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LiVTeO₅: a mid-infrared nonlinear optical vanadium tellurate crystal exhibiting enhanced second harmonic generation activities and notable birefringence†

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Mid-infrared (IR) nonlinear optical (NLO) crystals are of great interest for applications in laser science and technology. Nevertheless, the pursuit of mid-IR NLO crystals displaying exceptional comprehensive performance remains a challenge. Herein, an effective strategy is achieved by combining two distinct species of second-order Jahn–Teller distorted cations, Te^{4+} and V^{5+} , into one compound. A prospective NLO vanadium tellurate crystal, LiVTeO₅, was synthesized *via* spontaneous crystallization, which crystallizes in the orthorhombic space group $P2_12_12_1$ (no. 19) with a three-dimensional framework structure. The optical measurements revealed that LiVTeO₅ manifests a large band gap of 3.2 eV and an intense second-harmonic-generation (SHG) activity of 3.2 × KDP. Moreover, it displays an extensive transmittance region beyond 5.0 μ m, along with a discernible birefringence of 0.091, thereby facilitating the realization of the phase matching conditions. Additionally, structural analysis combined with first-principles calculations elucidate that the [TeO₃] and [VO₅] active building blocks primarily contribute to the intriguing macroscopic nonlinearity and notable birefringence observed in LiVTeO₅. These findings highlight the optimized overall properties of LiVTeO₅, which makes it a potential mid-IR NLO material.

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Introduction

Nonlinear optical (NLO) materials capable of generating coherent and tunable lasers through frequency conversion capability, particularly those with the mid-infrared (IR) region covering the crucial atmospheric window (3–5 μm), have garnered significant interest in science and technology. $^{1-5}$ In recent years, significant strides have been made in the realm of design, synthesis, property characterization, and structure-property elucidation of mid-IR NLO compounds. Prominent examples include commercially available chalcogenide- and pnictide-based crystals, $AgGaQ_2$ (Q = Se, S) and $ZnGeP_2.^{6-8}$ Nevertheless, the practical applications of these materials are hampered by inherent limitations, such as poor laser damage

To manifest a second-order NLO effect, the paramount precondition is that the compound must crystallize in a noncentrosymmetric (NCS) structural form. Previous findings have shown that incorporating transition metal-based cations, susceptible to the second-order Jahn–Teller (SOJT) effect, into the structure constitutes an effective strategy for inducing a desirable NCS configuration. $^{17-20}$ Among the SHG-active d^0 cations, the average distortive magnitude roughly adheres to the following hierarchy: $Mo^{6+} \approx V^{5+} > W^{6+} > Ti^{4+} > Nb^{5+} > Ta^{5+} > Zr^{4+}$. This suggests that the introduction of V^{5+} cations can enhance the probability of achieving an amplified SHG response. 19 Furthermore, V atoms have versatile coordination modes, encompassing $[VO_4]$, $[VO_5]$, and $[VO_6]$ polyhedral arrange-

thresholds for $AgGaQ_2$ and pronounced two-photon absorption at the conventional wavelength of 1 μm for $ZnGeP_2$. In contrast to chalcogenides and pnictides, oxides have emerged as extensively employed NLO materials in commercial applications, exemplified by KH_2PO_4 (KDP), β -BaB $_2O_4$ (β -BBO), LiB $_3O_5$ (LBO), KTiOPO $_4$ (KTP) and LiNbO $_3$ (LN), $^{10-15}$ attributed to their large band gaps and high laser damage thresholds. Regrettably, most oxide crystals are incapable of supporting the fabrication of NLO devices that operate beyond 5.0 μm . Consequently, there exists an urgent imperative to target novel IR NLO oxide crystals that offer favorable overall performance.

