Ga 4s/4p and Te 5s/5p orbitals. The strong hybridization in the wide energy range between these orbitals indicates the strong covalent interaction between Ga and Te atoms. From -10 to -8 eV, Te 5s orbitals play a crucial role. Meanwhile, Ga 4s and Te 5p orbitals make the main contribution to the hybridization interaction. Notably, the valence band maximum is mainly composed of Ga 3p and Te 5p orbitals. In addition, Li 1s orbitals are only located

Table 4 Binding energies of the constituent element core levels in LiGaTe₂

Binding energy (eV)

Core level	Binding energy, eV					
	${\rm LiGaTe}_2$	$LiGaS_2$	GaTe	PbTe(100)	Bi ₂ Te ₃ (001)	
Ga 3d	19.1	19.7	19.5	_	_	
Te 4d _{5/2}	39.6	_	_	_	_	
Te 4d _{3/2}	41.0	_	_	_	_	
Li 1s	54.8	54.9	_	_	_	
C 1s	Fixed at 284.8, 289.7	Fixed at 284.6, 288.4	_	_	Fixed at 284.8	
O 1s	531.5	531.5	530.8	530.0	_	
Te 3d _{5/2}	572.1	_	573.1	571.9	572.5	
Te 3d _{3/2}	582.5	_	583.5	_	582.8	
Ga 2p _{3/2}	1117.3	1117.6	1117.9	_	_	
Ga 2p _{1/2}	1144.2	114.5	_	_	_	
Reference	This study	49	50	51	52	

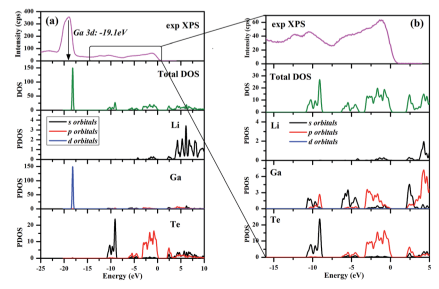


Fig. 12 Electronic total and partial densities of states (DOS and PDOS) in LiGaTe₂.

at the valence band maximum with a negligible contribution; (iv) the bottom of the conduction band minimum is mainly contributed from the Ga 4s and Te 5p orbitals. Since the optical effects of a crystal are mainly determined by the optical transition between the electronic states close to the bandgap, it is anticipated that they are dominantly contributed from [GaTe₄] tetrahedra, and the contribution from the orbitals of Li⁺ cations is negligibly small.

4.3. Optical properties

Paper

LiGaTe₂ is a direct gap crystal with the calculated bandgap (1.85 eV) (Fig. S5†), which is slightly smaller than the experimental value (2.30 eV) owing to the well-known band gap underestimation by the GGA exchange-correlation (XC) functional.⁵³ Therefore, a scissors operator⁵⁴ is usually introduced to shift all the conduction bands in order to agree to the experimental band gap value for calculating the optical coefficients. The calculated second order NLO coefficient is $d_{36} = -52.7 \text{ pm V}^{-1}$, which is comparable to other calculated results.55 If adopting GGA+U method, the calculated d_{36} is -49.6 pm V⁻¹. The Hubbard U mainly changes the position of Ga 3d orbitals but no influence on the electric bands near forbidden gap (see Fig. S6†). Because the optical effects of a crystal are mainly determined by the optical transition between the electronic states close to the band gap, accordingly, it is anticipated that the effects of Hubbard U are negligibly small on optical properties. In addition, the birefringence is calculated to be 0.035 (see Fig. S7†), which is smaller than the experimental results, 16 but it agrees well to other calculated results.34,55

4.4. Pressure effect

The SHG coefficients of NLO materials are highly dependent on the band gap energy. Band gap engineering is generally achieved through lattice distortion by the impurity doping that results in changing the material composition. Since this method often leads to some negative change that could not be controlled well, material compression under high pressure is a remarkable alternative for changing the lattice parameters while keeping the element composition. To identify the high pressure effects on the properties of the LiGaTe2 crystal, the lattice constants and band gap energy dependence on the pressure applied uniformly is also investigated. The crystal lattice constants (a and c axis) and the band gap energy of the LiGaTe₂ at different pressures are shown in Fig. 13. The a and c axes compression up to 5% and 3%, respectively, refers to the pressure uniformly applied on the crystal in the range of 0-5 GPa. The LiGaTe2 band gap decreases monotonically with the increasing pressure, which is similar to that of Li₂S⁵⁶ and As₂S₃.⁵⁷ For the As₂S₃ crystal, the optical band gap redshifts rapidly with the increasing pressure, decreasing from 2.7 eV at the ambient condition to 1.6 eV at 10 GPa. Owing to a positive correlation between the band gap and laser induced damage threshold (LIDT), it indicates that the LIDT value of LiGaTe2 will decrease with applying the external pressure.

