Orthorhombic distortion and orbital order in the vanadium spinel FeV₂O₄

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Using synchrotron and neutron diffraction measurements, we find a low-temperature orthorhombic phase in vanadium spinel FeV_2O_4 . The orbital order of V^{3+} ions with tetragonal normal modes occurs at 68 K, and this leads to an appearance of the pseudotetragonal phase at a noncollinear ferrimagnetic transition temperature. Below the magnetic transition temperature, unconventional behavior of the orbital state of Fe^{2+} ions accompanied by the emergence of the orthorhombic phase was observed by using the normal mode analysis. We have also studied the structural properties of orbitally diluted materials. The orthorhombic phase, which is significantly affected by the other ions, is intrinsic in FeV_2O_4 . We suggest the orthorhombic phase is strongly related with the double orbital states of Fe^{2+} and V^{3+} ions.

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I. INTRODUCTION

Obtaining an accurate crystal structure is critically important for understanding the interplay of orbital and spin degrees of freedom and lattice distortions, because it provides significant clues to the understanding of the behavior of its lattice degrees of freedom [1,2]. Recently, vanadium spinel oxides AV_2O_4 (A = Mn, Co, Fe, Zn, and Cd), which have the V³⁺ (3d²) ions in triply degenerate t_{2g} levels at octahedral sites (B sites), have been intensively investigated as one of the forefront topics in modern orbital physics.

Most of these compounds exhibit a cubic to compressed tetragonal (c < a) lattice distortion caused by the orbital order (OO) of V^{3+} ions at low temperature, the origins of which have been experimentally and theoretically studied [3-11]. However, FeV₂O₄ is an unconventional material because an elongated tetragonal phase (c > a) appears at low temperature, and the A sites occupied by Fe^{2+} ions have orbital degrees of freedom in the *e* states [12]. Hence, FeV_2O_4 has Jahn-Teller (JT) active ions at both the A sites and the B sites; thus, the competition of ions, the cooperation of ions, or both induce multistep structural and magnetic transitions [11,12] accompanied by unique physical properties, i.e., multiferroic nature, large magnetostriction, and anomalous magnetization [12–17]. Therefore, a detailed structural study of FeV_2O_4 is important for understanding the origin of physical properties and the contribution to orbital physics. However, the OO models of V^{3+} ions at T_{N2} , suggested by Nii *et al.* [11] and Kang et al. [18], are contradictory. Additionally, the origins and structural properties of the low-temperature phase including the elongated tetragonal phases [12] are still disputed despite numerous experimental and theoretical efforts [11–21]. In this paper, we tackle the accurate structure analysis with high-resolution synchrotron diffraction and neutron diffraction by using polycrystalline samples as well as single crystals of FeV_2O_4 and orbitally diluted materials for each A and B site.

II. EXPERIMENTAL

The powder samples of FeV_2O_4 , $Fe_{1-r}Mn_rV_2O_4$, and $FeV_{2-r}Fe_rO_4$ used in this study were prepared by solid-state reaction from a stoichiometric mixture of high-purity Fe₂O₃, Fe, V₂O₃, and MnO powder. The powder was heated in an evacuated quartz tube at 1323 K for 24 h in a furnace, and the resultant sample was subsequently ground and pressed into pellets under 35 MPa pressure. The pellet was sintered in an evacuated tube at 1373 K for 24 h. The single crystal was grown using a floating-zone method (see Appendix A). The compositions of these samples were measured by energy dispersive x-ray spectroscopy. The compositions in powder samples almost correspond to those for preparation; however, excess Fe of 5%-10% exists in the single crystal, which is consistent with the past studies [12,17]. In this paper, we express the single crystal of FeV₂O₄ as $Fe_{1+\delta}V_{2-\delta}O_4$ single crystal (see Appendix A).

Synchrotron powder diffraction (SPD) experiments were carried out with a large Debye-Scherrer camera installed at the beamline BL02B2 at SPring-8 [22]. The wavelength of the incident x ray was 0.79790 Å. The sample temperature was controlled by the He/N₂ gas flow devices. The single-crystalline synchrotron diffraction experiment was performed at BL-4C with an off-center-type four-circle diffractometer (Photon Factory, KEK). The x-ray beam was monochromatized to 10.33 keV (1.2 Å). The single crystal ($2.5 \times 2.0 \times 1.5 \text{ mm}^3$) was mounted with the [100]_c axis parallel to the scattering vector \boldsymbol{q} in the closed-type cryostat. The neutron powder diffraction (NPD) measurements were carried out at high-flux (G41) and high-resolution powder diffractometers

(3T2) at the Laboratoire Léon Brillouin and the Institut Laue-Langevin. The wavelengths of incident beams at G41 and 3T2 were 2.42834 and 1.22494 Å, respectively. The magnetization and specific heat measurements were performed by the SQUID magnetometer (MPMS-XL) and the physical property measurement system, respectively.

