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The spin-orbit-phonon coupling and crystalline elasticity of LaCoO₃ perovskite

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Based on an integrated study of magnetic susceptibility, specific heat, and thermal expansion of single-crystal LaCoO $_3$ free from cobalt and oxygen vacancies, two narrow spin gaps are identified before and after the phonon softening of gap size $\Delta E \sim 0.5$ meV in a CoO $_6$ -octahedral crystal electric field (CEF) and the thermally activated spin gap $Q \sim 25$ meV, respectively. Significant excitation of Co $^{3+}$ spins from a low-spin (LS) to a high-spin (HS) state is confirmed by the thermal activation behavior of spin susceptibility χ^S of energy gap $Q \sim 25$ meV, which follows a two-level Boltzmann distribution to saturate at a level of 50% LS/50% HS statistically above ~ 200 K, without the inclusion of a postulated intermediate spin (IS) state. A threefold increase in the thermal expansion; coefficient (α) across the same temperature range as that of thermally activated HS population growth is identified, which implies the non-trivial spin-orbit-phonon coupling caused the bond length of Co $^{3+(LS \leftrightarrow HS)}$ -O fluctuation and the local lattice distortion. The unusually narrow gap of $\Delta E \sim 0.5$ meV for the CoO $_6$ octahedral CEF between e $_9$ -t $_{29}$ indicates a more isotropic negative charge distribution within the octahedral CEF environment, which is verified by the Electron Energy Loss Spectroscopy (EELS) study to show nontrivial La-O covalency.

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Introduction

Following extensive research into ABO3-type transition metal oxide perovskites in the 1950s, 1,2 among the series LaTO₃ (T = Cr, Mn, Fe, Co, Ni), LaCoO3 was demonstrated to exhibit rich magnetic and structural properties. For example, LaCoO₃ does not possess the expected high-symmetry cubic structure found in most perovskites, but has a pseudo-cubic distortion best described with the rhombohedral space group $R\bar{3}c$ (no. 167) up to temperatures of \sim 650 K,³ as illustrated in Fig. 1. In addition, the magnetic susceptibility $\chi(T)$ of LaCoO₃ shows both thermal activation behavior and Curie-Weiss paramagnetic behavior arising presumably from the excitation of nonmagnetic Co³⁺ ions ([Ar]3d⁶) sitting in the CoO₆ octahedral CEF, which has been attributed to a thermally activated two-level spin state transition of Co^{3+} from the low-spin (LS = $\mathrm{t_{2g}^6}\mathrm{e_g^0}$) state with S=0 to the high-spin (HS = $t_{2g}^4 e_g^2$) state with S = 2,4,5 although orbital ordering for the intermediate-spin (IS = $t_{2g}^5 e_g^1$) state with

In their pursuit of a deeper understanding of the "transitional" nature of d-orbital electrons-from a localized to an itinerant character-in the transition metal perovskites, Goodenough and co-workers¹⁶⁻¹⁹ have raised many fundamentally important questions closely related to LaCoO₃, such as (1) why is it difficult to obtain phases having more than 50% of Co⁴⁺ through A-site substitution? (2) What is the origin of the localized spins showing "Curie tail" behavior below ~30 K? (3) Does disproportionation of Co³⁺ into Co²⁺–Co⁴⁺ occur? (4) Is there an ordered configuration of LS and HS? In addition, does the IS state really exist in LaCoO₃? To answer these questions conclusively, mixed valence of cobalt due to nonstoichiometry must first be ruled out, especially the oxygen vacancy.

By focusing on reducing the localized impurity spin contribution in the growth of single-crystal LaCoO₃, we have obtained vacancy-free single crystals using the optical traveling solvent floating-zone method following a high-oxygen-pressure and high-temperature post-annealing procedure. In particular, the vacancy-free sample shows no low-temperature $\chi(T)$ Curie tail, but saturates to a constant as $T \to 0$ K, which is in sharp contrast to the behavior resulting from cobalt and/or oxygen vacancy defects that has been reported in nearly all early publications.^{7,20,21} Integrated data analysis of the magnetic susceptibility $\chi(T)$, specific heat $C_P(T)$, and thermal expansion coefficient $\alpha(T)$ without complications arising from impurity

S=1 has also been proposed.^{6,7} Many conflicting reports about the spin state of LaCoO₃ have been made based on the DFT calculations, susceptibility model fittings, and spectroscopic studies since.^{4,8-15}

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(a) (b)

Fig. 1 (a) The expected cubic symmetry of common ABO₃-type perovskite structure. (b) The room temperature structure of LaCoO₃ is pseudocubic best described with a rhombohedral symmetry of space group R3c (Co in blue, O in red, and La in green).

spins and mixed valence leads to the conclusive identification of an HS population distribution across the two energy gaps of $\Delta E \sim 0.5$ meV and $Q \sim 25$ meV before and after phonon softening.

Current findings on defect-free $LaCoO_3$ are able to resolve most of the controversies raised in the literature, including the debate about the existence of an IS state and orbital ordering. In addition, the finding of three times increase of thermal expansion coefficient above the thermally activated phonon softening range strongly suggests that the nontrivial spin–orbit–phonon coupling is responsible for the pseudo-cubic symmetry of the ABO_3 -type perovskite due to constant Co–O bond length fluctuation at room temperature.

II. Experimental details

Single crystal LaCoO3 was grown using optical floating-zone growth method through direct melting and re-crystallization for its congruent melt nature with a melting point of 1750 °C. In the preparation of polycrystalline LaCoO₃ feed rod, Co and O loss is unavoidable during high temperature treatment in the air at ambient pressure, which is evidenced by the commonly found Al₂CoO₄ precipitate in blue color at the bottom of alumina annealing tray (shown in Fig. 2). It is found that a 5% Co excess in the precursor is required to compensate for the loss of Co for treatment above 990 °C. For the high pressure oxygen annealing process, the as-grown single crystal was mixed with Co₃O₄ powder in molar ratio of 1:1 sealed in a thick wall 310S stainless steel tubing with added KClO₄ oxidant. The sealed tubing was annealed at 1200 °C for one week. The quantitative chemical analysis has been applied to the as-prepared and annealed crystal samples with electron probe micro-analysis (EPMA) and the oxygen/nitrogen combustion analyzer (EMGA-920, Horiba).

