

Cite this: *Mater. Adv.*, 2023,  
4, 651

# Large-scale oxygen order phase transitions and fast ordering kinetics at moderate temperatures in $\text{Nd}_2\text{NiO}_{4+\delta}$ electrodes†

Sumit Ranjan Maity,<sup>\*ab</sup> Monica Ceretti,<sup>id c</sup> Ruben De Barros,<sup>c</sup> Lukas Keller,<sup>a</sup>  
Jürg Schefer,<sup>a</sup> Antonio Cervellino,<sup>d</sup> J. Alberto Rodríguez Velamazán<sup>id e</sup> and  
Werner Paulus<sup>id \*c</sup>

Non-stoichiometric 214-nickelates with Ruddlesden–Popper (RP) type frameworks emerged as potential candidates for mixed electronic/ionic conductors in the intermediate temperature range. In this work we investigated structural aspects of the oxygen ion mobility diffusion mechanisms in non-stoichiometric  $\text{Nd}_2\text{NiO}_{4+\delta}$  nickelates by X-ray (laboratory and synchrotron) as well by neutron diffraction. Temperature dependent synchrotron powder diffraction revealed a phase diagram of unprecedented complexity, involving a series of highly organized, 3D modulated phases related to oxygen ordering below 800 K. All phase transitions simply translational periodicities exceeding 100 Å, and are found to be of 1<sup>st</sup> order, together with fast ordering kinetics. These surprising structural correlations, induced by the presence of interstitial oxygen atoms, suggest a collective phason-like oxygen diffusion mechanism together with dynamical contributions from the aperiodical lattice creating shallow diffusion pathways down to room temperature.

Received 19th July 2022,  
Accepted 12th December 2022

DOI: 10.1039/d2ma00833e

rsc.li/materials-advances

## Introduction

Tailoring new materials with improved oxygen mobility especially at moderate temperatures is of great scientific interest owing to their potential applications in electrochemical energy conversion and energy storage devices, including intermediate temperature solid oxide fuel cells (IT-SOFCs), oxygen membranes, and solid oxide electrolysis cell devices.<sup>1–5</sup> In this regard, non-stoichiometric 214-nickelates ( $\text{Ln}_2\text{NiO}_{4+\delta}$ ; Ln = La, Pr or Nd) of the Ruddlesden–Popper (RP) series with layered  $\text{K}_2\text{NiF}_4$  structure appear as one of the most promising candidates due to their electronic complexity and enhanced oxygen mobility down to ambient temperature.<sup>5–15</sup> The crystal structure of these compounds, as shown in Fig. 1(a), can be described as a layered structure with alternate stacking of  $\text{NiO}_2$  layers and rock salt type  $\text{Ln}_2\text{O}_2$  bilayers along the stacking-axis (*c*-axis) of the

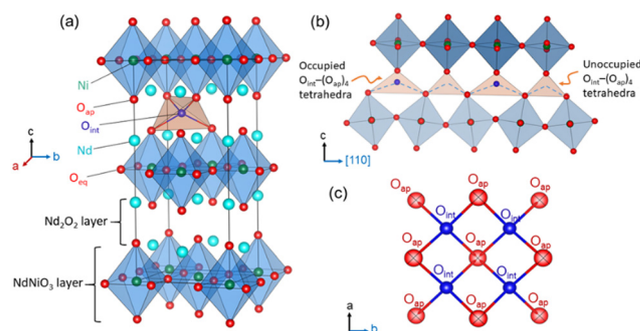


Fig. 1 (a)  $\text{K}_2\text{NiF}_4$  type layered crystal structure of  $\text{Nd}_2\text{NiO}_{4+\delta}$ , consisting of alternating perovskite and rock-salt layers along the stacking axis. Only one out of possible eight tetrahedral interstitial sites are occupied for a  $\delta$  value of  $\approx 0.23$ . (b) The tilting scheme of  $\text{NiO}_6$  octahedra induced by oxygen ordering in the interstitial sites, resulting in alternative occupied (expanded) and unoccupied (contracted)  $\text{O}_{\text{int}}-(\text{O}_{\text{ap}})_4$  tetrahedra along the diffusion pathway, promoting anisotropic oxygen diffusion along [110], involving  $\text{O}_{\text{int}}-\text{O}_{\text{ap}}-\text{O}_{\text{int}}$  tetrahedral chains as shown with blue dashed lines. (c) Sketch of the  $\text{O}_{\text{ap}}$  and  $\text{O}_{\text{int}}$  atoms within the rock salt layer projected along [001].