III. SYNCHROTRON AND NEUTRON DIFFRACTION MEASUREMENTS

Figure 1(a) shows the synchrotron powder diffraction (SPD) and neutron powder diffraction (NPD) patterns around the 440 cubic Bragg reflections at low temperatures. In the SPD patterns, the distinguishing feature is that three peaks are observed at 15 K, indicating an orthorhombic phase. In contrast, the peak splitting at other temperatures is similar to that in past studies [12]. In the NPD study, signs of these transitions were also observed, consistent with recent literature [13,20,21]. However, the peak splitting was not observed as clearly as in the SPD study because of the lower angular resolution of NPD. Therefore, we mainly focused on SPD experiments. We performed structural refinements by the Rietveld method and obtained the crystallographic parameters listed in Tables I and II. Figures 1(b) and 1(c) give the results of the Rietveld fitting in the SPD pattern at 15 K and the NPD pattern at 2 K, respectively. These refinements are best described by the orthorhombic model with the space group Fddd. This space group is a subgroup of the tetragonal space group $I4_1/amd$, and it has been often applied to describe structural ground states of the spinels with JT active ions located at the A site [23]. The characteristic atomic displacement (mentioned in Sec. IV) was found in the low-temperature orthorhombic (LTO) phase, whereas the spin alignments and moment sizes for the Fe^{2+} and V^{3+} cations [Fig. 1(d)] were almost the same as those in the canted-ferrimagnetic structure reported in the literature [13].

The temperature dependencies of magnetization and specific heat are shown in Fig. 1(e), and those of the lattice constants, which are calculated from SPD data, are shown in Fig. 1(f). These results indicate structural transitions from cubic to high-temperature tetragonal (HTT; c < a) at $T_{S1} = 137$ K, HTT to high-temperature orthorhombic (HTO) at $T_{\rm N1} = 106$ K, HTO to low-temperature tetragonal (LTT; c > a) at $T_{N2} = 68$ K, and LTT to LTO at $T_{S2} \approx 30$ K. Note that the crystal axes in the HTT and LTT phases are defined as $\vec{a}_{\text{HTT}} = (1/2)(\vec{a} - \vec{b}), \ \vec{b}_{\text{HTT}} = (1/2)(\vec{a} + \vec{b}), \ \vec{c}_{\text{HTT}} = \vec{c}, \ \vec{a}_{\text{LTT}} =$ $(-1/2)(\vec{c}+\vec{b}), \vec{b}_{LTT} = (-1/2)(\vec{c}-\vec{b}), \text{ and } \vec{c}_{LTT} = \vec{a}, \text{ where } \vec{a},$ \vec{b} , and \vec{c} are the cubic lattice vectors, and the lengths of \vec{a} , \vec{b} , and \vec{c} are shown in Fig. 1(f) as a, b, and c, respectively. Considering previous neutron diffraction experiments [13], T_{N1} and T_{N2} represent collinear ferrimagnetic (C-FM) and noncollinear ferrimagnetic (NC-FM) orders, respectively, accompanied by lattice distortions. The magnetic structures refined using our NPD data agree well with those in the literature [13]. However, an anomaly accompanied by the emergence of the LTO phase was not observed in the temperature dependencies of magnetization and specific heat; the reason for this will become evident later. We also performed single-crystalline synchrotron diffraction; however, an indication of lattice distortion from LTT to LTO below 30 K was not clearly detected (see



FIG. 1. Temperature dependence of the (a) SPD and NPD patterns around 440 cubic Bragg reflection. Rietveld fitting for (b) SPD pattern at 15 K and (c) NPD pattern at 2 K in FeV₂O₄. The crosses, solid lines, and bottom lines represent the observed intensities, the calculated ones, and the differences, respectively. The vertical bars show the position of Bragg peaks (for the NPD pattern, the long and short bars indicate the crystal and magnetic reflections, respectively) with the Fddd space group. The temperature dependencies of (d) the ordered moment size on the Fe^{2+} and V^{3+} cation sites, (e) the magnetization (left axis) as well as the heat capacity (right axis), and (f) the lattice constants in FeV₂O₄. The lattice constant in the temperature range of $T_{S2} \leq T \leq T_{N2}$ was calculated by using the LTT model with the 141/amd space group. Note that the lattice constants in the HTT and LTT phases are shown by the cubic axes for comparison. Typical errors of the lattice constants are less than the size of the data markers. Note that the moments below T_{N1} are calculated using the *Fddd* model.

Appendix B) presumably because of excess Fe in the single crystal due to evaporation of Fe and separation of V_2O_3 in the synthesis (see Appendix A).