The magnetic susceptibility, specific heat, and thermal expansion coefficient of single crystal LaCoO $_{3-\delta}$ samples were measured using SQUID-VSM (QD-USA) in both AC and DC modes, Physical Property Measurement System (PPMS, QD-

USA) with a pulsed heating method, and a capacitance-type device in the PPMS platform between 2 to 600 K, respectively. In particular, the linear thermal expansion coefficient $\alpha=(1/L_0)$ d $L/\mathrm{d}T$ of single-crystal samples was measured directly with a newly designed capacitance-type device incorporated into the PPMS.

The electron energy-loss spectroscopy (EELS) spectra were acquired using Transmission Electronic Microscope (TEM) with a JEOL 2100F field emission gun operated at 200 kV, which equipped with a Gatan image filter (GIF, Tridiem 863) of energy resolution \sim 0.85 eV. The TEM specimen was prepared with



Fig. 2 After LaCoO $_3$ crystal is annealed at 800 °C in the argon atmosphere, a blue color residue is coated, which indicates the formation of Al $_2$ CoO $_4$.

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focused ion beam (FIB, Tescan GAIA). The single scattering STEM-EELS spectra were deconvoluted from the raw data with subsequent Kramers-Kronig Analysis (KKA), and the calculation of effective electron number (n_{eff}) was conducted with the Digital Micrograph EELS package which was written on the basis of ref. 22.

III. Results and discussions

Crystal nonstoichiometry

The chemical analysis results of the as-prepared and annealed crystal samples are summarized in Table 1. Both the Co and O vacancies have been controlled through the initial 5% Co excess adjustment and the post high-pressure high-temperature annealing process. The magnetic susceptibilities $\chi(T)$ and the isothermal magnetization M(H) at 2 K are compared in Fig. 3 to show the impact of nonstoichiometry on the physical properties. It is noted that samples with either Co or O vacancies shows clear Curie-tail behavior for $\chi(T)$ at low temperature, which indicates the existence of localized spins from defects. Additional field scan of isothermal magnetization M(H) at 2 K [inset of Fig. 3(a)] has provided solid evidence that the impurity

Table 1 EPMA results for LaCoO₃ crystals prepared under various conditions, from the stoichiometric ratio, with added 5% excess of Co, and with additional post high-oxygen- pressure annealing. The formula unit (f. u.) is converted from the averaged raw data normalized

	La (at%)	Co (at%)	O (at%)	f. u. (±0.01)
La:Co	21.14(2)	20.08(5)	58.77(3)	LaCo _{0.95} O _{2.78}
Excess Co Excess Co + HP	20.36(8) 19.96(6)	20.36(4) 19.99(6)	59.26(6) 60.00(9)	$LaCoO_{2.92}$ $LaCoO_{3.00}$

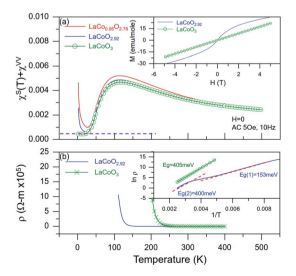


Fig. 3 (a) The magnetic susceptibilities of studied LaCoO_{3 $-\delta$} samples and the corresponding isothermal magnetizations at 2 K are shown in the inset. (b) Resistivities for the as-prepared and annealed LaCoO_{3- δ} are compared, and the inset shows analyzed activation gaps from the Arrhenius plot.

spin of Curie-tail behavior is consistent to the paramagnetic behavior of S = 1/2 from the isolated spins following the Brillouin function. The total number of spin per mole is estimated from the intercept of M (2 K, $H \rightarrow 0$) for LaCoO_{2.92} to be N ~ 3.4%, which is consistent to the vacancy level of $\delta \sim 0.08$ within error, i.e., each oxygen vacancy roughly generates an additional spin 1/2. In contrast, the vacancy-free sample shows no Curietail in $\chi(T)$ and its isothermal magnetization at 2 K shows the constant paramagnetic background without contribution from the additional S = 1/2 spins being aligned by the increasing field.

Yan et al.23 proposed that the low temperature upturn could be coming from the ferromagnetic (FM) behavior of the HS state in their study. However, our results have ruled out the FM interpretation in several aspects: (1) the upturn behavior is removed entirely after high pressure oxygen annealing, which supports the explanation of oxygen vacancy-induced PM behavior, (2) no FM phase transition is identified below \sim 20 K from $C_p(T)$ data to support the existence of a FM phase transition, (3) the size of LaCoO₃ single crystal grown by the FZ method is large enough to obtain a crack-free single crystal sample for the susceptibility measurement by slicing, which significantly reduced the surface-to-bulk ratio. Thus, the surface contribution can be neglected entirely in this study. On the other hand, Pimental et al.,24 pointed out that higher concentration point defects would prevent the valence number of Co ions to reach the perfect +3 valence. Although they proposed that antiferromagnetic coupling between Co³⁺/Co⁴⁺ may induce a weak canted-spin ferromagnetic moment, which lacks microscopic model, it's challenging to apply their interpretation to explain our observations based on current experimental results.

Besides the magnetic property difference shown in Fig. 3(a), an obvious difference between the as-prepared and annealed samples is revealed on the transport behavior also. The onset of metal-to-insulator transition is shown shifted from ~175 K to \sim 275 K after the post-annealing, and the Arrhenius plot shown in the inset of Fig. 3(b) suggests that besides the intrinsic gap of ~400 meV for both, the sample with oxygen vacancies has an additional gap opening of ~153 meV, which is probably related to the formation of a defect-generated impurity band.

B. Direct measurement of χ^{VV} and spin gap Q

Both homogeneous DC and AC spin susceptibilities of pulverized LaCoO₃ ($\delta \sim 0$) crystal samples without Co/O vacancies were measured, and Fig. 4(a) shows $\chi(T)$ data using an AC field of 5 Oe/10 Hz from 2 to 500 K after the core diamagnetic correction $(\chi^{\text{core}} = \chi_{\text{La}} + \chi_{\text{Co}} + 3\chi_{\text{O}} = -0.66 \times 10^{-4} \text{ cm}^3 \text{ mol}^{-1})^{25}$

$$\chi(T) = \chi^{\text{imp}}(T) + \chi^{\text{core}} + \chi^{\text{VV}} + \chi^{\text{S}}(T)$$

where $\chi^{\text{imp}}(T)$ is the impurity spin contribution, χ^{VV} is the Van Vleck paramagnetic contribution, and $\chi^{S}(T)$ is the actual spin contribution. Comparing susceptibilities for the high-oxygenpressure annealed sample LaCoO₃ ($\delta \rightarrow 0$) with those for the as-grown sample of LaCoO_{3- δ} ($\delta \sim 0.08$ -0.22) [see Fig. 3(a)],