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ments, thereby allowing for diverse structural configurations. ²¹ Apart from the d⁰ cations, the introduction of stereo-chemically active lone pair (SCALP) cations, exemplified by heavy element ions like Te4+, has garnered considerable interest owing to their capability to construct SHG-active chromophores and elicit a red-shift in the IR absorption cutoff. For instance, the NLO tellurate crystal Li₂ZrTeO₆, which replaces Nb in LiNbO3 with Te and Zr, extends the IR transmission cutoff edge to 7.4 μm . $^{22-26}$

Drawing upon these ideas, we propose that by incorporating both V⁵⁺ and Te⁴⁺ ions, both of which are susceptible to the SOIT effect, into one compound, a pronounced structural distortion and a wide infrared (IR) transmission range can be effectively maintained.²⁷ Consequently, the vanadium tellurate crystal LiVTeO5 (LVTO), which possesses an acentric orthorhombic structure analyzed through powder X-ray diffraction in 1973,28 has piqued our interest as it aligns with our analysis strategy. However, investigations regarding the NLO properties of LVTO have remained absent from the scientific discourse. Accordingly, we successfully fabricated a high-quality vanadium tellurate crystal, LVTO, through a spontaneous crystallization technique. By optimizing the structure constructed from the NLO-active functional units [VO₅] and [TeO₃], LVTO achieves a well-balanced optical performance. In this study, we report the preparation, crystal growth, structural analysis, thermal behavior, and optical characterization of LVTO by the combination of experimental and theoretical approaches.

Experimental

Polycrystalline synthesis

The polycrystalline LVTO was obtained utilizing a stoichiometric one-step solid-phase reaction technique. The starting materials including Li₂CO₃ (Alfa Aesar, 99.9%), V₂O₅ (Alfa Aesar, 99.9%) and TeO₂ (Aladdin, 99.9%) according to their stoichiometric ratio were manually ground in an agate mortar and preheated at 350 °C for 10 h in a muffle furnace, followed by calcination at 480 °C for 72 h with several intermediate regrinding rounds. Powder X-ray diffraction (PXRD) analysis was employed to examine the phase purity of the target product.

Powder X-ray diffraction (PXRD)

A Rigaku SmartLab 9 kW diffractometer equipped with a diffraction monochromator setting for Cu K_{α} radiation (λ = 1.5418 Å) was used to record the PXRD data of the target compound. The test was performed over the range $10^{\circ} \le 2\theta \le 70^{\circ}$ at a scan step width of 0.01° with a fixed counting time of 0.3 s.

Thermal property analysis

The thermal behavior was analyzed for the LVTO crystalline samples with a NETZSCH STA 449F5 TG/DTA thermal analyzer. LVTO polycrystalline specimens, weighing approximately 12.5 mg, were carefully introduced into an Al₂O₃ crucible. The

crucible was subjected to a controlled heating process, gradually raising the temperature from ambient conditions to 600 °C at a heating rate of 10 °C per minute under a flowing nitrogen gas atmosphere.

Energy-dispersive spectroscopy (EDS) analysis

To probe the elemental ratios in the LVTO crystal, elemental distribution maps and microprobe elemental analyses were recorded using a field-emission scanning electron microscope (FEI Quanta FEG 250).

Structural characterization

Single crystal data were collected through single crystal X-ray diffraction experiments on a Bruker D8 VENTURE CMOS X-ray source at a constant temperature with Mo K α radiation (λ = 0.71073 Å). Cell refinement, data integration, and absorption correction were implemented using the APEX III software. The preliminary crystal structure data were resolved by direct methods using the SHELXTL crystallographic software package. Comprehensive structural parameter data are presented and elucidated in Tables S1-S3 (ESI†), which involve the atomic coordinates, equivalent isotropic displacement parameters, interatomic distances and angles, and the bond valence sums (BVS) for all atoms. The corresponding experimental conditions and the refined crystallographic data are available in Table 1.