IV. ORBITAL STATES OF Fe²⁺ AND V³⁺ IONS

To understand the origin of structural phase transitions, we investigated the detailed orbital states of both Fe^{2+} and V^{3+} ions in the LTO phase as well as other phases, and we performed normal (*Q*) mode analysis, which is related

	Cubic	Cubic	HTT	HTO	LTT	LTT	LTO
T	300 K	150 K	115 K	100 K	60 K	40 K	15 K
$a(\text{\AA})$	8.45361(5)	8.44846(7)	6.0062(2)	8.5278(2)	5.93142(6)	5.93289(8)	8.5277(1)
b(Å)	8.45361	8.44846	6.0062	8.4703(2)	5.93142	5.93289	8.4091(1)
c(Å)	8.45361	8.44846	8.3472(2)	8.3398(2)	8.5299(1)	8.5264(1)	8.3687(1)
Ζ	8	8	4	8	4	4	8
Fe ^a	8a	8a	4a	8a	4a	4a	8 <i>a</i>
$B(\text{\AA}^2)$	0.75(1)	0.46(1)	0.46(2)	0.44(2)	0.39(2)	0.33(4)	0.27(2)
V ^b	16 <i>d</i>	16 <i>d</i>	8d	16 <i>d</i>	8d	8d	16 <i>d</i>
$B(\text{\AA}^2)$	0.56(1)	0.36(1)	0.50(2)	0.46(2)	0.38(1)	0.22(3)	0.32(2)
0	32 <i>e</i>	32 <i>e</i>	16h	32h	16 <i>h</i>	16 <i>h</i>	32h
x	0.3867(1)	0.3867(1)	0	0.3879(4)	0	0	0.3910(2)
у	0.3867	0.3867	0.4747(4)	0.3862(4)	0.4827(2)	0.4843(6)	0.3842(3)
z	0.3867	0.3867	0.2584(3)	0.3823(3)	0.2662(1)	0.2656(4)	0.3831(2)
$B(\text{\AA}^2)$	0.67(3)	0.59(3)	0.28(3)	0.31(3)	0.18(2)	0.10(6)	0.11(3)
$R_{\rm WP}$ (%)	4.75	5.50	4.81	5.03	2.44	6.54	3.09
$R_{\rm I}$ (%)	3.31	2.89	3.59	3.60	2.33	2.41	2.59

TABLE I. Structural parameters in FeV₂O₄. The cubic and orthorhombic phases have the space groups $Fd\overline{3}m$ and Fddd (origin choice 1), respectively, and then the tetragonal phases have the space group $I4_1/amd$ (origin choice 2).

^aFractional coordinates (x, y, z) in the 8a and 4a sites are (0, 0, 0) and (0, 3/4, 1/8), respectively.

^bFractional coordinates (x, y, z) in the 16d and 8d sites are (5/8, 5/8, 5/8) and (0, 0, 1/2), respectively.

to the local distortions of the ligands [11,24–28]. Figure 2 represents the distortions of the FeO₄ tetrahedron and the VO₆ octahedron in the two-dimensional Q_2 - Q_3 plane, where Q_2 and Q_3 correspond to orthorhombic and tetragonal distortions, respectively.

First, we focus on the distortion of the FeO₄ tetrahedron, considering the Q_2 and Q_3 modes related to the *e* orbital states. A negative Q_3 ($\theta = \pi$ related to the $3z^2 r^2$ type orbital) mode was observed at 115 K, and a positive Q_2 mode appeared at 100 K. Subsequently, at 60 K, the $y^2 r^2$ orbitals are stabilized where $\theta = 2\pi/3$. These behaviors are explained by the cooperative JT effect at T_{S1} and the relativistic spin-orbit (SO) coupling at T_{N1} of the Fe²⁺ ions, which stabilize the $3z^2 r^2$ and $y^2 r^2$ orbitals [11,25,27], respectively. Here, we introduce the deviation angle $\Delta\theta$ from $\theta = \pi$ (Fig. 2), which indicates the variation of orbital states between $3z^2 r^2$ and $y^2 r^2$ orbitals. The most distinguishing feature is the variation of $\Delta\theta$ from 60° at 60 K to 52° at 15 K. This unconventional behavior, which was not seen in other materials with orbital degrees of freedom, is likely related to the two orbitally active ions.

Next, the distortions of the VO₆ octahedron are discussed in the Q_2 - Q_3 plane. The VO₆ octahedron was compressed along the *x* direction below T_{N2} , similar to the behavior observed in the Fe_{1+ δ}V_{2- δ}O₄ single crystal [11]. This distortion is very similar to that derived from the ferro-OO type where one electron occupies the *yz* orbital and other occupies the $(xy + izx)/\sqrt{2}$ orbital [11]. We found that the Q_2 and Q_3 values of the VO₆ octahedron at 15 K are almost equal to those at 60 K.