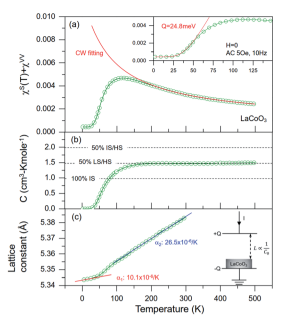


Fig. 4 (a) AC susceptibilities of pulverized LaCoO $_3$ crystal samples after the high-oxygen-pressure annealing. χ^{VV} can be read directly from $T \to 0$ K for an annealed sample of $\delta \to 0$ after the χ^{core} correction. The inset shows the thermal activation behavior of a spin gap $Q \sim 24.8$ meV. (b) The Curie constant is analyzed to show the thermally activated increase of HS population, which reaches $\sim 50\%$ LS/50% HS in a Boltzmann distribution. (c) The threefold increase in the thermal expansion coefficient α across the same temperature range as the spin gap opening is derived directly from the detected capacitance change due to dilation, as illustrated in the inset.

 ${
m Co}^{3+}({
m [Ar]}3{
m d}^6)$ in the ${
m CoO}_6$ octahedral CEF of ${
m LaCoO}_3$ is expected to remain in the nonmagnetic ground state of the LS (${
m t}_2^6{
m ge}_{
m g}^0$) configuration with S=0 below ~ 30 K. The dramatic increase in $\chi^S(T)$ between ~ 30 and ~ 100 K [see Fig. 4(a)] has often been attributed to the increasing number of ${
m Co}^{3+}$ ions being excited to a state of nonzero spin, ${
m ^{28}}$ but the question next is what types of excited spin state of corresponding populations are distributed. Without complications arising from the Curie tail correction, the thermal activation behavior of ${
m exp}(-Q/k_{\rm B}T)$ is fitted reliably for $\chi^S(T)$ below ~ 50 K to show a spin gap of $Q\sim 24.8$ meV [inset of Fig. 4(a)], which is comparable to the values estimated using DC magnetic susceptibility (25 meV), 29 ultrasound (11 meV), 11 and optical absorption (19 meV) on samples of unknown Co or O vacancy level. 30

C. Boltzmann distribution of HS

For LaCoO₃ of $\delta \sim 0$, the $1/\chi^S$ vs. T linear dependence between $\sim 200-500$ K implies a typical PM behavior following the Curie-Weiss law. The Curie-Weiss law fitting of $\chi^S(T) = C/(T-\Theta)$ for $\chi^{VV} + \chi^S(T)$ data between 200 and 500 K [Fig. 4(a)] used a constant fitting parameter of $\chi^{VV} = 4.2 \times 10^{-4}$ cm³ mol⁻¹ read directly from $T \to 0$ K, which satisfactorily returns a Curie constant C = 1.476(5) cm³ mol⁻¹ and a Weiss temperature $\Theta = -131.7(8)$ K. The magnitude and sign of Θ corresponds to the minimum thermal energy required to break the antiferromagnetically coupled spins.³¹ If we follow the conventional data analysis method by assuming there is only one type of spin specie that is responsible for the PM state described by the Curie-Weiss law, the Curie constant $C = N_0 \mu_{\rm eff}^2/3k_{\rm B}$ is contributed from the single spin specie of effective moment $\mu_{\rm eff} = 3.436~\mu_{\rm B}$.

Because the LS($t_{2g}^6 e_g^1$), IS($t_{2g}^5 e_g^1$), and HS($t_{2g}^4 e_g^2$) states have S =0, 1, and 2, respectively, g = 2.42 for IS and g = 1.39 for HS are calculated from $\mu_{\text{eff}} = g\sqrt{S(S+1)}\mu_{\text{B}} = 3.436\mu_{\text{B}}$ under the single specie spin-only assumption. The observation of g > 2 for the first row transition metal element of 3d⁶ filling could be attributed to the incomplete orbital quenching,32 mostly from the unpaired electrons in the triply degenerate t_{2g} $(d_{xy}/d_{yz}/d_{zx})$ orbitals which are operable with 90° orbital rotation. However, the degree of orbital contribution to the effective moment remains unknown theoretically, not to mention that the g < 2value for HS cannot be explained in this context. In addition, the three-fold degeneracy of t_{2g} for the octahedral CEF splitting of LaCoO₃ must have been lifted by the pseudo-cubic distortion already. In order to extract more meaningful information from the measured Curie constant, we propose to use a multicomponent spin-only model analysis that has been proven successful in the interpretation of the magnetic property of Na_rCoO₂ before.33

While the Curie constant represents the sum of all nonzero spins in the PM state without distinguishing the actual spin species of corresponding populations, there are multiple possible distributions of spin size/population that are consistent with the experimental value of

$$\sum_{i} N_{i}\mu_{i}/3k_{\rm B}$$

under the constraint of

$$N_0 = \sum_i N_i = N^{\mathrm{LS}} + N^{\mathrm{IS}} + N^{\mathrm{HS}}$$

where the effective moments $\mu_i^2 = g^2 S_i (S_i + 1) \mu_B^2$ (i = 1, 2, 3) correspond to LS, IS, and HS states, respectively. To analyze Curie constant with multiple spin species spin-only model, *i.e.*, g = 2 in the high-temperature PM regime (for $T > |\Theta| \sim 132$ K) of the full thermally activated stage, the theoretical Curie constants of four possible excited spin distributions for Co^{3+} are calculated to be C = 0.999, 1.999, 2.998, and 1.499 cm³ mol⁻¹ for mixtures of 100% IS, 50% IS/50% HS, 100% HS, and 50% LS/50% HS, respectively, as indicated by the dashed lines shown in Fig. 4(b).