UV-visible diffuse reflectance spectra

The ultraviolet-visible (UV-vis) diffuse reflectance spectra were gauged from 200 to 2000 nm utilizing a Hitachi UH4150 spectrophotometer at ambient temperature, while BaSO₄ was selected as the diffuse reflectance standard sample. The Kubelka-Munk remission function was employed to transform reflectance spectra into absorbance: $F(R) = (1 - R)^2/(2R) = K/S$, where R is the reflectance, K is the absorption and S is the scattering.29

Table 1 Crystallographic data and structural refinement of LVTO

Empirical formula	LiVTeO ₅
Formula weight	265.48
Temperature (K)	298(2)
Wavelength (Å)	0.71073
Crystal system	Orthorhombic
Space group	$P2_12_12_1$
a (Å)	5.2789(2)
b (Å)	8.3415(2)
$c(\mathring{A})$	9.5082(3)
Volume (ų)	418.68(2)
Z	4
Density (g cm ⁻³)	4.212
F(000)	472
R(int)	0.0451
Completeness	100%
GOF on (F^2)	1.079
Final R indices $[F_0^2 > 2\sigma(F_0^2)]^a$	$R_1 = 0.0133$, $wR_2 = 0.0341$
R indices (all data)	$R_1 = 0.0139, \text{w} R_2 = 0.0344$
CCDC number	2289244

 $^{a}R_{1} = \sum ||F_{o}| - |F_{c}||/\sum |F_{o}|; wR_{2} = [\sum w(F_{o}^{2} - F_{c}^{2})^{2}/\sum w(F_{o}^{2})^{2}]^{1/2}.$

IR spectroscopy and Raman spectra

The IR spectra of the LVTO powder samples were characterized using a Nicolet iS50 FT-IR infrared spectrometer at ambient temperature across a spectral range of 400-4000 cm⁻¹. About 3.5 mg samples were placed on the test platform for testing. Raman spectroscopy measurements were recorded on a WITECalpha300R spectrometer equipped with a CCD detector using 532 nm excitation radiation at ambient temperature. The measurements were performed in the range of 100 to 1100 cm⁻¹.

Second-harmonic-generation tests

Second harmonic generation (SHG) tests were performed on the polycrystalline samples of LVTO using a Q-switched Nd: YAG laser with a fundamental wavelength of 1064 nm, following the Kurtz-Perry method. A powdered microcrystalline sample of LVTO was thoroughly ground and then sieved into several distinct particle size ranges of 50-75, 75-109, 109-120, 120-150, 150-180, and 180-212 µm. For comparison of SHG activities, KH₂PO₄ (KDP) with the same particle size ranges were also prepared as the measured reference patterns.

Birefringence characterization

The optical birefringence of the target compound was experimentally examined through a polarizing microscope (Nikon Eclipse E200MV POL) with a visible light source. The corresponding birefringence value can be collected following the equation: $R = \Delta n \times D = |N_e - N_o| \times D$, where Δn denotes the birefringence, D refers to the crystal thickness, R signifies the retardation, and N_e and N_o represent the birefringence of the extraordinary wave and the ordinary wave, respectively.30 A pristine crystal was chosen and its thickness was ascertained employing a Bruker SMART APEX III CCD diffractometer.

Computational details

We performed first-principles calculations using density functional theory (DFT) calculations within the Cambridge sequential total energy package (CASTEP) software to analyze the electronic density difference and band structure of the LVTO crystal.31 In order to accurately comprehend the microcosmic mechanism of the LVTO crystal, the exchange-correlation effects were described using the Perdew-Burke-Ernzerhof (PBE) functional with the generalized gradient approximation (GGA) scheme. 32-34 Norm-conserving pseudopotentials were utilized with the following valence electron configuration: Li 2s¹, Te 5s²5p⁴, V 3d³4s², and O 2s²2p⁴. To ensure reliable simulation results, the plane-wave cutoff energy was set to 810 eV, and a Monkhorst-Pack scheme k-point mesh of $6 \times 4 \times 4$ in the first Brillouin zone was used for sampling and numerical integration.

Results and discussion

Powder X-ray diffraction (PXRD) and thermal analysis

Polycrystalline samples of LVTO were synthesized employing solid-state methodologies. The phase purity of LVTO was authenticated through PXRD Rietveld refinement, utilizing the Toolbar FullProf Suite program. As shown in Fig. 1a, the PXRD patterns of the as-synthesized samples match well with the calculated observations derived from the structural data. Additionally, the thermal durability of LVTO was assessed via precise measurements of DSC and TGA. Fig. 1b demonstrates that the DSC curves exhibit one appreciable endothermic peak at approximately 498 °C upon heating, while the TG curves demonstrate no weight loss over the test temperature range. These results suggest that LVTO possesses good thermal stability without undergoing any phase transition. Moreover, the powder PXRD patterns of the melted solid agree well with