V. ORTHORHOMBIC DISTORTION

To investigate the nature of the orthorhombic lattice at the ground state and the variation of $\Delta \theta$ below T_{N2} , we examined the SPD profiles of FeV₂O₄ in detail. The crosses in Fig. 3(a)

	Cubic	Cubic	HTT	HTO	LTT	LTT	LTO
T	300 K	150 K	115 K	100 K	60 K	40 K	15 K
O-Fe-O angle (deg)	109.47(7)	109.47(7)	103.43(9)	107.3(1)	105.56(7)	105.5(1)	105.65(9)
			108.1(1)	108.9(1)	111.46(6)	111.5(1)	110.91(9)
				112.4(1)			111.93(9)
V-O distance (Å)	2.0189(9)	2.0181(9)	2.020(2)	2.025(4)	1.997(1)	2.001(4)	1.998(2)
			2.022(3)	2.027(4)	2.0307(8)	2.037(3)	2.031(2)
				2.029(4)			2.031(2)
O-V-O angle (deg)	84.22(7)	84.25(7)	83.83(9)	84.1(1)	83.96(7)	84.3(1)	83.86(9)
	95.78(7)	95.75(7)	84.8(1)	85.1(1)	85.66(6)	86.1(1)	84.12(9)
			95.2(1)	85.4(1)	94.34(6)	93.9(1)	85.65(9)
			96.17(9)	94.6(1)	96.04(7)	95.7(1)	94.35(9)
				95.0(1)			95.88(9)
				95.8(1)			96.14(9)

TABLE II. O-Fe-O bond angle, V-O bond distance, and O-V-O bond angle in FeV_2O_4 .



FIG. 2. Normal mode analysis in the Q_2 - Q_3 plane of the FeO₄ tetrahedron (solid circles) and VO₆ octahedron (squares) in the powder sample of FeV₂O₄. The triangles represent the Q_2 and Q_3 modes of the orthorhombic model for the P-tetra phase at $T_{S2} < T < T_{N2}$. Note that the Q_2 and Q_3 values, which are calculated from the SPD data, are only shown for clarity.

show the profile at 40 K around the 440 cubic Bragg peak. The 440_c reflection is significantly broader than that of 044_c , where the subscript "c" indicates the Miller indices based on the cubic axis. Additionally, the temperature dependence of the ratio of the full width at half maximum (FWHM) for the 440_c to 044_c reflections is shown in Fig. 3(b). The FWHM of 440_c is the same as that of 044_c at $T_{N1} < T < T_{S1}$, whereas the FWHM ratio below T_{N1} (even in the LTT phase) is significantly larger than that in the HTT phase. This result indicates that the crystal structure in the LTT phase has orthorhombic symmetry with similar lattice constants for b and c ($b \approx c$). Furthermore, all Bragg reflections were expressed by Rietveld fitting, using the Fddd model, as shown in Figs. 3(a) and 3(c). The fitting is fairly good, and the detailed structural parameters are listed in Table III. Here, we regard the LTT phase as a pseudotetragonal (P-tetra) system with an orthorhombic lattice. The temperature dependencies of the lattice constants considering the P-tetra phase are shown in Fig. 3(d).

Structural refinements and normal mode analysis were also performed by using the orthorhombic model for the P-tetra (LTT) phase (Fig. 2; see triangles). We found only a slight change (from $\Delta\theta = 60^{\circ}$ to 58°) for the FeO₄ tetrahedron at 68 K, as reflected by the orthorhombic structural models. However, the uniaxial distortion of the VO₆ octahedron, which is associated with tetragonal normal modes and induced by the OO of V³⁺ ions, was not significantly influenced by the models. Here, we suggest that the behavior of $\Delta\theta$ up to near 60° is caused by the ferro-OO of V³⁺ ions accompanied by the NC-FM order. The OO of V³⁺ ions likely affects the orbital state of the Fe²⁺ ion and leads to stabilization of the y^2-z^2 orbital because it is associated with tetragonal normal modes at T_{N2} . For the variation of the lattice, the decreasing of the *a*