It is interesting to note that the high-temperature saturated value of the Curie constant follows the 50% LS/50% HS distribution perfectly. The 50% LS/50% HS statistical distribution at high temperature is illustrated clearly in Fig. 3(b) by the plot of the temperature dependence of the Curie constant C(T) = N(T) $\mu_{\rm eff}^2/3k_{\rm B} = \chi^{\rm S}(T)x(T-\Theta)$ derived from the Curie–Weiss behavior of $\chi^{\rm S}(T)$, where

$$\frac{N(T)}{N_0} = \frac{N^{\text{HS}}}{N^{\text{HS}} + N^{\text{LS}}} \rightarrow \frac{1}{2}$$

above $\sim 300 \text{ K}$; *i.e.*, $N^{\text{HS}} = N^{\text{LS}} \approx 1 \text{ when } T > 300 \text{ K}$ with thermal energy of $k_{\rm B}T > Q \sim 25$ meV. Although any other transient intermediate state like the IS state cannot be ruled out completely without spectroscopic evidence, the nearly perfect demonstration of thermally activated Boltzmann distribution of a two-level spin system strongly suggests that the spin state of Co³⁺ fluctuates between the LS-HS states statistically. Above all, even when the supplied thermal energy is larger than the gap size, a thermodynamic system must reach its equilibrium state of Boltzmann statistical distribution, instead of activating all spins to the excited state all at once under the consideration of enthalpy reduction only, because the increased entropy is able to lower the Gibbs free energy through $-T\Delta S$ as well.^{5,34} The single spin state assumption without considering the statistical nature of Boltzmann distribution could lead an error of unphysical spin-only moment and overpopulation for the excited state.21

D. Phonon softening and crystalline elasticity

Because of the close relationship between spin state and crystal lattice, there have been a number of detailed investigations reported on the structural properties of nominally stoichiometric LaCoO₃ using temperature-dependent X-ray and neutron diffraction techniques.8,35 with the aim of obtaining a more precise determination of this relationship. Instead of using a diffraction technique for the averaged lattice parameters, we have extracted the linear thermal expansion coefficient $\alpha = (1/2)$ L_0)dL/dT of a single-crystal sample directly with a newly designed capacitance-type device incorporated into the PPMS, as illustrated schematically in the inset of Fig. 4(c). L(T) data were extracted from measurements of the capacitance $C_0 = \varepsilon_0 A/\epsilon_0$ L between 5 and 300 K, which are consistent with the lattice parameters obtained from high-resolution neutron diffraction data with L_0 calibrated to the room-temperature lattice parameter.³⁵ The α values calculated from the slope of L(T) for $T \ge 135$ K ($\alpha_2 \sim 26.5 \times 10^{-6} \text{ K}^{-1}$) are nearly three times larger than those extracted from $T \leq 30 \text{ K} (\alpha_1 \sim 10.1 \times 10^{-6} \text{ K}^{-1})$. It is also

interesting to see that the variation of $\alpha(T)$ is consistent with the temperature dependence of $\chi(T)$ [Fig. 4(a) and Table 2] in three ranges: α is low for $T \leq 30$ K, corresponding to the T-independent $\chi^{\rm VV}$; α is high between Weiss temperature $|\Theta| \sim 132$ K and Debye temperature $\theta_{\rm D} \sim 512$ K, corresponding to the PM behavior of HS spins saturating to the 50% LS/50% HS level; and α behaves nonlinearly between ~ 30 and $|\Theta| \sim 132$ K, corresponding to the growing thermally activated HS population. A strong spin–orbit–phonon coupling is hinted on crossing the temperature range of thermally activated phonon softening.

The bond length of Co^{3+(LS ↔ HS)}-O fluctuation had been observed by Zhang et al. with resonance ultrasound spectroscopy.35 Their discussion about the relaxation processes in terms of thermally activated Debye-like freezing processes is exactly in line with our proposed model for interpretation in this study. Although most of the early works treated the spin states of LS and HS to be a simple thermal activation process for the electrons excited to the unoccupied CEF state, Zhang et al. had examined elastic and inelastic oscillation modes to demonstrate not only spin-spin relaxation but also spin-lattice relaxation dynamically and randomly. The proposed complex local Jahn-Teller deformation with concomitant local FM field is consistent with our proposed symmetry breaking related to the Co³⁺ size variation in time and positions. Although the thermal expansion coefficient (α) goes down to zero at 0 K theoretically without considering the quantum nature of the solid state, the observation of α changes three times in the two regimes in our experiments is consistent with the high resolution synchrotron X-ray results reported by Bull and Knight.36

E. Magnetic specific heat contribution from HS

The specific heats of LaCoO3 between 2 and 300 K were measured with the PPMS using a pulsed heating method, and those between 300 and 600 K were measured with a differential scanning calorimeter (DSC) (STA 449F3, NETZSCH), as shown in Fig. 5(a). The Dulong-Petit limit $3RZ = 3 \times 8.314 \text{ J (mol K)}^{-1}$ \times 5 = 124.71 J (mol K)⁻¹ (Z = 5 for LaCoO₃) is approached above \sim 500 K to be near the Debye temperature $\theta_{\rm D}$. The total speci_c heat includes electronic (C_p^{el}) , phonon (C_p^{ph}) , and magnetic (C_p^{mag}) contributions. Although no long-range magnetic ordering has been identified in LaCoO3, except for the spingapped ground state below ~30 K, the magnetic contribution $C_{\rm p}^{\rm mag}(T)$ in the PM regime should not be excluded, because each spin in the excited HS state must carry magnetic energy higher than its ground LS state. Since the phonon contribution $C_{\mathrm{p}}^{\mathrm{ph}}(T)$ is expected to be reduced after the phonon is softened above ~ 30 K, the loss of $C_p^{\rm ph}(T)[\Delta C_p^{\rm ph}(T)]$ could be transferred to

Table 2 Strongly correlated crystalline, magnetic, and thermodynamic behaviors of LaCoO₃

	$T \leq 30 \text{ K}$	$30 \le T \le \theta \sim 132 \text{ K}$	$ \theta \le T \le \theta_{\rm D}$	$T \ge \theta_{\mathrm{D}} \sim 521 \mathrm{\ K}$
$\alpha(T)$	$lpha_1 \sim 10.1 imes 10^{-6} \; ext{K}^{-1} \ \gamma^{ ext{core}} + \gamma^{ ext{VV}}$	Nonlinear $\gamma^{S}(T) \propto \exp(-Q/T)$	$lpha_2 \sim 26.5 imes 10^{-6} ext{ K}^{-1} \ \gamma^{ ext{S}}(T) \sim C/(T-\Theta)$	
$\chi(T)$ HS%	$\chi + \chi$ ~ 0	$\chi(I) \propto \exp(-Q/I)$ Boltzmann distribution	χ (1) \sim C/(1 - Θ) \sim 50% LS/50% HS	
$C_{ m p}$	$C_{ m p}^{ m el}$ + $C_{ m p}^{ m ph}(T)$	Phonon softening	$C_{ m p}^{ m el}(T)$ + $\left[C_{ m p}^{ m ph}$ $ \Delta C_{ m p}^{ m ph} ight]\!(T)$	Dulong-Pettit limit