<sup>a</sup> Laboratory for Neutron Scattering and Imaging, Paul Scherrer Institut, Villigen, CH-5232, Switzerland. E-mail: sumit050491@gmail.com

<sup>b</sup> University of Geneva, Department of Quantum Matter Physics (DQMP) 24, Quai Ernest Ansermet, CH-1211, Genève 4, Switzerland

<sup>c</sup> ICGM, Univ. Montpellier, CNRS, ENSCM, Montpellier, France. E-mail: werner.paulus@umontpellier.fr

<sup>d</sup> Swiss Light Source, Paul Scherrer Institut, Villigen, CH-5232, Switzerland

<sup>e</sup> Institut Laue-Langevin, 71 Avenue des Martyrs, F-38042, Grenoble, France

† Electronic supplementary information (ESI) available: Laboratory X-ray and synchrotron powder diffraction patterns (Fig. S1–S10); twin domains scheme (Fig. S2). See DOI: <https://doi.org/10.1039/d2ma00833e>

tetragonal unit cell.<sup>16</sup> Oxygen diffusion in these materials has been shown to be highly anisotropic, following a push–pull diffusion mechanism between apical ( $\text{O}_{\text{ap}}$ ) and vacant interstitial oxygen sites ( $\text{O}_{\text{int}}$ ) inside the rock salt layer, promoted by large displacements of the  $\text{O}_{\text{ap}}$  atoms.<sup>17–21</sup>



Our present work focuses on the  $\text{Nd}_2\text{NiO}_{4+\delta}$  (NNO) system that shows a complex structural and electronic phase diagram as a function of temperature and the excess oxygen content  $\delta$ .<sup>22–24</sup> As for other Ruddlesden–Popper type oxides, structural strain is present due to a large lattice mismatch between the  $\text{NiO}_2$  and  $\text{Nd}_2\text{O}_2$  rock salt layers. While the high-temperature tetragonal (HTT) structure with space group  $F4/mmm$  yields a sufficiently good matching between these layers,<sup>25</sup> the strain is released by the cooperative tilt around [100] of  $\text{NiO}_6$  octahedra, lowering the symmetry from tetragonal to orthorhombic. It consequently creates a lattice instability for the apical oxygen atoms which are significantly shifted from their equilibrium positions, left on a shallow potential.<sup>19,26</sup>

The incorporation of excess oxygen ( $\text{O}_{\text{int}}$ ) atoms into the vacant tetrahedral interstitial sites inside the  $\text{Nd}_2\text{O}_2$  bilayers leads to oxygen hyper-stoichiometric NNO with  $0 \leq \delta \leq 0.25$ <sup>27</sup> (see Fig. 1(a)). This partially relieves the structural strain as the oxidation of  $\text{Ni}^{2+}$  towards  $\text{Ni}^{3+}$  results in shorter Ni–O distances.<sup>16</sup> The strain is also reduced due to the presence of interstitial oxygen atoms ( $\text{O}_{\text{int}}$ ) causing important shifts of all adjacent oxygen apical atoms ( $\text{O}_{\text{ap}}$ ) towards [110], forming symmetrically increased  $\text{O}_{\text{int}}(\text{O}_{\text{ap}})_4$  tetrahedra. The as shifted  $\text{O}_{\text{ap}}$  atoms thereby directly point towards adjacent vacant  $\text{O}_{\text{int}}$  lattice sites, thus promoting the push–pull diffusion mechanism between  $\text{O}_{\text{ap}}$  and  $\text{O}_{\text{int}}$  atoms. The excess oxygen atoms can be reversibly inserted into the tetrahedral lattice sites at room temperature by electrochemical red/ox reactions in an aqueous alkaline solution, clearly demonstrating high oxygen mobility to proceed down to room temperature.<sup>24,27</sup> The related RT phase diagram of  $\text{Nd}_2\text{NiO}_{4+\delta}$  shows in addition to two orthorhombic line phases with an extra oxygen stoichiometry of  $\delta = 0.23$  and  $0.00$ , a non-stoichiometric phase with  $0.08 \leq \delta \leq 0.12$ . It thus involves two miscibility gaps outside this stoichiometry range, *i.e.*, for  $0 \leq \delta \leq 0.08$  and  $0.12 \leq \delta \leq 0.23$  corresponding to the fully reduced and oxidized phases.