FIG. 3. (a) Rietveld fitting by using the orthorhombic model around the 440 cubic Bragg peak for SPD patterns in FeV₂O₄ at 40 K. The subscripts "c" and "o" indicate the cubic and orthorhombic axes, respectively. (b) Temperature dependence of the ratio of the FWHM for 440_c to that for 044_c reflections. In the orthorhombic phases, the FWHM of the 440_c reflection is expressed by overlapping the 440_c and 404_c reflections. (c) Rietveld analysis of FeV₂O₄ for the SPD pattern in the P-tetra phase at 40 K. The profiled-weighted *R* factor *R*_{WP} and the *R*-Bragg factor *R*₁ represent 4.45% and 2.55%, respectively. (d) Temperature dependencies of lattice constants for FeV₂O₄ considering the orthorhombic lattice in the P-tetra phase. Note that the lattice constant in the temperature range of $T_{52} \leq T$ $\leq T_{N2}$ was calculated by using P-tetra model with the *Fddd* space group. The orbital states of Fe²⁺ ions and the spin alignments of Fe²⁺ and V³⁺ ions are also illustrated schematically.

and *b* axes and the increasing of the *c* axis [see arrows in Fig. 3(d)] around T_{N2} with decreasing temperature are most likely influenced by the OO of V³⁺ ions with the tetragonal normal mode of the VO₆ octahedron at T_{N2} . Below T_{N2} , the orbital state of Fe²⁺ ions (see orbital states at 60 K and 15 K in Fig. 2) gradually changes with decreasing temperature, and then the orthorhombic structure is stabilized at the ground state. These results indicate that the lattice constants continuously change below T_{N2} [Fig. 3(d)]. This continuous change of the lattice for

TABLE III. Structural parameters of FeV_2O_4 for the P-tetra phase, where structural refinements are performed using the orthorhombic model with the space group *Fddd* (origin choice 1).

	P-tetra 60 (K)	P-tetra 40 (K)		
a (Å)	8.53240(8)	8.5344(2)		
b (Å)	8.39482(9)	8.4045(2)		
<i>c</i> (Å)	8.3838(1)	8.3823(2)		
Fe	8a(0, 0, 0)	8a(0, 0, 0)		
$B(\text{\AA}^2)$	0.38(2)	0.28(2)		
V	16 <i>d</i> (5/8, 5/8, 5/8)	16 <i>d</i> (5/8, 5/8, 5/8)		
$B(\text{\AA}^2)$	0.28(2)	0.26(2)		
0	32h	32h		
x	0.3912(2)	0.3911(2)		
у	0.3836(6)	0.3841(6)		
z	0.3834(6)	0.3833(6)		
$B(\text{\AA}^2)$	0.44(4)	0.25(4)		

the temperature is consistent with the results of magnetization and specific heat measurements, which show no anomalies at T_{S2} in Fig. 1(e). The summary of the phase transitions and orbital states of both Fe²⁺ and V³⁺ ions are also schematically illustrated in Fig. 3(d).

VI. ORBITALLY DILUTED MATERIALS

To ensure our suggestion that the crystal system of FeV₂O₄ at the ground state has the orthorhombic lattice, we consider the SPD profiles in the substituted materials $(Fe_{1-x}Mn_x)[V_2]O_4$ and $(Fe)[V_{2-x}Fe_x]O_4$ (Here, parentheses and square brackets indicate the A- and B sites, respectively, in the spinel structure.). The detailed physical properties of these compounds are provided in Refs. [29-32]. Figure 4(a) represents the SPD patterns around the 440 cubic Bragg peaks at 15 K for $(Fe_{1-x}Mn_x)[V_2]O_4$. As x increases, the $440_{\rm c}$ reflection becomes sharper than that for x = 0.05. The compositional dependencies of the FWHM ratio of the $440_{\rm c}$ to 044_c reflections are shown in Fig. 4(b). The ratio decreases as x increases and is significantly different from unity when $x \leq 0.2$, whereas the FWHM for the 440_c reflection is almost equal to that for the 044_c reflection when x > 0.2. Note that the orthorhombic and compressed tetragonal phases are absent when $x \ge 0.3$ in these materials [29–32]. From these results, the crystal system is logically regarded as the P-tetra rather than the elongated tetragonal (c > a) lattice until x = 0.2.

We examined the distortions of the FeO₄ tetrahedron and the VO₆ octahedron of Fe[V_{1.9}Fe_{0.1}]O₄ in the Q_2 - Q_3 plane. Note that the structural details are provided in a previous study [29]. Figure 5 gives the distortion of the FeO₄ tetrahedron and the VO₆ octahedron in the Q_2 - Q_3 plane for polycrystalline samples of FeV₂O₄ and Fe[V_{1.9}Fe_{0.1}]O₄ and the Fe_{1+ δ}V_{2- δ}O₄ single crystal. Note that the structural parameters of the single crystal are found in Ref. [29]. The distortion of the FeO₄ tetrahedron of polycrystalline samples of FeV₂O₄ below T_{N2} is significantly different from that of the single crystal, and the unprecedented behavior ($\Delta\theta$ from 60° to 52°) disappears in the single crystal. The most distinguished feature is that