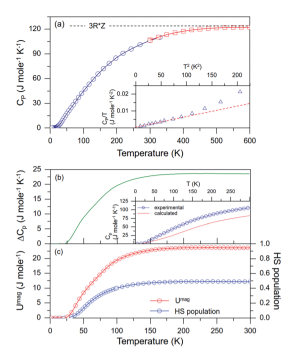


Fig. 5 (a) Specific heat of LaCoO $_3$ from 2 to 600 K, where the Dulong–Petit limit is indicated. The value for $C_p^{\rm e}(T)$ is extracted from T < 10 K as shown in the inset. (b) Significant difference between the measured total $C_p(T)$ and the Debye model calculated $C_p^{\rm ph}(T)$ is found. The difference $\Delta C_p^{\rm ph}(T)$ is attributed to the gain of $C_p^{\rm mag}(T)$. (c) The magnetic internal energy calculated from $U^{\rm mag}(T) = \int_{C_p}^{\rm mag}(T) \mathrm{d}T$ shows a similar temperature dependence to the HS population growth being saturated at the 50% LS/HS level, as estimated independently from the Curie constant data shown in Fig. 4(b).

 $C_{\rm p}^{\rm mag}(T)$ owing to the strong spin-orbit-phonon coupling to reflect the HS population growth, *i.e.*,

$$C_{\rm p}(T) = C_{\rm p}^{\rm el}(T) + C_{\rm p}^{\rm ph}(T)$$

for $T \le 30$ K. On the other hand,

$$C_{\rm p}(T) = C_{\rm p}^{\rm el}(T) + [C_{\rm p}^{\rm ph} - \Delta C_{\rm p}^{\rm ph}](T)$$

and

$$\Delta C_{\rm p}^{\rm ph}(T) \to C_{\rm p}^{\rm mag}(T)$$

for $T \ge 30$ K.

Although it is common to extract the β and Sommerfeld coefficient γ values from the low-temperature limit of $C_{\rm p}$ ($T \leq 10$ K) through the approximations of $C_{\rm p}^{\rm el} \sim \gamma T$ and $C_{\rm p}^{\rm ph} \sim \beta T^3$ ($\gamma \sim 0.3447~{\rm mJ~mol^{-1}~K^{-2}}$ and $\beta \sim 0.068626~{\rm mJ~mol^{-1}~K^{-4}}$), $C_{\rm p}^{\rm ph}(T)$ data after the phonon softening $[C_{\rm p}^{\rm ph} - \Delta C_{\rm p}^{\rm ph}](T)$ above _30 K are not accessible this way. However, we may use the available β value before phonon softening (and thus without magnetic contribution) to construct a $C_{\rm p}^{\rm ph}(T)$ spectrum using the Debye model for the whole temperature range as a baseline. The expected loss of $C_{\rm p}^{\rm ph}(T)$ after phonon softening $[\Delta C_{\rm p}(T)]$ is proposed to be the $C_{\rm p}^{\rm mag}(T)$ gain after phonon softening with HS excitation, i.e., $C_{\rm p}^{\rm mag}(T) = \Delta C_{\rm p}^{\rm ph}(T)$ for $T \gg 30$ K, which can then

be estimated by subtracting the "Debye model constructed $C_{\rm p}^{\rm ph}(T)$ before phonon softening and without magnetic contribution" from the experimental total $C_{\rm p}(T)$.

It should be noted that each data point of $C_{\rm p}^{\rm ph}(T)$ described by the Debye model is non-analytical and can only be obtained through numerical evaluation of the Debye integral using $\theta_{\rm D}/T$ as the upper bound, and the only parameter $\theta_{\rm D} \sim 521$ K is derived from $\beta \sim 12\pi^4RZ/5\theta_{\rm D}^3$ for $T \leq 10$ K before phonon softening. The phonon contribution of $C_{\rm p}^{\rm ph}(T)$ at each temperature could be described with³⁷

$$C_{\rm p}^{\rm ph}(T) = 3RZ \left[\frac{3}{x_{\rm D}^3} \int_0^{x_{\rm D}} \frac{x^4 e^x dx}{(e^x - 1)^2} \right]$$

where $R = N_A k_B$, $x = h\nu/k_B T$, and $x_D = h\nu_D/k_B T = \theta_D/T$. A complete $C_p^{\rm ph}(T)$ spectrum without phonon softening, and thus without $C_p^{\rm mag}(T)$ contribution, is constructed up to 300 K.

 $C_{
m p}^{
m el}(T)+C_{
m p}^{
m ph}(T)$ calculated using the low-temperature estimate of γ and $\beta(\theta_{
m D})$ is shown together with the experimental total $C_{
m p}(T)$ in the inset of Fig. 4(b). Although LaCoO3 has been categorized as an insulator according to its large calculated band gap and the value of γ seems negligible, $C_{
m p}^{
m el}(T){\sim}\gamma T{
m cannot}$ be ignored in the high T range within the Fermi–Dirac statistics [see inset of Fig. 5(a)], otherwise the magnitude of $C_{
m p}^{
m mag}(T)$ would be underestimated. As shown in Fig. 5(b), a significant difference is found between the calculated $C_{
m p}^{
m el}(T)+C_{
m p}^{
m ph}(T)$ without magnetic contribution and the experimental values of total $C_{
m p}(T)$ for $T \geq 30$ K, which can be attributed to the loss of $C_{
m p}^{
m ph}(T)$ after phonon softening, or equivalently, to the gain of $C_{
m p}^{
m mag}(T)$ after phonon softening spin–orbit-phonon coupling. The magnetic internal energy $U^{
m mag}(T)$ can then be evaluated from $C_{
m p}^{
m mag}(T)$ through the integral of

$$U^{
m mag} = \int C^{
m mag}_{
m p}(T) {
m d}T$$

as shown in Fig. 5(c).