With increasing oxygen uptake the local distortions of the  $\text{O}_{\text{int}}(\text{O}_{\text{ap}})_4$  tetrahedra get organized on a larger length scale, as evidenced by neutron and synchrotron diffraction experiments.<sup>7,19,28–31</sup> Ordering schemes of unprecedented structural complexity have been reported for the homologous  $\text{Pr}_2\text{NiO}_{4+\delta}$ , where sub-mesoscopic ordering is observed for  $\delta = 0.25$  attaining translational periodicities of *ca.* 100 Å.<sup>31</sup> Thereby puzzling correlations are established between oxygen, charge, and spin ordering. In this context, we could recently also unveil a complex 3D modulated structure related to oxygen ordering at RT for  $\text{Nd}_2\text{NiO}_{4.1}$ , the related modulation vectors being  $\mathbf{Q}_n = \pm 2/13\mathbf{a}^* \pm 3/13\mathbf{b}^*$  and  $\mathbf{Q}_n = \pm 1/5\mathbf{a}^* \pm 1/2\mathbf{c}^*$ . Below  $T_N \cong 50$  K, antiferromagnetic ordering is observed, with magnetic modulation vectors identical to those observed for the oxygen ordering.<sup>7</sup> This clearly suggests that the magnetic ordering is directly influenced by the oxygen ordering. It underlines that strong correlations exist between structural and electronic ordering schemes, which condition in the case of  $\text{Nd}_2\text{NiO}_{4.10}$  the degree of freedom for electronic, *i.e.*, spin ordering, as oxygen ordering sets in at much higher temperatures compared to magnetic ordering. The simple presence of  $\text{O}_{\text{int}}$  becomes

then decisive to create strong structural correlations, which have also been demonstrated for  $\text{Pr}_2\text{NiO}_{4.25}$  to persist upon heating, attaining temperatures of almost 700 K. Even above this temperature, the correlations between  $\text{O}_{\text{int}}$  and  $\text{O}_{\text{ap}}$  atoms partially continue to exist, as indicated by strong diffuse scattering contributions, and involved correlation lengths of about 25 Å.<sup>31</sup> While the real structure of modulated, oxygen doped Ruddlesden–Popper type oxides remains hitherto unexplored in terms of atomic positions and octahedra tilting *etc.*, no further coupling mechanism between lattice and electronic ordering has been reported so far in detail. Our understanding for such lattice–electronic coupling schemes is thus limited to a more phenomenological description, which is already interesting by itself.

While for oxygen-doped Ruddlesden–Popper type oxides, structural ordering goes along with electronic ordering schemes towards lower  $T$ , only little information is known on the evolution of these complex ordering schemes and respective stabilities with increasing  $T$ . Thus, our intention is to further explore the stability range of oxygen ordering but also the emergence of possible new ordering schemes as a function of temperature, together with their dependence on the oxygen stoichiometry. Such studies are highly relevant for a better understanding of oxygen diffusion mechanisms as they allow to discriminate up to which extent such mechanisms may be described as single particle events or based on the long-range oxygen ordering, need to be understood as a correlated process similar to phonons.

To further deepen structure and dynamics of oxygen doped Ruddlesden Popper phases, we report here on order/order phase transitions evidenced in  $\text{Nd}_2\text{NiO}_{4+\delta}$ , for which a complex 3D incommensurate oxygen ordering has already been reported to be present at ambient temperature.<sup>32</sup>

Using synchrotron X-ray powder diffraction (SXRPD), we were able to evidence a series of 1<sup>st</sup> order phase transitions related to oxygen ordering below the phase transition into the tetragonal high- $T$  parent phase at 800 K.