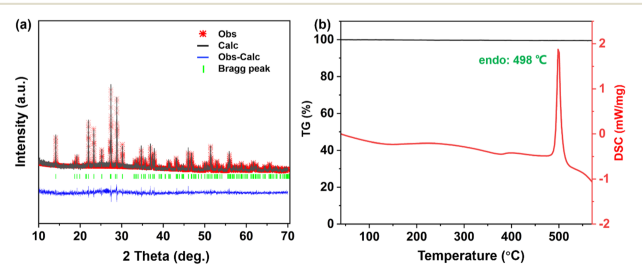


Fig. 1 (a) Rietveld refinement patterns: calculated data (black line), experimental data (red asterisk), and the difference (blue trace) between them. The tick marks (green) display the expected Bragg reflections. (b) DSC and TG curves for LVTO.

those of the initial LVTO, revealing that LVTO is a congruent compound (Fig. S1†).

Crystal growth and the structure of LVTO

Research Article

LVTO crystals were prepared using an optimized spontaneous nucleation technique. After conducting several rounds of experiments, we successfully cultivated bulk single crystals of LVTO with dimensions of $2 \times 2 \times 1.5 \text{ mm}^3$, as exemplified in Fig. S2.† The structural information of LVTO was first determined through powder X-ray diffraction in 1973.²⁸ To delve deeper into the intricate interplay between the structure and properties, single-crystal XRD was then employed to reconfirm the structure of LVTO. The title compound crystallized in an acentric orthorhombic configuration with the space group $P2_12_12_1$ (no. 19). The refined unit cell parameters were discerned to be a = 5.2789(2) Å, b = 8.3415(2) Å, c = 9.5082(3) Å, and $\beta = 90^{\circ}$. One independent Li atom, one independent Te atom, one independent V atom, and five O atoms were observed in the asymmetric unit of LVTO. As shown in Fig. 2a, the V atom is surrounded by five O atoms, forming a distorted trigonal bipyramidal geometry [VO₅] with V-O lengths ranging from 1.642(3) to 2.055(3) Å. The Te⁴⁺ cation adopts a triangular out-of-plane coordination with three O atoms, and the lengths of the three Te-O bonds are in the range of 1.852(3) to 1.912(3) Å due to the presence of Te⁴⁺ lone pair electrons. Each [TeO₃] triangular pyramid intertwines with three neighboring [VO₅]

square pyramids via shared oxygen atoms, orchestrating the construction of $[\text{TeV}_3\text{O}_{15}]^{11-}$ zigzag grids (Fig. 2b). These $[\text{TeV}_3\text{O}_{15}]^{11-}$ zigzag grids are further interconnected to form a three-dimensional (3D) framework structure, in which all Li⁺ cations are bonded in a 5-coordinate geometry (Fig. S3†) and are located in the cavities to balance the charge (Fig. 2c). Notably, this remarkable 3D framework unveils a quintessential honeycomb-like configuration, with tunnels that stretch indefinitely along the a axis as presented in Fig. 2d.

In terms of the formula, LiVTeO₅ and LiNbTeO₅ can be considered as LiATeO₅ (A = V, Nb).²⁴ However, by substituting V⁵⁺ cations in LTVO with Nb5+ cations of the same main group in the NLO crystal LiNbTeO5, it crystallizes in the P21 space group, which is non-isotypic to LVTO. To further elucidate this situation, a comparative analysis of their structural features was conducted. In LVTO, V^{5+} cations are all five-coordinated and adopt a distorted trigonal bipyramidal geometry, while the Nb5+ cations in LiNbTeO5 are located in a distorted octahedral environment. Additionally, one [TeO₃] triangular pyramid and three adjacent [NbO₆] octahedra in LiNbTeO₅ are linked by corner sharing to form $[\text{TeNb}_3\text{O}_{17}]^{15-}$ and $[\text{TeNb}_3\text{O}_{18}]^{17-}$ zigzag grids, respectively (Fig. 2e). As shown in Fig. 2f and g, the 3D structure of LiNbTeO5 also displays characteristics of honeycomb-like channels similar to those in LVTO. Furthermore, unlike the independent $[VO_5]$ polyhedra in LVTO, the $[NbO_6]$ octahedra are interconnected to form [Nb₂O₁₀]_∞ zigzag chains