FIG. 4. (a) SPD patterns around the 440 cubic Bragg peak for $Fe_{1-x}Mn_xV_2O_4$ at 15 K. (b) Ratios of FWHM for the 440_c reflection to that for the 044_c reflection in $Fe_{1-x}Mn_xV_2O_4$ and $Fe[V_{2-x}Fe_x]O_4$.

the behavior of the Q_2 and Q_3 modes in Fe[V_{1.9}Fe_{0.1}]O₄ is very similar to that of the Fe_{1+ δ}V_{2- δ}O₄ single crystal. This is explained by excess Fe in FeV₂O₄, which is often observed in the single crystal and occupies the *B* site until x = 0.35 [12,17,29,33,34]. Additionally, the orbital degrees of freedom of t_{2g} states at the V³⁺ site are diluted by the excess Fe. The excess Fe in FeV₂O₄ makes the magnetic phase transition temperature T_{N2} decrease. When the excess Fe of 10% is occupied in FeV₂O₄, T_{N2} becomes 61 K [29]. This temperature well agrees with that in our Fe_{1+ δ}V_{2- δ}O₄ single crystal and a reported one [13,17]. Therefore, it is conceivable that the excess Fe always exists in the single crystal, and crystal growth of the single crystal with stoichiometry is very difficult because of the evaporation of Fe and the separation of V₂O₃ (see Appendix A).

On the other hand, we investigated the behavior of the Q_2 and Q_3 modes in Fe_{1-x}Mn_xV₂O₄. The aim of this investigation is opposed to that in (Fe)[V_{2-x}Fe_x]O₄ because the *e* states at the Fe²⁺ site are diluted by the Mn²⁺ ion with nonorbital orbital degrees of freedom. Figure 5 shows the distortion of the FeO₄ tetrahedron in the Q_2 - Q_3 plane for polycrystalline samples of Fe_{0.9}Mn_{0.1}V₂O₄. Note that the details of the structural parameters are listed in the literature [31]. Below T_{N2} , the behavior of $\Delta\theta$ from 60° at 60 K to 52° was not shown, and this result is similar to results in Fe[V_{1.9}Fe_{0.1}]O₄ and the Fe_{1+δ}V_{2-δ}O₄ single crystal. Here, we focus on the magnitude



FIG. 5. (a) Normal mode analysis in the Q_2 - Q_3 plane of the FeO₄ tetrahedron and the VO₆ octahedron for Fe[V_{1.9}Fe_{0.1}]O₄ and the Fe_{1+ δ}V_{2- δ}O₄ single crystal. Note that the Q_2 and Q_3 values in the single crystal are from Ref [11]. (b) Q_2 and Q_3 modes of the FeO₄ tetrahedron for FeV₂O₄, Fe[V_{1.9}Fe_{0.1}]O₄, and Fe_{0.9}Mn_{0.1}V₂O₄. PC and SC denote the polycrystalline and the single-crystal samples, respectively.

of distortions of the FeO₄ tetrahedron and the VO₆ octahedron at 15 K in $Fe_{1-x}Mn_xV_2O_4$ [31]. The magnitudes of distortions of not only the FeO₄ tetrahedron but also the VO₆ octahedron are changed by the quantity of the substitution of the Mn^{2+} ion. This result indicates that the orbital state of the Fe^{2+} ion is affected by that of the V3+ ions. In these viewpoints, we suggest that not only OO of the Fe²⁺ ions but also that of the V^{3+} ions plays an important role in the appearance of the LTO phase. The other possible factors for suppression of the LTO phase by the substitution are the change in ionic sizes and that in valence states. In the vanadium spinel oxides, it is well known that the bond length between vanadium ions V-V is strongly related with the localized-itinerant electronic transition, e.g., enhancement of the ferrimagnetic transition and suppression of the structural distortion [35-37]. In this study, when the ionic sizes of Fe^{2+} and V^{3+} ions are changed by the substitution of the Mn^{2+} and Fe^{3+} ions, respectively, the V-V lengths also change. These variations can affect the electronic states in FeV₂O₄. On the other hand, the effect of the change in the valence state may be much smaller because the ionic valences at both the *A* site and the *B* site do not change from those of FeV₂O₄ by the small amount of the Mn^{2+} and Fe³⁺ substitution (i.e., those of the *A* site and the *B* site are 2+ and 3+, respectively) [31,34]. Therefore, the appearance of the LTO phase was possibly suppressed by the effects of not only the orbital dilution but also the change in the ionic sizes by the chemical doping. Thus, the orthorhombic lattice distortion accompanied by unprecedented behavior of orbital states of the Fe²⁺ ions at the lowest temperatures is suppressed by an impurity, and we conclude that the orthorhombic system at 15 K is an intrinsic phase in FeV₂O₄.