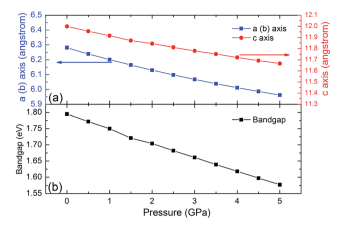


Fig. 13 The crystal lattice constants (a and c) and band gap energy of LiGaTe₂ as a function of pressure.

5. Conclusions

In this study, the big-sized high-quality LiGaTe2 crystal was grown by the Bridgman-Stockbarger method. The anisotropic thermal expansion behavior of the LiGaTe₂ crystal was measured for the first time. Combining the experimental characterization and theoretical calculations, the negative thermal expansion along the c axis was assigned to the rotational vibration mode of (GaTe₄) tetrahedra. The negative thermal expansion was first found in chalcopyrite-type chalcogenides and, respectively, searching for similar structural effects in other materials from this wide crystal family is topical. In addition, the electronic structure of LiGaTe2 crystal was measured by XPS and the recorded valence band is in a good agreement with the theoretical electronic density of states. Moreover, the optical properties and pressure response from 0 to 5 GPa were also calculated in the theory. These results indicate that LiGaTe2, besides its well known pronounced linear and nonlinear optical properties in IR spectral range, possesses the specific structural effects that may be a way to new applications of this telluride material.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the state assignment project (0330-2016-0008), Russian Foundation for Basic Research (Grant No. 17-45-540775, 16-52-48010, 17-52-53031), National Natural Science Foundation of China under Grant No. 91622118, 91622124, 11174297, 51602318, 11611530680 and China "863" project (Grant No. 2015AA034203). Besides, this study was partly supported by the Russian Science Foundation (Grant No. 17-12-01047, in the part of XPS measurements).