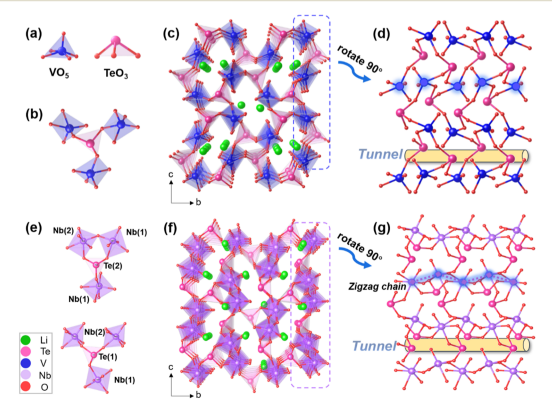


Fig. 2 Structural characterization. (a) $[VO_5]$ and $[TeO_3]$ motifs. (b) The $[TeV_3O_{15}]^{11-}$ fundamental building block of LVTO. (c) 3D honeycomb-like framework of LVTO. (d) The crystal structure of LVTO viewed along the *ac* plane. (e) The $[TeNb_3O_{17}]^{15-}$ and $[TeNb_3O_{18}]^{17-}$ building blocks of LiNbTeO₅. (f) 3D honeycomb-like framework of LiNbTeO₅. (g) The crystal structure of LiNbTeO₅ viewed along the *ac* plane.

in LiNbTeO₅. It is suspected that the disparate evolutionary trajectories of these structures give rise to variations in their symmetrical attributes.

In addition, bond valence sum (BVS) calculations reveal the expected oxidation states of 3.80, 5.09 and 2.03 for Te, V and O, respectively, which are consistent with those derived from the structural data of LVTO. Energy-dispersive spectroscopy (EDS) measurements corroborated the existence of V/Te/O with an average atomic ratio in a reasonable range for LVTO (Fig. S4†), which aligns well with the observations from the structural analysis.

Optical performance

The UV-vis diffuse reflectance spectrum of the polycrystalline LVTO revealed that the UV cut-off edge occurs at approximately 357 nm, corresponding to an experimental band gap of 3.2 eV (Fig. 3a), as determined utilizing the Kubelka–Munk formula equation. Moreover, the IR spectroscopy of LVTO is presented in Fig. 3b, demonstrating its high mid-IR transparency up to

914 cm $^{-1}$, which covers a critical atmospheric transparent window (3–5 µm). Additionally, the absorption peak characteristics associated with V–O and Te–O vibrations in the IR spectrum were analyzed. The bands located at 796, 865, and 914 cm $^{-1}$ predominantly originate from the V–O vibration. The bands at around 600–740 cm $^{-1}$ are mainly attributed to the Te–O vibration. These peak assignments are consistent with those reported in previous studies.

Furthermore, Raman spectrum tests were recorded to comprehensively identify vibration modes in structural analysis. As shown in Fig. S5,† the absorption bands located at 365, 645, and 737 cm⁻¹ can be assigned to the characteristic symmetric and asymmetric stretching vibrations, as well as the bending vibration of [TeO₃], according to previous findings. The bands at around 750 cm⁻¹ are attributed to the symmetric stretching vibration of [TeO₃] units. The Raman frequency-shifted peaks observed in the range of 790–920 cm⁻¹ are mainly induced by the V–O stretching vibration. The bands ranging from 400 to 495 cm⁻¹ are dominated by the Li–O vibrations. These above observations confirm

