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Interdependent scaling of long-range oxygen and magnetic ordering in nonstoichiometric Nd$_2$NiO$_{4.10}$

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Hole doping in Nd$_2$NiO$_{4.00}$ can be achieved either by substituting the trivalent Nd atoms by bivalent alkaline-earth metals or by oxygen doping, yielding Nd$_2$NiO$_{4.10}$. While the alkaline-earth-metal atoms are statistically distributed on the rare-earth sites, the extra oxygen atoms in the interstitial lattice remain mobile down to ambient temperature and allow complex ordering scenarios depending on δ and T. Thereby the oxygen ordering, usually setting in far above room temperature, adds an additional degree of freedom on top of charge, spin, and orbital ordering, which appear at much lower temperatures. In this study, we investigated the interplay between oxygen and spin ordering for a low oxygen doping concentration, i.e., Nd$_2$NiO$_{4.10}$. Although the extra oxygen doping level remains rather modest with only 1 out of 20 possible interstitial tetrahedral sites occupied, we observed by single-crystal neutron diffraction the presence of a complex three-dimensional (3D) modulated structure related to oxygen ordering already at ambient, the modulation vectors being ±2/13±3/13*, ±3/13±2/13*, and ±1/5a±1/2c*, and satellite reflections up to fourth order. Temperature-dependent neutron-diffraction studies indicate the coexistence of oxygen and magnetic ordering below $T_N$ ≃ 48 K, the wave vector of the Ni sublattice being $k = (100)$. In addition, magnetic satellite reflections adapt exactly the same modulation vectors as found for the oxygen ordering, evidencing a unique coexistence of 3D modulated ordering for spin and oxygen ordering in Nd$_2$NiO$_{4.10}$. Temperature-dependent measurements of magnetic intensities suggest two magnetic phase transitions below 48 and 20 K, indicating two distinct onsets of magnetic ordering for the Ni and Nd sublattices, respectively.

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I. INTRODUCTION

Perovskite-type layered rare-earth Ruddlesden-Popper nickelates with the chemical formula Ln$_2$NiO$_4$ (Ln = La, Pr, Nd) are one of the many strongly correlated electronic systems which show interesting physical and electronic properties as a function of temperature [1–8]. These materials are complex because of the different active degrees of freedom such as lattice, spin, charge, and orbital ordering which interact in a competitive and synergetic way and open a wide field for applications. The crystal structures of these compounds can be described as a layered structure with alternate stacking of NiO$_2$ layers and rocksalt-type Ln$_2$O$_2$ bilayers along the long axis (c axis) of the unit cell (cf., Fig. 1) [9]. The undoped compounds are Mott insulators. Interesting physical properties like a metal-insulator transition appear when nickelates are doped with excess charge carriers. Holes can be incorporated either by partial substitution of Ln$^{3+}$ cations with divalent cations like Sr$^{2+}$, Ca$^{2+}$, and Ba$^{2+}$ or by doping with excess oxygen atoms. However, a metal-insulator transition occurs at a relatively large hole concentration ($n_h$) of 1 hole/formula unit [10]. In the insulating state, holes segregate inside the NiO$_2$ planes as an ordered array of parallel regions separating the antiferromagnetic (AFM) spin order as a domain boundary, known as the stripe or checkerboard charge ordered state [2,10–14]. The incorporated holes are in a diluted region for $n_h < 0.25$ and strong correlations among the holes appear for $0.25 < n_h < 1$ that consequently result in stripe or checkerboard charge ordered states, which are experimentally observed with neutron diffraction in Sr- and O-doped Ln$_2$NiO$_4$.

The substitution of Ln$^{3+}$ ions with smaller divalent cations is random and has a relatively weak effect on the lattice. In contrast, for oxygen-doped systems, excess oxygen atoms take interstitial lattice sites inside the Ln$_2$O$_2$ bilayer and introduce local disorder in the long-range tilting pattern of NiO$_6$ octahedra by systematically pushing adjacent apical oxygen atoms towards the nearest vacant interstitial sites [15–18]. Such apical oxygen displacements are strongly anharmonic in character and are important for the oxygen mobility close to room temperature, resembling a successively in- and exhaling behavior of the involved empty and occupied...
The presence of oxygen superstructure reflections appearing from the ordering of excess oxygen atoms has been observed down to 2 K with powder-diffraction studies. In addition, a commensurate magnetic structure has been proposed for Nd$_2$NiO$_{4+\delta}$ with propagation vector $k = (100)$ [18]. The commensurate magnetic order coexists with an additional magnetic phase which has been assumed to be incommensurate but significantly different from the stripe spin ordering [18].