References

1 A. Rogalski, Rep. Prog. Phys., 2005, 68, 2267-2336.

- 2 J.-F. Li, W.-S. Liu, Li-D. Zhao and M. Zhou, *NPG Asia Mater.*, 2010, 2, 152–158.
- 3 L. Isaenko, A. Yelisseyev, S. Lobanov, A. Titov, V. Petrov, J.-J. Zondy, P. Krinitsin, A. Merkulov, V. Vedenyapin and J. Smirnova, *Cryst. Res. Technol.*, 2003, 38, 379–387.
- 4 W. Yin, W. Wang, L. Bai, K. Feng, Y. o Shi, W. Hao, J. Yao and Y. Wu, *Inorg. Chem.*, 2012, **51**, 11736–11744.
- 5 V. V. Atuchin, T. A. Gavrilova, K. A. Kokh, N. V. Kuratieva, N. V. Pervukhina and N. V. Surovtsev, *Solid State Commun.*, 2012, **152**, 1119–1122.
- 6 Y. Tanaka, Z. Ren, T. Sato, K. Nakayama, S. Souma, T. Takahashi, K. Segawa and Y. Ando, *Nat. Phys.*, 2012, **8**, 800–803.
- 7 C. D. Malliakas, D. Young Chung, H. Claus and M. G. Kanatzidis, *J. Am. Chem. Soc.*, 2013, 135, 14540–14543.
- 8 A. Yasuda, K. Suto and J.-i. Nashizawa, *Mater. Sci. Semicond. Process.*, 2014, 27, 159–162.
- 9 K. A. Kokh, V. V. Atuchin, T. A. Gavrilova, N. V. Kuratieva, N. V. Pervukhina and N. V. Surovtsev, *Solid State Commun.*, 2014, 177, 16–19.
- 10 S. A. Avanesov, D. V. Badikov, V. V. Badikov, V. L. Panyutin, V. Petrov, G. S. Shevyrdyaeva, A. A. Martynov and K. V. Mitin, J. Alloys Compd., 2014, 612, 386–391.
- 11 S. Welzmiller, F. Hennersdorf, R. Schlegel, A. Fitch, G. Wagner and O. Oekler, *Inorg. Chem.*, 2015, 54, 5745–5756.
- 12 V. V. Atuchin, V. A. Golyashov, K. A. Kokh, I. V. Korolkov, A. S. Kozhukhov, V. N. Kruchinin, I. D. Loshkarev, L. D. Pokrovsky, I. P. Prosvirin, K. N. Romanyuk and O. E. Tereshchenko, J. Solid State Chem., 2016, 236, 203–208.
- 13 L. I. Isaenko and A. P. Yelisseyev, *Semicond. Sci. Technol.*, 2016, **31**, 123001.
- 14 M. C. Ohmer, J. T. Goldstein, D. E. Zelton, A. W. Saxler, S. M. Hegde, J. D. Wolf, P. G. Schuneman and T. M. Pollak, J. Appl. Phys., 1999, 86, 94–99.
- 15 A. H. Reshak, Linear, nonlinear optical properties and birefringence of AgGaX₂ (X = S, Se, Te) compounds, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2005, **369**, 243–253.
- 16 L. Isaenko, P. Krinitsin, V. Vedenyapin, A. Yelisseyev, A. Merkulov, J.-J. Zondy and V. Petrov, *Cryst. Growth Des.*, 2005, 5, 1325–1329.
- 17 J.-J. Zondy, F. Bielsa, A. Douillet, L. Helico, O. Acef, V. Petrov, A. Yelisseyev, L. Isaenko and P. Krinitsin, *Opt. Lett.*, 2007, 32, 1722–1724.
- 18 S. Avanesov, V. Badikov, A. Tyazhev, D. Badikov, V. Panyutin, G. Marchev, G. Shevyrdyaeva, K. Mitin, F. Noack, P. Vinogradova, N. Schebetova, V. Petrov and A. Kwasniewski, Opt. Mater. Express, 2011, 1, 1286–1291.
- S. Wang, G. Liu, Q. Shi, X. Zhang, X. Zhang, C. Li, Z. Gao,
 C. Dong, Q. Lu and X. Tao, RSC Adv., 2014, 4, 27830–27836.
- 20 A. Yelisseyev, P. Krinitsin, L. Isaenko and S. Grazhdannikov, *Opt. Mater.*, 2015, **42**, 276–280.
- 21 A. A. Ionin, I. O. Kinyaevskiy, Y. M. Klimachev, A. A. Kotkov, V. V. Badikov and K. V. Mitin, *Opt. Lett.*, 2016, 41, 2390–2393.
- 22 H. Lin, Y. Liu, L.-J. Zhou, H.-J. Zhao and L. Chen, *Inorg. Chem.*, 2016, **55**, 4470–4475.