Meanwhile, a trigonal distortion of the VO₆ octahedron has been suggested to play an important role in the OO of the V^{3+} ions and the NC-FM order at T_{N2} [20]. Therefore, the appearance of the P-tetra lattice accompanied by the OO of V³⁺ ions may be related to the trigonal distortion. However, we did not observe such a distortion at T_{N2} , consistent with a previous study [11]. Our crystallographic studies experimentally and/or theoretically call for a reinvestigation of the physical properties in FeV_2O_4 because the orbital state of V^{3+} ions and the origin of the magnetostructural transition are considered only in tetragonal models [4,5,9,10]. Taking into account the orthorhombic structure, a detailed calculation of electronic states, local structure analysis by using highresolution transmission electron spectroscopy, and extended x-ray absorption fine structure analysis and high-resolution neutron diffraction experiments will contribute to a further understanding of the orbital physics in vanadium spinels.

VII. CONCLUSIONS

We found the LTO phase below 30 K in the polycrystalline sample of FeV_2O_4 . The LTO phase was intrinsic in FeV_2O_4 because the formation of this phase was significantly affected by the impurity. We performed the structural refinements for SPD and NPD patterns using the Fddd model, and we investigated the orbital states of the Fe^{2+} and V^{3+} ions using the normal mode analysis. The unprecedented behavior of orbital states of Fe²⁺ ions below T_{N2} was observed. The LTO phase accompanied by this behavior disappears when the Fe²⁺ site and the V^{3+} site are occupied by the excess Fe and Mn²⁺ ions with nonorbital degrees of freedom, respectively. Hence, the formation of the LTO phase is related with double orbital states of Fe^{2+} ions and \tilde{V}^{3+} ions. These double degenerate OOs of Fe^{2+} and V^{3+} ions induce the unconventional structural properties below T_{N1} accompanied by continuous variation of the orbital state of Fe^{2+} ions.

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APPENDIX A: SINGLE CRYSTAL GROWTH

The crystal growth of FeV₂O₄ was carried out in an argon atmosphere in an infrared-heated image furnace installed with four halogen lamps having a power of 1500 W and four ellipsoidal mirrors. Figure 6 shows the quartz tube and samples in the floating-zone (FZ) system before and after crystal growth. After the growth, the black powder attaches to the inner surface of the quartz tube. This is derived from the evaporation from the rods during the crystal growth. Kismarahardja et al. [15] claimed that the evaporated powder is V_2O_3 with the corundum structure, and extra V_2O_3 in the starting materials is critical to obtain high-quality samples. However, the x-ray diffraction pattern for the black powder indicates the body-centered cubic structure with a lattice constant of 2.86 Å, as shown in Fig. 7(a). Additionally, we confirmed using a ferrite magnet that the black powder represents ferromagnetism at room temperature. Considering the chemical species of FeV₂O₄, we concluded that the black powder is not V_2O_3 but rather the single phase of pure Fe.

Next, the inset of Fig. 7(a) represents the section of the seed rods after the crystal growth. The crystallinity in the inner region of growth crystal is better than that in the outer region. Accordingly, we performed the x-ray diffraction experiments using the growth crystal, which is ground into a fine powder, as shown in Fig. 7(a). In the outer of the seed rod, the peaks derived from the impurity of V_2O_3 are observed. This impurity may be caused by the evaporation of Fe. On the other hand, the inner of the crystal shows a single phase of the spinel structure. Therefore, we removed the outer region of the seed rod, and the inner of crystal was only used for this study. However, the chemical composition analysis indicates the excess Fe



FIG. 6. The FZ equipment (a) before and (b) after crystal growth. The black powder attached on the surface of the quartz tube after the crystal growth.



FIG. 7. (a) SPD patterns for the black powder and the outer and inner regions of the crystal. The Bragg peaks of black powder can be indexed as a cubic system with the space group $Im\overline{3}m$. The asterisks indicate the Bragg peaks derived from the impurity of V_2O_3 . The inset shows the section of seed rods after crystal growth. (b) The ratio of Fe (α) and V (β) cations was determined by the chemical composition analysis. Note that PC and SC denote the polycrystalline and the single-crystal samples, respectively.

(5%-10%) even in the inner region of the growth crystal. This excess Fe in the single crystal may originate from the phase separation of V₂O₃ in the outer region of the crystal, as shown in Fig. 7(a). As a result, the off-stoichiometry in the single crystal is derived from both the evaporation of Fe and the precipitation of V₂O₃ during the crystal growth.