In this work, we aim to understand this additional magnetic phase by employing neutron-diffraction studies on a Nd$_2$NiO$_{4+\delta}$ single crystal with similar oxygen nonstoichiometry of about $\delta \simeq 0.10$. In addition, we search for possible superstructure reflections appearing from the ordering of excess oxygen atoms, spin, and charge in the LTT phase of Nd$_2$NiO$_{4.10}$.

II. EXPERIMENTAL METHODS

A. Sample synthesis

A large single crystal was grown at Institute Charles Gerhardt Montpellier (ICGM) using a two-mirror optical floating zone furnace at high temperature in oxygen atmosphere as described elsewhere [28,29]. The as-grown crystal has an overall oxygen nonstoichiometry ($\delta$) in the range 0.23–0.25. Consequently, further heat treatment was performed to obtain a single crystal with reduced excess oxygen content $\delta$. For that, a small section (~200 mg) was cut and annealed in a dynamic vacuum at 1073 K for 6 h. The details are reported in Ref. [18]. The overall oxygen stoichiometry in the single crystal was determined to be 4.097(10) with the least-squares refinement of the room-temperature single-crystal neutron-diffraction data [18]. A powder sample was separately synthesized and reduced under identical conditions used for the single crystal. The overall oxygen stoichiometry in the powder sample was determined using thermogravimetric analysis (TGA) at high temperature as well as with neutron powder-diffraction data at 300 K to be 0.11(2), and thus very similar to the one used for the single crystal using neutron-diffraction experiments. Both the single-crystal and powder samples are thus identical to those reported in Ref. [18]. In between different measurements, the samples were stored inside a glove box with Ar atmosphere to minimize the oxidizing effects.

B. X-ray and neutron-diffraction measurements

Synchrotron x-ray powder-diffraction (XRPD) measurements were performed at different temperatures in between 5 and 300 K at the Material Sciences (MS) beamline X04SA at the Swiss Light Source (SLS) of the Paul Scherrer Institute (PSI), Switzerland. The wavelength ($\lambda = 0.6208(1)$ Å) and instrumental resolution parameters were determined from a
standard LaB₆ powder (NIST), measured under identical experimental conditions. Samples were placed into thin glass capillaries of 0.3 mm diameter and continuously rotated to reduce preferred orientation during the data collection. Neutron powder-diffraction (NPD) studies were carried out in the temperature range 2–300 K using the cold neutron powder diffractometer DMC [30,31] at SINQ, PSI, in Switzerland. Both x-ray and neutron-diffraction data were analyzed with the FULLPROF Suite [32]. DMC was further used to perform the reciprocal plane mappings on the single crystal at variable temperatures between 2 and 300 K. A preoriented single-crystal sample was mounted on top of a thin aluminum pin and the reciprocal (hk0) and (h0l) planes were mapped out by rotating the single crystal around the respective zone axis. The reciprocal maps are particularly useful to locate weak superstructure peaks or diffuse scattering in the reciprocal space up to a momentum transfer (Q) of about 4.15 Å⁻¹. Moreover, λ/2 contamination could be excluded at this wavelength on DMC. The reciprocal planes were reconstructed from the raw data using the TVTUEB program [33]. Integrated intensities of selected structural and magnetic Bragg reflections were measured using the hot-neutron four-circle diffractometer HEIDI (equipped with a point detector) at the Heinz Maier-Leibnitz Zentrum (MLZ) at the FRM-II reactor in Garching, Germany [34]. For these measurements, transverse, i.e., ω, scans were performed with the crystal oriented on the (hk0) scattering plane with the c axis vertically aligned. A short wavelength of 0.793(1) Å was used for the data collection up to a momentum transfer of about 11.30 Å⁻¹. The data sets were collected at 300, 55, and 2 K. Structural least-squares refinements of the integrated and Lorentz corrected structure factors of P-type Bragg reflections were performed with the JANA2006 software package [35]. The crystal structure is visualized using VESTA freeware [36].