It is gratifying to find that a similar temperature dependence is found for the HS population growth from the $\chi(T)$ data analysis independently [Fig. 4(b)], as if the total $U^{\rm mag}$ were coming from the population growth of spins at the HS state carrying a fixed amount of excited energy, *i.e.*, ΔE between eg-t_{2g}. We may trace the magnitude of ΔE using the magnitude of $U^{\rm mag}(300~{\rm K})$ from the population of HS (50% LS/50% HS at 300 K) based on the assumption that each Co³⁺ at HS state contributes ΔE to the magnetic system. ΔE is estimated to be $\sim 0.488~{\rm meV}$ from $U^{\rm mag}(300~{\rm K}) = 23.5~{\rm J}~{\rm mol}^{-1}$ *via*

$$U^{\text{mag}} = 1/2N_{\text{A}} \text{ (mol}^{-1}) \times \Delta E \text{ (eV)} \times 1.6 \times 10^{-19} \text{ (J eV}^{-1})$$

which is in excellent agreement with the value of activation energy $\Delta E = 0.5$ meV fitted from the Schottky-type $C_{\rm p}^{\rm mag}(T)$ data analysis before, ³⁸ although it was viewed "striking" and puzzling by assuming being solely contributed from the \sim 0.1% Co at $T \leq$ 10 K. As a supporting evidence for the identification of such a narrow gap, we find that a similar value of $\Delta E \sim$ 0.6 meV has also been obtained from the zero-field splitting energy of D = 4.9 cm⁻¹ detected using an electron spin resonance (ESR) technique.³⁹

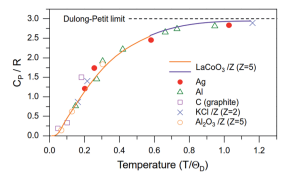


Fig. 6 The reliability of phonon specific heat estimated with the Debye model is confirmed by the scaling of the experimental data of LaCoO $_3$ in $C_{\rm p}(T)/R$ vs. $T/\theta_{\rm D}$, where several representative compounds of different chemical bonding types (reproduced following ref. 43) collapse onto the same scaling relationship.

The difference between the experimentally measured total $C_p(T)$ and the theoretically estimated phonon contribution of $C_p^{ph}(T)$ following the Debye model [see Fig. 5(b)] could be partly coming from the imperfect description of Debye model. However, we find that the reliability and accuracy of Debye model has been demonstrated by the universality of the scaled experimental C_p values collapsed on the theoretical $C_p(T)/R \nu s$. $T/\theta_{\rm D}$ model, 40 which covers samples of chemical bonding types ranging from ionic, metallic, to covalent bondings. We have plotted the experimental $C_p(T)$ data of LaCoO₃ in the scaled form of $C_p(T)/R$ vs. T/θ_D ($\theta_D = 521$ K) as shown in Fig. 6. The universality is clearly demonstrated by the collapsed data points of several compounds with different chemical bonding types after scaling,40 which also overlap with the scaled theoretical Debye model nearly perfectly. The universality of scaled reduced temperature T/θ_D also suggests that the phonon softening induced $\Delta C_{\rm p}^{\rm ph}$ -to- $C_{\rm p}^{\rm mag}$ transfer is absorbed by the parameter of $\theta_{\rm D}$ completely as $\theta_{\rm D} > |\theta|$ (see Table 2), which is consistent to the definition of θ_D as the upper bound of the normal mode for lattice vibration.37 It is also noteworthy to find that the ratio of Debye temperature (θ_D) to melting point (m.p.) for LaCoO₃ $(\theta_{\rm D}/{\rm m.p.}=521/1750~\sim 0.3)$ is only half of that for diamond as a typical rigid covalent compound ($\theta_D/\text{m.p.} = 2230/3700 \sim 0.6$), which once again reacts the unique elastic nature of crystallinity for LaCoO₃.

F. Band gaps and spin gaps

The resistivity as a function of temperature for two representative nonstoichiometric and stoichiometric samples are compared in Fig. 3(b). The onset of sharp resistivity increase as a signature of metal-to-insulator transition for sample with oxygen vacancies is reduced significantly, presumably as a result of the carrier doping created by the oxygen vacancy. According to the Arrhenius plot shown in the inset of Fig. 3(b), besides the high temperature larger activation gap of $\sim\!400$ meV extracted between 200–400 K for both, the sample with oxygen vacancy shows additional activation gap of $\sim\!150$ meV at lower temperature between 80–200 K, which is consistent to a band picture of larger intrinsic CB–VB band gap $\sim\!400$ meV plus

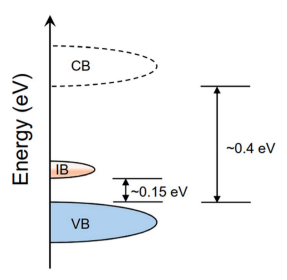


Fig. 7 A proposed band picture of $LaCoO_{3-\delta}$ with an oxygen-deficiency induced p-type impurity band (IB) between the valence band (VB) and conduction band (CB).

a smaller gap \sim 150 meV across the impurity band above VB (ptype doping) or below CB (n-type doping). Based on the Seebeck measurement results (not shown),⁴¹ the carrier doping from oxygen vacancy is confirmed to be p-type, and a schematic band picture is proposed as shown in Fig. 7. The band gap near \sim 150 meV is close to that reported by English *et al.* on a sample having similar Curie-tail behavior.⁴² On the other hand, the larger gap of \sim 400 meV is consistent to the optical gap size extracted from the synchrotron X-ray reflectivity spectrum and that calculated using the density functional theory (DFT) under GGA+U approximation.^{44,45} Mostly importantly, the oxygen vacancy-free sample shows single gap of \sim 400 meV, in contrast to the sample having oxygen vacancy to show two gaps, including the intrinsic CB–VB gap of \sim 400 meV and an additional smaller gap between IB–VB of \sim 150 meV.

Besides the band gaps extracted from the thermal activation behavior of resistivity shown above, two more narrow gaps, including the CEF gap $\Delta E \sim 0.5$ meV estimated from the magnetic internal energy $U^{\rm mag}$ gain [Fig. 5(c)] due to increasing HS population, and the spin gap $Q \sim 25$ meV extracted from the thermal activation behavior of $\chi(T)$ [Fig. 4(a)], are closely related

Table 3 Summary of energy gaps identified in LaCoO $_{3-\delta r}$ including the CEF gap (ΔE) from $C_p(T)$, the thermally activated spin gap (ΩE) from $\chi(T)$, and the band gaps E_{gi} from resistivity data [Fig. 3(b)] of this work. Several other values reported in the literature are compared

(meV)	ΔE	Q	E _{g1} (<80 K)	$E_{ m g2}~{ m (80-200}~{ m K)}$	Едз (200-400 К)
LaCoO ₃ LaCoO _{2,92}	0.5	25		153	405 400
$LaCoO_{3-\delta}$			38 ^a	146^a	$340^b, 310^c$

 a Ref. 45 from resistivity measurement. b Ref. 46 from impedance analysis. c Ref. 48 from DFT calculations.