In order to facilitate the complex structural behaviour presented in more detail below, we briefly anticipate our findings, evidencing a series of several order/order phase transitions of  $\text{Nd}_2\text{NiO}_{4+\delta}$  in the moderate  $T$ -range below 800 K, which are of extreme complexity and well-defined phase definition, all showing 3D modulated structures. Surprisingly, ordering kinetics are found to proceed instantly, despite the translational periodicities observed exceeding 100 Å for all involved phases. It thus implies that oxygen is not only exceptionally mobile, but also shows the ability for large-scale correlations, indicative to promote collective, phonon-assisted diffusion behaviour as a result of lattice instabilities following phonon softening.<sup>33</sup>

## Experimental methods

### Sample synthesis

$\text{Nd}_2\text{NiO}_{4+\delta}$  single crystal growth was done at the Institut Charles Gerhardt Montpellier (ICGM) using a two-mirror optical floating zone furnace at high temperature in an oxygen atmosphere as described elsewhere.<sup>32</sup> The crystal was further



annealed in air at 670 K for 7 days to obtain a uniform oxygen non-stoichiometry  $\delta$ . A representative powder sample was prepared by crushing a small section of the single crystal. The single crystal and powder samples were kept at ambient for more than six months prior to all the measurements.

### Thermogravimetric measurements

The average oxygen content  $\delta$  in the powder sample was determined by thermogravimetric (TG) measurements in 5%-H<sub>2</sub>/95%-He gas atmosphere in the temperature range of 300–1273 K. The measurement was carried out at the Laboratory for Multiscale Materials Experiments (LDM) of the Paul Scherrer Institute (PSI) using a Netzsch STA 449 C analyzer. The powder sample was fully decomposed into Nd<sub>2</sub>O<sub>3</sub> and metallic Ni at high temperatures. The  $\delta$  content in the powder sample was determined to be 0.23(2) from the TG weight loss curve as displayed in Fig. 2(a), in agreement with the previously reported result.<sup>32</sup> Further TG measurements on the powder sample of Nd<sub>2</sub>NiO<sub>4.23</sub> were carried out in 33%O<sub>2</sub>/67%Ar atmosphere in the temperature range of 300–1273 K with a heating/cooling rate of 5 K min<sup>-1</sup> using PerkinElmer TG8000 at ICGM (weight accuracy of 0.1  $\mu$ g). A baseline correction with an empty sample holder was applied to all the measurements.

### X-ray diffraction measurements

Temperature-dependent laboratory X-ray powder diffraction (XRPD) measurements were carried out at the Bioenergy and

Catalysis Laboratory (LBK) of PSI using a Bruker D8 advanced diffractometer using CuK<sub>α1,2</sub> radiation in the temperature range of 300–1273 K in air atmosphere. The XRPD patterns were collected every 50 K during heating while a temperature step of 100 K was used for cooling to check for reversibility. Synchrotron X-ray powder diffraction (SXRPD) measurements were performed at different temperatures between RT–873 K at the Material Sciences (MS) beamline X04SA at the Swiss Light Source (SLS) of the Paul Scherrer Institute, Switzerland.<sup>34</sup> The wavelength and instrumental resolution parameters were determined from a standard LaB6 powder (NIST), measured under identical experimental conditions. Samples were filled into thin quartz capillaries of 0.3 mm diameter and continuously rotated to reduce the effect of preferred orientation during data collection. The pattern at each temperature was collected roughly for two minutes. A hot air blower was used for the high-temperature SXRPD measurements. SXRPD patterns in the temperature range of 853 K < *T* < 723 K during cooling could not be measured due to a detector problem. All powder diffraction patterns were analyzed using the FullProf suite program.<sup>35</sup> The crystal structure is visualized using the VESTA freeware.<sup>36</sup> Single crystal X-ray diffraction studies at room temperature (298 K) were carried out at the ICGM using an STOE-StadiVari (MoK<sub>α</sub> Xenocs Microfocus tube) diffractometer system with a Dectris 200K Pilatus pixel-detector.