III. RESULTS AND DISCUSSIONS

A. Average crystal structure

The LTT crystal structure of the moderately oxygen-doped Nd₂NiO₄₋ₓδ compound with δ ~ 0.11 has been previously revealed with powder and single-crystal neutron-diffraction studies in the temperature range 2–300 K [18]. The unit cell of the LTT phase is shown in Fig. 1(a) obtained from room-temperature neutron-diffraction data. In general, the Nd₂O₂ layer is in the extensional strain whereas the NiO₂ layer undergoes a compressional strain. The structural strain is released through structural distortions that involve a systematic tilting of NiO₆ octahedra about a fixed crystallographic axis. In the LTT phase, NiO₆ octahedra are canted around the [1T₀] direction in the basal plane such that in-plane nearest octahedra are contrarotated. The rotation axis alternates between the [110] and [1T₀] directions in the neighboring NiO₂ planes along the c axis. Such an octahedral tilting gives rise to extra P-type superstructure reflections compared to the fully disordered high-temperature tetragonal (HTT) phase with space group F4/mmm. Our neutron-diffraction measurements give evidence of sharp P-type superstructure reflections already at room temperature, indicating the LTT phase. The ω-scan profile of such P-type structural Bragg peaks measured at room temperature is presented in Fig. 2(a) for the (104) reflection. The specific octahedral tilting in the LTT phase allows favorable openings for the tetrahedral 4b (3/4, 1/4, 1/4) and 4e (1/4, 1/4, z ~ 1/4) interstitial sites as shown in Fig. 1(b). The refinement of the integrated intensity data of the main structural Bragg peaks at room temperature demonstrates that the excess oxygen atoms selectively occupy the 4b interstitial sites, in accordance with previously reported results [17,18]. It is worth noting that the LTT structure has been reported for the stoichiometric Nd₂NiO₄₀₀ compound below 135 K [37]. Here, the LTT structure becomes stable already at ambient temperature related to additional oxygen doping. We note that only 1 of 20 possible tetrahedral sites is occupied in Nd₂NiO₄₁₀; i.e., 0.4 oxygen atoms are present on average per unit cell.

B. Commensurate magnetic structure

The dc magnetic susceptibility data, recorded for the Nd₂NiO₄₁₀ single crystal with a field of 1 T applied parallel to an arbitrary crystallographic axis in zero-field-cooled (ZFC) configuration, is presented in Fig. S1 of the Supplemental Material [38]. An antiferromagnetic transition at Tₐ ≃ 48 K can be clearly discerned from the magnetic susceptibility plot. The magnetic structure of Nd₂NiO₄₁₀ below Tₐ can be described with a commensurate propagation vector k = (100) as previously revealed by NPD data. Consequently, the intensities of all Bragg peaks significantly increase below Tₐ due to the magnetic contribution, and commensurate magnetic peaks become evident with decreasing temperature from 55 to 2 K. The ω-scan profile of a pure magnetic Bragg peak such as the (101) is presented in Fig. 2(b). The temperature evolution of integrated intensities of the (104) and (101) reflections are shown.
in Figs. 2(c) and 2(d), respectively. Two magnetic transitions can clearly be distinguished from the temperature dependence. The first starts below \( T \approx 48 \text{ K} \) in accordance with the dc magnetic susceptibility data and indicates the onset of antiferromagnetic order of the Ni sublattice. The one indicated by a change of the slope below \( T \approx 20 \text{ K} \), while intensities of all the Bragg reflections are rapidly increasing, is associated to the onset of magnetic polarization of the Nd sublattice. The commensurate magnetic structure of the average structure corresponding to the LTT phase has been solved by least-squares refinement from a set of integrated intensities recorded at 2 K. In this magnetic structure, Ni and Nd spins are coupled antiferromagnetically and oriented parallel to the crystallographic [110] or [1
d] axis was reported for \( \text{La}_2\text{NiO}_4 \) [18].

The total magnetic moments for the Ni and Nd sites were in good agreement with previous results [18]. These magnetic superstructure reflections can be rationalized with the same modulation vectors that define the excess oxygen ordering reflections at 300 K. For moderately oxygen-doped \( \text{Nd}_2\text{NiO}_4 \), a commensurate magnetic order coexists along with long-range oxygen ordering, which was not previously observed in homologous \( \text{La}_2\text{NiO}_{4+x} \) compounds. Furthermore, extra commensurate reflections such as the (210) and the (105), which are symmetry forbidden in \( P4_2/mnc \), were evidenced already at ambient temperature, suggesting that the true symmetry of \( \text{Nd}_2\text{NiO}_{4.10} \) is possibly lower while inconsistent with the structural model proposed for homologous \( \text{La}_2\text{NiO}_{4.105} \). In this context, it is worth noting that the commensurate charge-ordering reflections in hole-doped \( \text{La}_{2-x}\text{Sr}_x\text{NiO}_4 \) are found to emerge in such commensurate positions [13]. However, we do not expect stripe- or checkerboard-type charge orders to occur in our sample as the required hole concentration for such correlation to appear is considerably higher than the incorporated hole concentration in our sample, only about 0.2 per formula unit. Furthermore, in the presence of checkerboard-type charge ordering, magnetic Bragg peaks at \((h = 0.5, 0, 0)\) positions would also be expected, which are clearly not observed here [13]. Therefore, the commensurate reflections are most likely due to a lowering of the symmetry from \( P4_2/mnc \).