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to the quantum nature being reflected on the spin state. The band gaps and spin gaps deducted from this work and those reported in the literature are summarized in Table 3. The difference between ΔE and Q is not just one order of magnitude on size, the former is so small that allows spontaneous HS excitation of quantum nature even for $T \to 0$ K. The narrow ΔE is also responsible for the temperature-independent paramagnetic contribution of χ^{VV} as an orbital Zeeman effect for the whole temperature range, ⁴⁶ mostly because the dominant contribution of χ^{VV} must be coming from the narrow CEF gap ΔE of the energy spectrum in the denominator of the second order perturbation for $\chi = \partial^2 E/\partial H^2$ as

$$\chi^{\text{VV}} \propto \sum_{n \neq 0} \frac{|\langle n|\mathcal{H}|0\rangle|^2}{E_n - E_0}$$

where \mathcal{H} is the Zeeman operator.³² In particular, the thermally activated gap Q must have little contribution to χ^{VV} for its much larger gap size and thus little contribution to the perturbation.

It must be stressed that the size of actual CEF-induced energy splitting is often unknown theoretically,⁴⁷ let alone its additional fine splitting due to the octahedral CEF distortion, *e.g.*, the Jahn–Teller effect. Current work using LaCoO₃ sample without low temperature Curie-tail behavior provided a unique opportunity to obtain a reliable χ^{VV} value which is contributed from the quantum fluctuation across the extremely narrow CEF gap ΔE . On the other hand, the thermally activated gap Q is reflected on the HS population growth with the assistance of phonon softening mechanism, which indicates energy transfer from the external thermal energy to the system magnetic energy

by way of spin–orbit–phonon coupling, as also reflected on the tripled thermal expansion coefficient $\alpha(T)$ across the same temperature range [see Fig. 4(c)]. We might view the phenomenon of phonon softening as a prerequisite for the HS population growth that requires longer $(\text{Co-O})^{\text{HS}}$; otherwise, the stronger Coulomb repulsion among electrons would suppress the HS excitation in a rigid lattice of shorter $(\text{Co-O})^{\text{LS}}$ bond length.

G. EELS study: why is the CEF gap so small?

Because the CEF splitting for Co³⁺ ion sitting in the center of a slightly distorted CoO6 octahedral environment remains unknown and has only been deduced to be near 1 eV,32,48 the actual CEF-induced eg-t2g splitting is not likely to be any one of the gaps summarized in Table 3. However, we argue that the actual e_g - t_{2g} splitting corresponds to the narrow gap of $\Delta E \sim 0.5$ meV for LaCoO3, and the thermal activation gap Q for HS excitation is achieved through thermal-strain-magnetic energy exchange with the help of a strong spin-orbit-phonon coupling after the phonon softening. It is reasonable to interpret the Hund's rule for the HS configuration that two electrons are excited to the eg level so that the Coulomb repulsion among electrons in 3d-orbitals is minimized (see Fig. 8). The softened phonon of expanded average interatomic distance is preferable for the HS state of larger ionic size, which has been confirmed by the study thermal expansion as shown in Fig. 4(c). This direction of interpretation basically follows the developmental concepts of small polaron and the breathing type distortion of

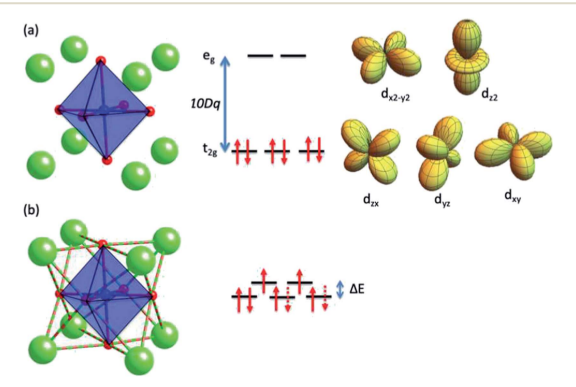


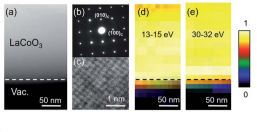
Fig. 8 The schematic plot of 3d orbitals split by the octahedral crystal electric field, (a) without La—O covalency, and (b) with La—O covalency that leads to a more isotropic charge distribution and thus a smaller CEF gap.

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ligands consistently. 5,17,43 The immediate question following would be: why is the CEF splitting (ΔE) so small for LaCoO₃?

Since the origin of an octahedral CEF splitting is coming from the static electric field created by the uneven electron distribution of six ligands surrounding the Co center, a more isotropic environment of negative effective charge of ligands is expected to reduce the CEF size significantly. On examining the ligands surround the Co center of LaCoO₃, although Krapek et al. have proposed that Co-O covalency is responsible for the size reduction of the effective moment for the HS state,49 the covalency of Co-O is not sufficient to assist a more isotropic negative charge distribution, as illustrated by the six dp hybridized orbitals still pointing toward the six neighboring oxygens (see Fig. 8). The most likely source to create less effective negative charge of oxygen ligands is that partial valence electrons of O²⁻ is pulled away from the region along the six Co-O bonding directions and re-distribute more evenly toward the eight La-O bonding directions, as illustrated in Fig. 8(b). It is likely that the La-O bonding is different to the more ionic nature of Na–O and Sr–O in NaCoO₂ and SrCoO_{3- δ}, ^{50,51} *i.e.*, the La-O covalency may not be ignored anymore in LaCoO3. In the study of LaCoO₃, the impact of Sr substitution to the La site has been examined repeatedly in the literature. 16,45,52 However, the possibility of Sr substitution induced oxygen vacancy has often been ignored, which complicates the Co spin state interpretation due to the intertwined factors among local strain, electron count, and vacancy ordering. In order to examine the possibility of La-O covalency, the EELS spectrum of LaCoO3 has been explored for the evidence of plasmonic resonance coming from the paired covalent electrons among La-O.