### Neutron diffraction measurements

Single crystal neutron diffraction measurements at 20 K and 300 K were performed using ZEBRA/TriCS single crystal diffractometer (equipped with a point detector) at SINQ, PSI.<sup>37</sup> An incident wavelength of  $\lambda = 1.178(1)$  Å was used to access a reciprocal space up to  $Q = 4\pi \sin \theta / \lambda \approx 9.5$  Å<sup>-1</sup>. The high-temperature measurements at 673 K and 1073 K were carried out on D9 (equipped with a 2D detector) at ILL (France) using a hot air blower.<sup>38</sup> A monochromatic beam of wavelength  $\lambda = 0.837(1)$  Å was produced using the (220) plane of a Cu crystal an Er absorption filter in the transmission geometry to access a large reciprocal space up to  $Q \approx 11.3$  Å<sup>-1</sup>. Integrated intensities of structural Bragg peaks were collected with standard transverse scans ( $\omega$ -scans) with the *c*-axis of the crystal aligned vertically to the scattering plane of the diffractometer. Least-square refinements of the Lorentz corrected structure factors were carried out using the JANA2006 program.<sup>39</sup>

Nuclear densities were reconstructed in real space through the Maximum Entropy Method (MEM) *via* Dynomia<sup>40</sup> and nuclear density distributions were visualized by using the VESTA program.

## Results

### The room temperature phase and oxygen stoichiometry

Before discussing the structural changes in detail, we anticipate a brief discussion on the *O*-stoichiometry of as-synthesized Nd<sub>2</sub>NiO<sub>4+ $\delta$</sub>  samples (NNO), as this plays an important role for its modulation, and it is also important for the choice of

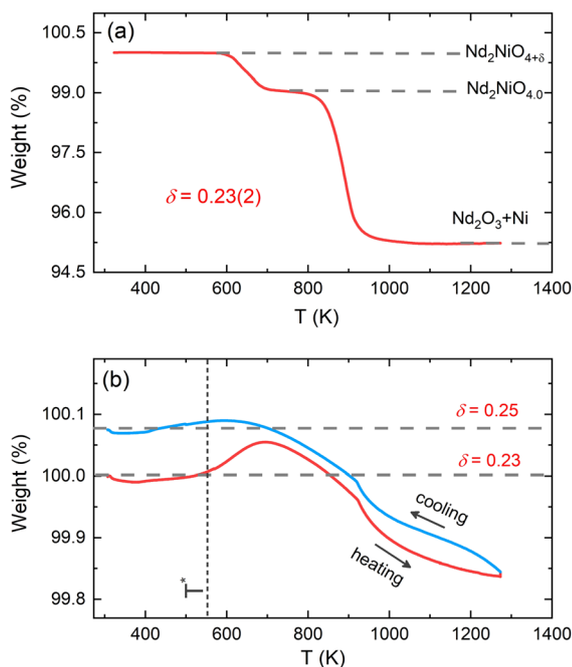


Fig. 2 (a) TG weight loss curve of the crushed powder sample of Nd<sub>2</sub>NiO<sub>4+ $\delta$</sub>  measured in 5%-H<sub>2</sub>/95%-He gas atmosphere with a heating rate of 5 K min<sup>-1</sup>. The average excess oxygen content  $\delta$  has been determined to be  $\approx 0.23(2)$ . (b) The temperature evolution of sample weight in 33%O<sub>2</sub>/67%Ar atmosphere. Vertical dashed lines denote the temperature  $T^* \approx 550$  K where the modulation vector of the excess oxygen ordering changes during cooling.