We have attempted to index all the superstructure reflections appearing in the 2 K maps. This is a complicated task especially in the case of the presence of a large number of satellites, i.e., higher-order harmonics. It is found that all the superstructure reflections could be indexed assuming modulation vectors of the type \( q_{\nu} = \alpha \alpha^* + \beta b^* + \gamma e^* \) added to the position of each Bragg reflection \( G = h\alpha + k\beta + l\epsilon + m\alpha \), with \( m \) being an integer number representing the order of corresponding superstructure reflections. The underlying modulation vectors, which describe both the excess oxygen ordering and magnetic ordering in the \((hk0)\) and \((h0l)\) planes, are found to be \( \pm 2/3 a^* \pm \pm \pm 3/5 b^* \pm \pm\pm \pm 3/13 a^* \pm \pm \pm \pm 3/2 b^* \), and \( \pm 1/5 a^* \pm \pm 1/2 c^* \). In addition, structural Bragg peaks from epitaxially intergrown NiO as reported in Ref. [25] were also observed at all the investigated temperatures. An example of the manually indexed experimental and idealized \((hk0)\) plane is presented in Figs. S2 and Fig. S3 of the Supplemental Material, respectively [38]. The simulated reconstructions for possible superstructure reflection positions were undertaken using these modulation vectors. However, it is essential to consider all the symmetry-related directions of the modulation vectors to satisfactorily describe all the satellite positions in the \((hk0)\) and \((h0l)\) plane maps. This is clearly evident from the \((hk0)\) plane map in which all the satellite positions at 2 K are excellently represented by all eight directions of the modulation vectors given by \( \pm 2/13 a^* \pm \pm \pm \pm 3/13 b^* \) and \( \pm 1/13 a^* \pm \pm \pm 2/13 c^* \). In a similar way, the \((h0l)\) plane can be described by the four directions of the modulation vectors which are characterized by \( \pm 1/5 a^* \pm \pm 1/2 c^* \). It is furthermore necessary to consider up to fourth-order harmonics for a successful indexation of all the satellites appearing in both reciprocal planes at 2 K. The simulated reconstructions, which excellently represent the experimental maps, are shown in Figs. S4 and S5 of the Supplemental Material [38]. The
FIG. 3. Reconstructed single-crystal neutron-diffraction plane maps of Nd$_2$NiO$_4$-10. Excerpts of (a–c) ($h0l$) and (d–f) ($hk0$) reciprocal planes measured at 300, 55, and 2 K. The dashed blue circles show the reflections that emerged from the NiO impurity phase while the squares and solid circles represent characteristic superstructure reflections appearing from the ordering of excess oxygen atoms and spins in Nd$_2$NiO$_4$-10, respectively. Rectangular boxes in orange in the 2 K maps display the sections which are magnified in Fig. 4. The measurements were carried out on DMC at SINQ, PSI, with $\lambda$ = 2.4586(3) Å.

magnified sections of the experimental and simulated zones of the ($h0l$) and ($hk0$) planes are displayed in Figs. 4(a)–4(d). It is found that the superstructure peaks in the ($h0l$) plane are symmetrically positioned in such a way that they form a hexagon around a commensurate magnetic peak of ($h0l$) type with $l$ = odd as shown in Fig. 4(b). An important finding of these simulations is that a substantial overlap of intensities from different harmonics associated to different Bragg peaks

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FIG. 4. Magnified zones of the (a) experimentally measured \((h0l)\) plane presented together with the (b) simulated zone. Magnified sections of the (c) experimentally measured \((hk0)\) plane presented along with the (d) simulated zone. Integer numbers represent different orders of the satellites corresponding to a Bragg peak. \(Q\) scans through the superstructure peaks observed in the \((hk0)\) and \((h0l)\) plane maps of \(Nd_2NiO_4\). (e) The \(Q\) scan is starting from \((102)\) along the blue line up to the fourth-order satellite as shown in (a); (f) the cut is starting from \((110)\) along the green line up to the fourth-order satellite as shown in (c). A constant \(y\) offset is used to plot the 55 and 2 K data.