- 23 V. Petrov, L. Isaenko, A. Yelisseyev, P. Krinitsin, V. Vedenyapin, A. Merkulov and J.-J. Zondy, J. Non-Cryst. Solids, 2006, 352, 2434–2438.
- 24 F. Liang, L. Kang, Z. Lin, Y. Wu and C. Chen, *Coord. Chem. Rev.*, 2017, 333, 57–70.
- 25 T. C. Ozawa and S. J. Kang, J. Appl. Crystallogr., 2014, 37, 679.
- 26 A. C. Stowe, J. S. Morrell, P. Bhattacharya, E. Tupitsyn and A. Burger, *Proc. SPIE*, 2011, **8142**, 81421H.
- 27 E. Tupitsyn, P. Bhattacharya, E. Rowe, L. Matei, Y. Cui, V. Buliga, M. Groza, B. Wiggins, A. Burger and A. Stowe, *J. Cryst. Growth*, 2014, 393, 23–27.
- 28 L. Isaenko, I. Vasilieva, A. Merkulov, A. Yelisseyev and S. Lobanov, *J. Cryst. Growth*, 2005, 275, 217–223.
- 29 L. I. Isaenko and I. G. Vasilyeva, J. Cryst. Growth, 2008, 310, 1954–1960.
- 30 I. G. Vasilyeva, R. E. Nikolaev, P. G. Krinitsin and L. I. Isaenko, J. Phys. Chem. C, 2017, 121, 17429–17435.
- 31 Yu. M. Basalaev, Yu. N. Zhuravlev, E. B. Kitova and A. S. Poplavnoi, *J. Struct. Chem.*, 2007, **48**, 1001–1005.
- 32 V. V. Atuchin, L. I. Isaenko, V. G. Kesler, S. Lobanov, H. Huang and Z. S. Lin, *Solid State Commun.*, 2009, **149**, 572–575.
- 33 A. V. Kosobutsky, Y. M. Basalaev and A. S. . Poplavnoi, *Phys. Status Solidi B*, 2009, **246**, 364–371.
- 34 A. V. Kosobutsky and Y. M. Basalaev, *J. Phys. Chem. Solids*, 2010, **71**, 854–861.
- 35 T. Ma, L. Sun, C. Xu and Y. Chen, *J. Alloys Compd.*, 2011, **509**, 9733–9741.
- 36 M. Tian-Hui, Z. Zhi-Ping and R. n Yu-Lan, *Acta Phys. Sin.*, 2012, **61**, 197101.
- 37 S. A. Grazhdannikov, P. G. Krinitsyn, A. F. Kurus', L. I. Isaenko, A. P. Yelisseyev and M. S. Molokeev, *Mater. Sci. Semicond. Process.*, 2017, 72, 52–59.
- 38 D. V. Demidov, I. P. Prosvirin, A. M. Sorokin and V. I. Bukhtiyarov, *Catal. Sci. Technol.*, 2011, 1, 1432–1439.
- 39 V. V. Atuchin, V. A. Golyashov, K. A. Kokh, I. V. Korolkov, A. S. Kozhukhov, V. N. Kruchinin, S. V. Makarenko, L. D. Pokrovsky, I. P. Prosvirin, K. N. Romanyuk and O. E. Tereshchenko, *Cryst. Growth Des.*, 2011, 11, 5507–5514.
- E. J. Rubio, V. V. Atuchin, V. N. Kruchinin, L. D. Pokrovsky,
 I. P. Prosvirin and C. V. Ramana, *J. Phys. Chem. C*, 2014,
 118, 13644–13651.
- 41 J. H. Scofield, J. Electron Spectrosc. Relat. Phenom., 1976, 8, 129–137.
- 42 S. J. Clark, M. D. Segall, C. J. Pickard, P. J. Hasnip, M. J. Probert, K. Refson and M. C. Payne, *Z. Kristallogr.*, 2005, 220, 567–570.
- 43 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865–3868.
- 44 A. M. Rappe, K. M. Rabe, E. Kaxiras and J. D. Joannopoulos, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1990, **41**, 1227–1230.
- 45 H. J. Monkhorst and J. D. Pack, *Phys. Rev. B: Solid State*, 1976, 13, 5188–5192.
- 46 S. Baroni, S. de Gironcoli, A. Dal Corso and P. Giannozzi, *Rev. Mod. Phys.*, 2001, 73, 515–562.
- 47 L. A. Solovyov, J. Appl. Crystallogr., 2004, 37, 743-749.

Paper

- 48 J. Haines, C. Chateau, J. M. Léger, C. Bogicevic, S. Hull, D. D. Klug and J. S. Tse, Phys. Rev. Lett., 2003, 91, 015503.
- 49 V. V. Atuchin, L. I. Isaenko, V. G. Kesler and S. I. Lobanov, J. Alloys Compd., 2010, 497, 244-248.
- 50 K. S. Mandal, R. M. Krishna, T. C. Hayes, P. G. Muzykov, S. Das, T. S. Sudarshan and S. Ma, IEEE Trans. Nucl. Sci., 2011, 58, 1981-1986.
- 51 L. V. Yashina, E. V. Tikhonov, V. S. Neudachina, T. S. Zyubina, A. N. Chaika, V. I. Shtanov, S. P. Kobeleva and Yu. A. Dobrovolsky, Surf. Interface Anal., 2004, 36, 993-996.
- 52 V. V. Atuchin, V. A. Golyashov, K. A. Kokh, I. V. Korolkov, A. S. Kozhukhov, V. N. Kruchinin, I. D. Loshkarev,

- L. D. Pokrovsky, I. P. Prosvirin, K. N. Romanyuk and O. E. Tereshenko, J. Solid State Chem., 2016, 236, 203-208.
- 53 J. P. Perdew and Y. Wang, Phys. Rev. B: Condens. Matter Mater. Phys., 1992, 45, 13244-13249.
- 54 C. S. Wang and B. M. Klein, Phys. Rev. B: Condens. Matter Mater. Phys., 1981, 24, 3417-3429.
- 55 L. Bai, Z. S. Lin, Z. Z. Wang and C. T. Chen, J. Appl. Phys., 2008, 103, 083111.
- 56 R. D. Eithiraj, G. Jaiganesh, G. Kalpana and M. Rajagopalan, Phys. Status Solidi B, 2007, 244, 1337-1346.
- 57 R. Zallen, High Pressure Res., 2004, 24, 117-118.