material for the  $T$ -dependent SXRPD studies. The situation of NNO is pretty much similar to its homologue  $\text{Pr}_2\text{NiO}_{4+\delta}$  (PNO), for which a topotactic oxygen release of as-grown single crystals has been discussed, changing its oxygen stoichiometry from  $\delta = 0.25$  to  $\delta = 0.225$  within several months, when stored under ambient atmosphere.<sup>31</sup> The reason for such an unusual behaviour is that the extra oxygen uptake is stabilized at high temperatures as a consequence of the strain release discussed above, while at room temperature the structure slowly relaxes towards another oxygen ordering scheme, involving a small but significant change in the stoichiometry as well as the modulation vector. The change in the  $O$ -stoichiometry also goes along with minor but significant changes in the lattice parameters.<sup>31</sup> It is therefore essential to associate to any crystallographic characterization of NNO samples their thermal history. While the relaxation time for oxygen release may require more than a year on large single crystals, 50  $\mu\text{m}$  size crystals need approximately 2–3 months to relax towards their ground state in terms of oxygen stoichiometry and related modulation vectors. For this reason, the powder samples used for the SXRPD experiment were obtained from a single crystal that has been exposed at 673 K in air for 7 days, before grinding it into powder with a typical grain size of a few  $\mu\text{m}$ . This powder was then stored for 6 months at ambient conditions prior to the diffraction studies. The  $O$ -stoichiometry of the as-treated samples has been thoroughly analysed by TG measurements in hydrogen atmosphere, revealing an overall stoichiometry of  $\text{Nd}_2\text{NiO}_{4.23}$  as shown in Fig. 2a. Further on, we studied the thermal evolution of the  $O$ -stoichiometry of  $\text{Nd}_2\text{NiO}_{4.23}$  as outlined in Fig. 2b. Upon heating, an  $O$ -uptake starts to set in slightly before 600 K, while beyond 700 K  $O$ -release is observed, reaching a value of  $\delta = 0.19$  at 1273 K. Upon cooling a reversible  $O$ -uptake sets in, showing a maximum at 600 K, and maintaining a final stoichiometry corresponding to  $\text{Nd}_2\text{NiO}_{4.25}$  at room temperature. We thus note that the interstitial oxygen atoms remain confined inside the structure at  $\delta = 0.25$ , while getting released towards  $\text{Nd}_2\text{NiO}_{4.23}$  on a timescale of several weeks.

From laboratory XRPD measurements on a polycrystalline sample that was kept in ambient conditions for more than 6 months, NNO appears to adopt the standard  $\text{K}_2\text{NiF}_4$  structure for Ruddlesden–Popper type oxides, resembling the high temperature orthorhombic (HTO) phase reported previously for  $\text{Nd}_2\text{NiO}_{4+\delta}$  with  $\delta \approx 0.21$ – $0.224$ .<sup>23</sup> A closer inspection, however, evidences a subtle splitting of all ( $hhl$ )-type reflections, related to a small monoclinic distortion (see Fig. S1, ESI<sup>†</sup>). The monoclinic splitting is also illustrated in the inset of Fig. 3(a) at room temperature for (448) and ( $\bar{4}$ 48) structural Bragg peaks, while its  $T$ -dependence is plotted in Fig. 3(b). The monoclinic space group  $F112/m$  (a non-standard setting of space group  $C112/m$ ), corresponding to the low temperature monoclinic (LTM) phase, satisfactorily describes the positions of the main structural reflections as well as the monoclinic splitting of ( $hhl$ )-type Bragg reflections. Rietveld refinements are illustrated in Fig. 3(a) and Fig. S1 (ESI<sup>†</sup>) for the 300 K SXRPD data and laboratory XRPD data, respectively, showing a good agreement between the experimental and calculated patterns. The cell