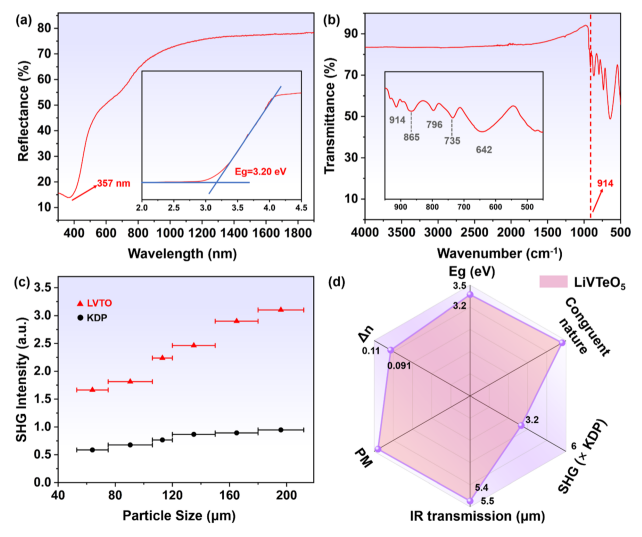


Fig. 3 Optical characterization of LVTO. (a) UV-vis diffuse-reflectance spectrum and the band gap. (b) IR transmission spectra. (c) Phase-matching curves for LVTO and KDP, respectively. (d) The overall NLO performance of LVTO.

the presence of [TeO $_3$] and [VO $_5$] groups, which corresponds well to the single crystal structure of LVTO. 23,35,37

SHG measurement

As the LVTO compound crystallizes in the $P2_12_12_1$ space group and lacks an inversion center within the crystal structure, the Kurtz-Perry method was employed to evaluate its powder SHG performance under a Q-switched Nd:YAG laser (1064 nm).³⁸ As depicted in Fig. 3c, the SHG signals of LVTO exhibit enhancement with increasing particle size, and eventually reaching saturation, which implies that LVTO follows the phase-matchable rule. Within the particle size range of 180-212 μm, it is noteworthy that the LVTO compound exhibits a pronounced SHG intensity of 3.2 times that of KDP. Moreover, the desirable SHG activities of LVTO are comparable to those of other NLO tellurates, such as $\text{Li}_2\text{ZrTe}_2\text{O}_6$ (2.5 × KDP) and $\text{Li}_2\text{SnTeO}_6$ (2.5 × KDP).^{26,39} Hence, it is evident that the title compound exhibits commendable attributes (Fig. 3d), including a large band gap, a broad transmission window, and favorable SHG effects, making it a promising mid-IR NLO crystal.

Birefringence characterization

In order to achieve coherent light output, birefringence is a critical parameter for satisfying phase-matching conditions. The experimental birefringence of LVTO was quantified employing a cross-polarizing prism methodology. The precise thickness of the crystal plane was determined to be 11.7 μ m utilizing a Bruker single crystal diffractometer, as depicted in Fig. 4a. When incident light passes through an anisotropic single crystal, it undergoes a division into two polarized beams. The refractive indices of the two polarized lights are not equal and the propagation speeds are distinct (N_0 and N_e). The original interference color and the complete extinction of LVTO under orthogonally polarized light were confirmed to be second-order orange according to the Michel-Levy color chart in the inset of

Fig. 4b, and it achieves complete extinction through the employment of a Berek compensator. In accordance with the equation $R = \Delta n \times D$, the experimental measurement yielded a noteworthy value of 0.091 at the visible wavelength, suggesting that the birefringence is sufficiently substantial to facilitate phase matching. Moreover, the wavelength-dependent refractive indices for LVTO were also calculated, and the theoretical value of the birefringence was found to be 0.090 at 1064 nm as presented in Fig. 4b, in parallel with the experimental one. This value is also comparable to that of the commercial NLO and birefringent crystal LiNbO₃ (0.0786). 40,41