The chemical compositions of the single-crystal and polycrystalline samples were determined by energy dispersion x-ray spectroscopy by using Horiba EMAX ENERGY equipment. Figure 7(b) shows the ratio of Fe (α) and V (β) cations, where $\alpha + \beta = 3$, in the single crystal in inner region of growth crystal and the polycrystalline sample of FeV₂O₄ and of (Fe)[$V_{2-x}Fe_x$]O₄. The α values for the polycrystalline and single-crystal samples of FeV_2O_4 are 1.001 (8) and 1.100 (9), respectively. These results indicate a stoichiometric mixture for the polycrystalline sample of FeV₂O₄ and the existence of excess Fe in the single-crystal sample. The chemical composition of the single-crystal sample is similar to that of the polycrystalline sample of $Fe[V_{1,9}Fe_{0,1}]O_4$. This result is consistent with the results of the measurements of physical property. Hence, we regard the single crystal of FeV₂O₄ as $Fe_{1+\delta}V_{2-\delta}O_4$, and the δ is about 0.1. Note that we focused on the ratio of the transition metals in this study because it was difficult to quantitatively evaluate the oxygen atom.

APPENDIX B: PHYSICAL PROPERTIES IN $Fe_{1+\delta}V_{2-\delta}O_4$ SINGLE CRYSTAL

We have investigated the structural properties in the $Fe_{1+\delta}V_{2-\delta}O_4$ single crystal at low temperatures using the synchrotron diffraction technique. Figure 8 represents the synchrotron diffraction patterns around the 800 cubic Bragg peak. At 160 K, the 800 reflection is a single peak, and the peak is split into two peaks at 110 K and three peaks at 100 K. Furthermore, the three peaks are merged into two peaks at 60 K. For the $Fe_{1+\delta}V_{2-\delta}O_4$ single crystal, the temperature dependencies of the lattice constants, which are estimated from the Bragg peak positions, magnetization, and specific heat are shown in Figs. 8(b) and 8(c). Note that the experimental data of the temperature dependencies of the magnetization and the specific heat are from Ref. [17]. These results agree perfectly with recent studies using the single crystal [11,13]. On the other hand, below 40 K, we could not clearly observe a peak splitting caused by orthorhombic distortion. The reason is that the excess Fe in the single crystal suppresses the orthorhombic distortion at the lowest temperature. Indeed, the structural properties and phase transition temperatures for the single crystal are very similar to those for $Fe[V_{1.9}Fe_{0.1}]O_4$.

The excess Fe in the single crystal can affect the transitions at T_{N1} and T_{N2} . In practice, in the specific heat, the peak at T_{N2} in the polycrystalline sample is much sharper and shifts to higher temperature than that in the single crystal, as shown in Fig. 8(d). Additionally, the lattice constants around T_{N2} in the polycrystalline sample exhibit a sharper drop with decreasing temperature than those in the single crystal, as shown in Fig. 8(e). These differences of the behaviors between the polycrystalline sample and the single crystal can be explained by the excess Fe with nonorbital degrees of freedom at the B site in the spinel structure in the single crystal [34]. The phase transition at T_{N2} originates from the magnetic and orbital orders of V^{3+} ions at the *B* site, which play an important role in this transition; therefore, the excess Fe in the single crystal suppresses the long-range magnetic and the orbital orders at T_{N2} , leading to the difference in the behaviors of the specific heat between the polycrystalline and single-crystal samples. On the other hand, the peak of the specific heat at T_{N1} in the single crystal is slightly sharper than that in the polycrystalline sample. At present, the origin of this behavior is unclear. However, the transition temperature slightly increases. The increase in the transition temperature was also observed in a Fe-rich system, $Fe_{1+x}V_{2-x}O_4$. In this



FIG. 8. (a) Single-crystalline synchrotron diffraction patterns around the 800 cubic Bragg peak. (b) Temperature dependencies of lattice constants. (c) Temperature dependencies of the magnetization and the specific heat for $Fe_{1+\delta}V_{2-\delta}O_4$ single crystal [17]. Temperature dependencies of (d) the specific heat and (e) the lattice constant of the *a* axis for single-crystal (SC) and polycrystalline (PC) FeV₂O₄ samples.

viewpoint, the magnetic interaction between the *A* site and the *B* site in the spinel structure may be enhanced by the excess Fe with S = 5/2 [34]. In FeV₂O₄, the magnetic states are strongly related to the orbital states at T_{N1} ; therefore, the excess Fe in the single crystal affects the transitions related with orbital and spin degrees of freedom. Note that the change of temperature dependence of the magnetization at T_{N1} and T_{N2} in the Fe_{1+ δ}V_{2- δ}O₄ single crystal is much sharper than that in the polycrystalline sample. This behavior probably comes from the strong magnetic anisotropy in FeV₂O₄ [17]. In this study, the magnetic field was applied along the [001]_c direction of the crystal, where this direction is along the easy axis of the magnetization in FeV₂O₄ [14,17].

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