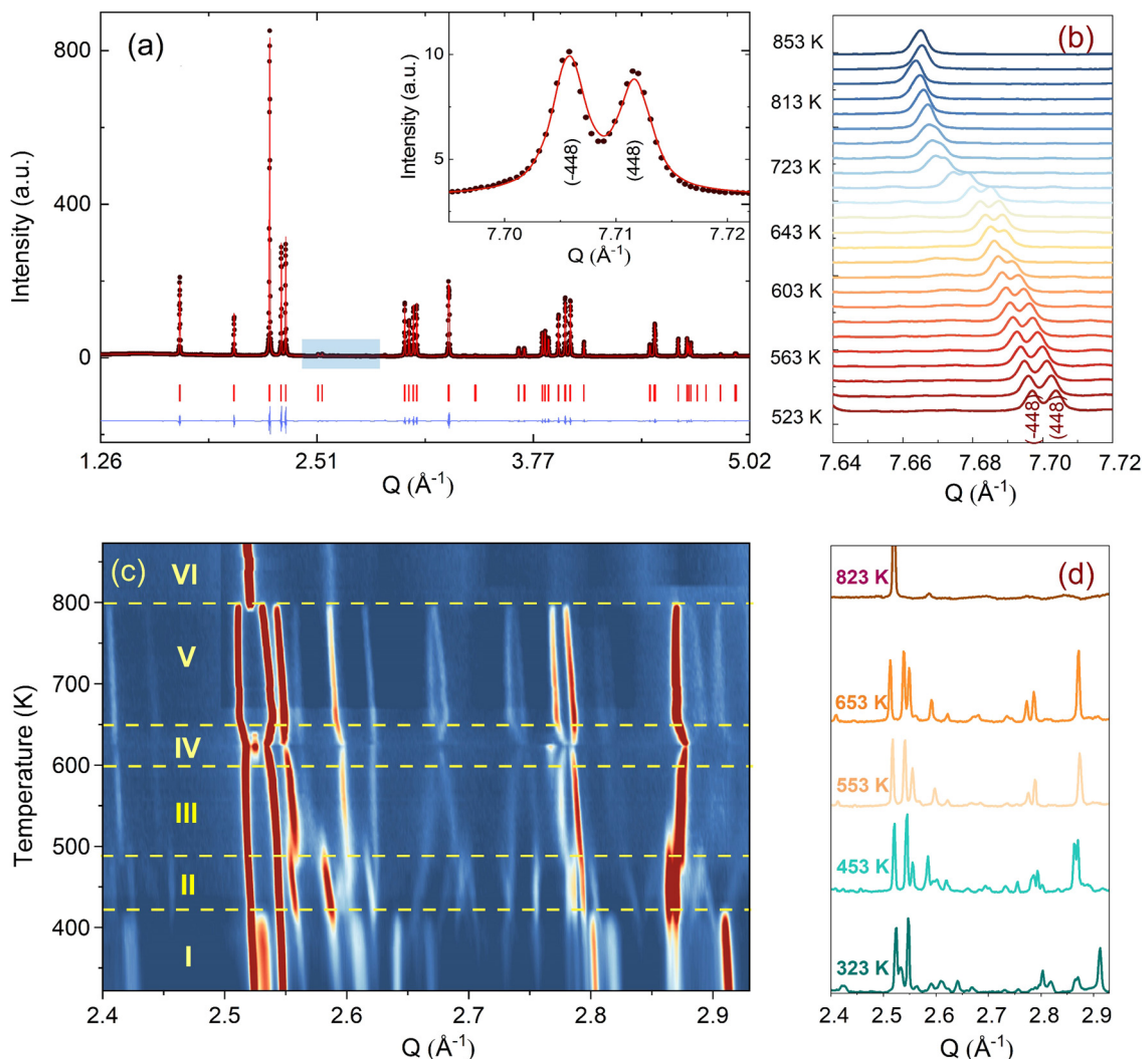
parameters at RT are refined to be  $a = 5.3908(1)$  Å,  $b = 5.4510(1)$  Å,  $c = 12.3695(2)$  Å,  $\alpha = \beta = 90^\circ$  with  $\gamma = 90.076(4)$ .<sup>19,41</sup>

From the SXRPD data the presence of extra reflections become evident (*cf.* Fig. 3(c and d)), which are 3 orders of magnitude lower compared to the strongest Bragg peaks. These reflections are not due to a simple increase of the unit cell but indicative for a 3D modulated oxygen ordering. For the diffraction data at RT, they could be indexed using reconstructed diffraction planes obtained from single crystal X-ray diffraction as further outlined below.