Structure-property relationship

To elucidate the underlying relationship between the structural characteristics and optical properties, theoretical calculation was carried out for LVTO. 42 As shown in Fig. 5a, the band structures indicate that LVTO belongs to an indirect band gap compound with a theoretical band gap value of 2.60 eV, which is consistent with the experimental observation. Fig. 5b illustrates the detailed total and partial density of states (PDOS), revealing that the near-forbidden-gap states are mainly contributed by Te, V and O atoms, while the contribution of the Li atom is relatively limited. Specifically, the electron states near the bottom of the conduction bands (CB) ranging from 2.0 to 7.0 eV are predominantly dominated by the 5p orbitals of tellurium and the 3d orbitals of vanadium, with the 2s orbitals of lithium making a smaller contribution. The behavior of Li orbitals can be ascribed to their coordination in a five-coordinated [LiO5] environment, which induces a structural distortion. The band positions in proximity to the maximum of the valence band (VB) spanning from -5.0 to 0.0 eV predominantly derive from oxygen 2p orbitals, showcasing extensive hybridization with tellurium 5p and vanadium 3d orbitals. The electron states in the range of -7 to -10 eV are mainly composed of the hybridizations of Te 5s and O 2p states.

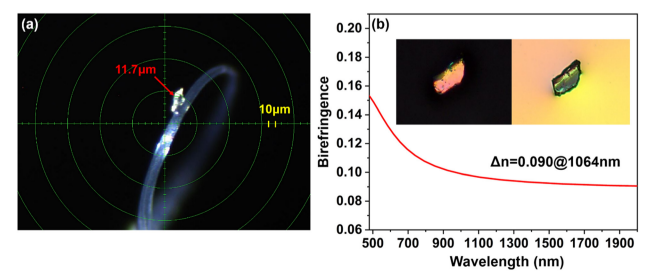


Fig. 4 Birefringence measurements for LVTO. (a) The measured thickness of the LVTO crystal. (b) The theoretical refractive index dispersion curves with the insets representing the original interference color and the complete extinction state of LVTO.

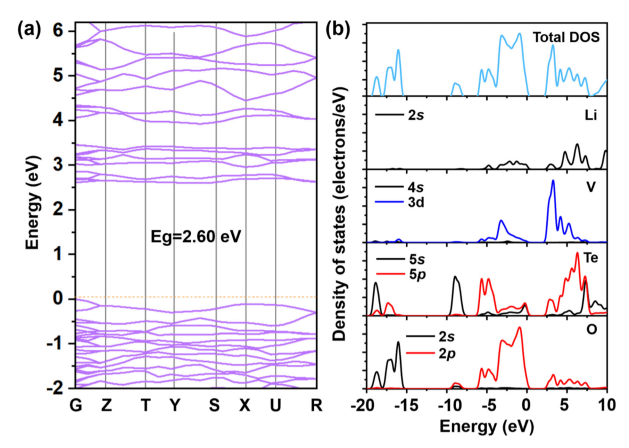


Fig. 5 Calculated electronic structure of LVTO. (a) Energy band structure. (b) Total DOS and partial DOS.

Based on these analyses, it can be perceived that the SOJT distortive building motifs [TeO₃] and [VO₅] are mainly responsible for the appreciable NLO activity of LVTO.

To further quantitatively elucidate the contribution of these groups to NLO performance, the local dipole moments of structural motifs in LVTO were examined utilizing the bondvalence approach methodology. As illustrated in Fig. S6,† the orientations of the dipole moments for these groups within one unit cell exhibit inconsistency, which is distinct from those observed in LiNbTeO₅. The magnitudes of the dipole moments for the individual [TeO₃], [VO₅] and [LiO₅] polyhedra were calculated and found to be 21.98, 4.42 and 2.87 Debye, respectively (Table S4†).

Conclusions

In summary, a prospective NLO vanadium tellurate crystal, LiVTeO₅, was designed by combining SOJT distorted cations Te⁴⁺, with stereochemically-active lone pairs, and V⁵⁺ cations characterized by a d⁰ electronic configuration. Optical characterization demonstrated that the aforementioned compound achieves a favorable balance of crucial optical properties, including an amplified SHG response of 3.2 × KDP, a large band gap of (3.2 eV), an extensive transmittance region, and a sufficient birefringence of 0.091 at the visible wavelength, enabling phase matching capability. First-principles calculations implied that the NLO and birefringent performance can be intuitively ascribed to the highly distortive units [TeO₃] and [VO₅]. Our findings unveil the promising potential of LiVTeO5 in NLO applications and pave the way for the exploration of novel mid-IR NLO materials exhibiting desirable performance.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

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