The STEM-EELS study results of LaCoO₃ are shown in Fig. 9. In contrast to the X-ray photoemission spectroscopy (XPS) which reveals the information of the excitation from the occupied states of valence band, the electron energy-loss spectroscopy (EELS) can provide the information of excitation from both the valence and conduction bands.53 The EELS spectrum contains the excitations of valence electrons in collective modes, including surface and volume plasmons, single-particle excitations, inter-band transitions, and low-lying core-level ionizations.²² Fig. 9(a) and (c) shows the low magnification and atomic resolved high-angle annular dark field (HAADF) images in scanning TEM (STEM), respectively. Fig. 9(b) shows the electron diffraction pattern correlated to Fig. 9(a). Indeed, the atomic resolved HAADF image recorded along [001] direction revealed a pseudo-cubic symmetry which is consistent with literature report.54 Although the ferroelastic domain walls in LaCoO₃ had widely observed in literature, 55,56 however, our HP-LaCoO₃ single crystal exhibits high crystal quality including the crystallinity and domain-free in this study. The EELS spectrum for LaCoO₃ [Fig. 9(f)] reveals several spectral features of incident beam energy loss at \sim 13.9, \sim 30.4 eV, and the \sim 63.3 eV (not shown) from Co M_{2,3} edge excitation. Within the framework of macroscopic dielectric-response theory,22,53 we derived the frequency-dependent dielectric function ($\varepsilon = \varepsilon_1 + i\varepsilon_2$) of LaCoO₃ by performing the Kramers-Kronig Analysis (KKA) as shown in Fig. 9(g). Since ε_1 passes through zero at \sim 13.9 and \sim 30.4 eV with accompanied decrease of ε_2 , which gives rise to the



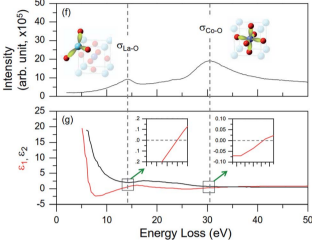


Fig. 9 The STEM-EELS study results of LaCoO $_3$. (a)–(c) The scanned area, diffraction pattern, and atomic scale real lattice image of LaCoO $_3$ along one of the axes of pseudo-cubic symmetry. (d)–(e) The EELS map of two specific energy absorption windows near \sim 13–15 and 30–32 eV. (f) The EELS spectrum of LaCoO $_3$ shows energy absorption of two volume plasmons corresponding to the collective resonance of σ -bond electrons of La–O and Co–O, respectively. (g) The real and imaginary part of the dielectric constant ($\varepsilon = \varepsilon_1 + i\varepsilon_2$) with a blow-up view to show that ε_1 passes zero.

maxima of the volume loss function Im $-1/\varepsilon(\omega)$ and the absorptions near ~ 13.9 and ~ 30.4 eV show characteristics of volume-plasmon excitations and also confirm with highest intensity localized in bulk interior in Fig. 9(d) and (e), respectively, performed by spectrum imaging. The physical origin of the volume plasmons near ~ 13.9 and ~ 30.4 eV are proposed coming from the collective plasmonic resonance of valence electrons among La–O and Co–O, respectively. The EELS evidence of nontrivial La–O covalency supports the scenario that the reduced and isotropic effective negative charge of oxygen ligands is responsible for the significantly reduced octahedral CEF gap size of $\Delta E \sim 0.5$ meV for LaCoO₃.

H. The implication of crystalline elasticity

It is reasonable to assume that the bond length of $(Co-O)^{HS}$ is slightly longer than that of $(Co-O)^{LS}$ for electron filling in the higher e_g energy level of HS $(t_{2g}^4 e_g^2)$ state, which could lead to direct correlation between the magnetic and phonon energies via spin-orbit-phonon coupling, as shown in Fig. 4 and the published high resolution lattice parameters.^{8,35} In particular, when a 3d-electron is excited from t_{2g} to the higher e_g level, concomitant local distortion must occur (see Fig. 10),²⁶ and

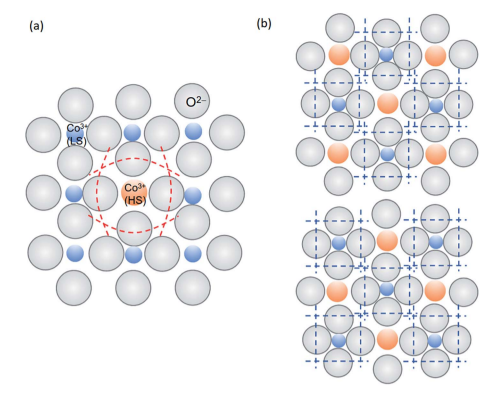


Fig. 10 2D illustrations to show that (a) low percentage of HS state of larger ion size would create local distortion and strain within the LS state background, and (b) the strain is relieved when the HS population is reaching 50%, which allows a breathing mode energy exchange between the magnon and phonon.

a local breathing mode is established between the magnetic Zeeman energy and the phonon energy of ferroelectric origin. The dilute distribution of HS spins below $\sim \! \! 30$ K exhibits a low α with a rigid and homogeneous distribution of shorter (CoO) bonds. As the HS population grows with increasing temperature, the initial randomly distributed low population of HS would create high strain to the lattice, until nearly half of the Co ions are excited to the HS state at high temperature.

A more homogeneous magnetic-phonon energy exchange is achieved for the 50% HS/50% LS distribution of minimum lattice strain with well-defined threefold increase of α (see Fig. 4). The reduction of rigidity is prerequisite for the 50%(Co–O)^HS/50%(Co–O)^LS statistical distribution under thermal fluctuations, which is also reflected on the increase of electron entropy. 58

Yamaguchi *et al.* have proposed that e_g orbital ordering (OO) may exist in the Co^{3+} IS state $(t_{2g}^5e_g^1)$ of $LaCoO_3$ having lifted orbital degeneracy in e_g , 7 *i.e.*, the electron in e_g prefers to occupy either $d_{x^2-y^2}$ or d_{z^2} orbital in an ordered pattern (see Fig. 8), which has also been applied to explain the observed split Co–O stretching mode as local distortion arising from the Jahn–Teller effect. Comparing these two proposed model interpretations for the local distortion due to Co–O bond length fluctuation, the controversial IS state could be explained equally well by either the OO of an IS state, or the 50% LS/50% HS statistical distribution in time average. Above all, the distinction between HS and IS is not possible in the high temperature metallic phase anyway. ⁴⁹

IV. Conclusion

In conclusion, based on an integrated study of the magnetic susceptibility, specific heat, and thermal expansion of a single crystal LaCoO₃ nearly free of cobalt and oxygen vacancies, we propose a consistent interpretation of strongly correlated spin state and crystal lattice for the pseudo-cubic LaCoO₃ perovskite. The puzzling deviation of the crystal symmetry from cubic to rhombohedral symmetry as a relaxed pseudo-cubic structure is related to the LS/HS spin state transition with corresponding statistically distributed (Co–O)^{LS}–(Co–O)^{HS} bond length fluctuation.

Data availablity

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

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