occurs in the reciprocal plane. For example, in the \((h0l)\) plane [cf. Fig. 4(b)], the first- and second-order satellites related to the \(1/5a^*+1/2c^*\) vector starting from the \((101)\) Bragg peak are identical with the first- and second-order satellites corresponding to the \(1/5a^*-1/2c^*\) vector starting from \((102)\) and \((100)\) Bragg peaks, respectively. However, in the \((hk0)\) plane [cf. Fig. 4(d)], the satellites are sufficiently separated which allows to determine their structure factor satellites separately. Figures 4(d) and 4(e) represent specific \(Q\) scans through the superstructure reflections in the reciprocal maps obtained at different temperatures. It is evident that the intensities of some superstructure reflections like the first- and second-order harmonics related to the \((102)\) and all the harmonics associated to \((110)\) are strongly enhanced below \(T_N\) owing to strong magnetic contributions. Furthermore, the modulation vectors are invariant in the temperature range 2–300 K; i.e., the same modulation vectors satisfactorily define the excess oxygen ordering peaks at room temperature. Moreover, no significant change in the full widths at half maximum (FWHM) of the oxygen superstructure reflections is detected as a function of temperature.

The excerpts of XRPD and NPD data recorded at 5 and 2 K are shown in Figs. 5(a) and 5(b), respectively. We note that superstructure reflections are observed in the NPD data only below \(T_N\) as shown in Fig. S6 of the Supplemental Material [38]. In strong contrast, superstructure reflections are present in the XRPD data already at room temperature and their positions remain unchanged down to 5 K. The satellites in the XRPD data are very weak owing to the small x-ray form factor of oxygen atoms but are clearly distinguishable from the background. The integrated intensities of two characteristic superstructure reflections observed in the NPD and XRPD data are plotted in the insets of Figs. 5(a) and 5(b). It can be seen that their integrated intensities in the NPD data are strongly enhanced with decreasing temperature down to 2 K while in the XRPD data they remain invariant in the temperature range 2–300 K. Therefore, the superlattice peaks in the XRPD data are essentially due to the ordering of excess oxygen atoms whereas those observed in the NPD data below \(T_N\) can be attributed to the magnetic ordering. Even though neutrons are quite sensitive to light oxygen atoms, the superstructure reflections originating from the excess oxygen ordering appear in the reciprocal plane maps only at higher \(Q\) but are not observed with NPD data probably because of their superimposition with the main Bragg reflections (see Fig. S7 [38]). Finally, we have performed Le Bail fits of the XRPD and the NPD data in order to index the superstructure reflections using the modulation vectors which are obtained from the plane maps. All the superstructure peaks observed by x-ray and neutron powder diffraction can be indexed satisfactorily by these modulation vectors as shown in Figs. 5(a) and 5(b). The fully refined NPD pattern is presented in Fig. S7 of the Supplemental Material [38]. This result indicates that the modulation vectors obtained from the plane mappings are equally valid for the description of satellites observed in the powder data and further confirms that the magnetic order is correlated to the excess oxygen order in moderately oxygen-doped \(Nd_2NiO_{4+\delta}\) with \(\delta \simeq 0.10\).
IV. CONCLUSIONS

Large-scale oxygen and magnetic ordering have been evidenced by single-crystal neutron-diffraction studies on Nd$_2$NiO$_{4.10}$ to reveal identical modulation vectors resulting in a 3D modulated structure. The commensurate antiferromagnetic order in Nd$_2$NiO$_{4.10}$ sets in at $T_N \simeq 48$ K with a propagation vector $\mathbf{k} = (100)$. The most peculiar point here is that the magnetic ordering modulation vectors are no longer independent and exactly follow the modulation vectors of the oxygen ordering. This is evidenced unambiguously from the selected planes explored with single-crystal neutron diffraction and also from the refinement of powder neutron-diffraction data. The magnetic order thus coexists below $T_N$ with the 3D ordering of excess oxygen atoms in Nd$_2$NiO$_{4.10}$, which is atypical so far, and not previously encountered for the homologous nickelates. Nd$_2$NiO$_{4.10}$ thus seems to present a special case in terms of structural and magnetic ordering phenomena, compared to other nickelates and especially to the La$_2$NiO$_{4.10}$. We note that no evidence for charge ordering has been identified in this study. Although the oxygen stoichiometry remains quite diluted with only 1 out of 20 possible tetrahedral sites occupied (i.e., 0.4 oxygen atoms on average per $P$-type unit cell), a complex 3D ordering gets established. This clearly indicates the potential of the interstitial oxygen atoms to enable inter- and intralayer coupling for structural and magnetic ordering. As a phenomenon, this becomes important with respect to the recently evidenced sub-mesoscale oxygen ordering in oxygen-rich Pr$_2$NiO$_{4+\delta}$ ($\delta \simeq 0.25$), revealing translational periodicities of almost 100 Å. In this context, the mediation achieved between magnetic and oxygen ordering thus yields a promising concept to establish long-range magnetic ordering, essentially based on long-range oxygen ordering.

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