A correct indexation of the satellite reflections is, however, mandatory for the interpretation of the complex  $T$ -dependent structural changes obtained for NNO (see Fig. 3c). Fig. 4(a) shows the reconstructed ( $hk1$ )-plane of a single crystal, which has been stored at ambient conditions for more than 4 years. It reveals a 3D modulated structure, similar to what has been recently reported for the homologous  $\text{Pr}_2\text{NiO}_{4+\delta}$  system.<sup>31</sup> The reconstructed ( $hk1$ )-plane shows the typical splitting of the main reflections, related to the twinning as a consequence of the tetragonal/orthorhombic symmetry reduction, involving two out of four possible orthorhombic twin domains sharing a common ( $\bar{1}10$ )-plane. Thereby each orthorhombic twin domain can further on split up into four monoclinic twin domains, following the twin relations described in Fig. S2 (ESI<sup>†</sup>). The possible maximum number of monoclinic twin domains which may be expected here is consequently eight. The important number of satellite reflections as illustrated in Fig. 4a can then be understood as the overlay of all eight monoclinic twin domains with a modulation vector of  $\mathbf{Q}_n = \pm 0.813\mathbf{a}^* \pm 0.519\mathbf{b}^*$  at RT. The simulation of the peak positions, indicated by white circles, considers the generation of satellite reflections up to 2<sup>nd</sup> order harmonics in space group  $F112/m$ . It also involves the separation due to the orthorhombic twin domains, finally demonstrating an excellent agreement with the experimental data. While the positions of the satellite reflections can be precisely simulated, their intensities cannot. A decrease in the intensity is expected to go along with the order of the satellites, so the simulations consider a relative intensity description with the diameter of the circles being largest for the basic reflection, while decreasing from the 1<sup>st</sup> order, becoming smallest for the 2<sup>nd</sup> order satellites.

We note that the observed modulation vector for NNO at RT up to  $\approx 400$  K is close to a lock-in transition at  $\mathbf{Q}_n = \pm 0.833\mathbf{a}^* \pm 0.5\mathbf{b}^*$ , and thus equivalent to  $\mathbf{Q}_n = \pm 5/6\mathbf{a}^* \pm 0.5\mathbf{b}^*$  as found for  $\text{Pr}_2\text{NiO}_{4.225}$ .<sup>31</sup> The latter involves in fine a commensurate description with a corresponding unit cell of  $10a \times 6b$ . The incommensurate modulation vector  $\mathbf{Q}_n = \pm 0.813\mathbf{a}^* \pm 0.519\mathbf{b}^*$ , obtained from the single crystal at RT, could have been applied successfully as starting values, to index the SXRPD data of phase (I) of NNO, as shown in Fig. S3 (ESI<sup>†</sup>). Due to the significant intensity reduction between the 1<sup>st</sup> and the 2<sup>nd</sup> order satellite reflections, the pattern could already be fitted considering only the 1<sup>st</sup> order satellite reflections, yielding a refined modulation vector for phase (I) at 373 K being  $\mathbf{Q}_n = \pm 0.8245(1)\mathbf{a}^* \pm 0.5230(1)\mathbf{b}^*$ .





**Fig. 3** (a) A representative Rietveld fit refinement of the room temperature SXRPD pattern of  $\text{Nd}_2\text{NiO}_{4.23}$  in space group  $F112/m$ . The experimental data are shown with filled circles while the calculated pattern is represented by a continuous line. The blue continuous line represents the difference curve. Red ticks show the positions of the structural Bragg peaks. The inset shows the splitting of  $(448)$  and  $(\bar{4}48)$  structural Bragg peaks resulting from the monoclinic distortion at room temperature. (b) The temperature evolution of the splitting between  $(448)$  and  $(\bar{4}48)$  structural Bragg peaks. Higher temperature data are shifted by along the  $y$ -axis for clarity. (c) The 2D contour plot representing the thermal evolution of oxygen superstructure reflections in the  $Q$  range of 2.4–2.95  $\text{\AA}^{-1}$ , as shown with a blue box in (a). (d) Illustration of oxygen superstructure reflections at different temperatures in the powder data. The measurement was performed at MS-X04SA in SLS, PSI with  $\lambda = 0.5646(3)$   $\text{\AA}$ .

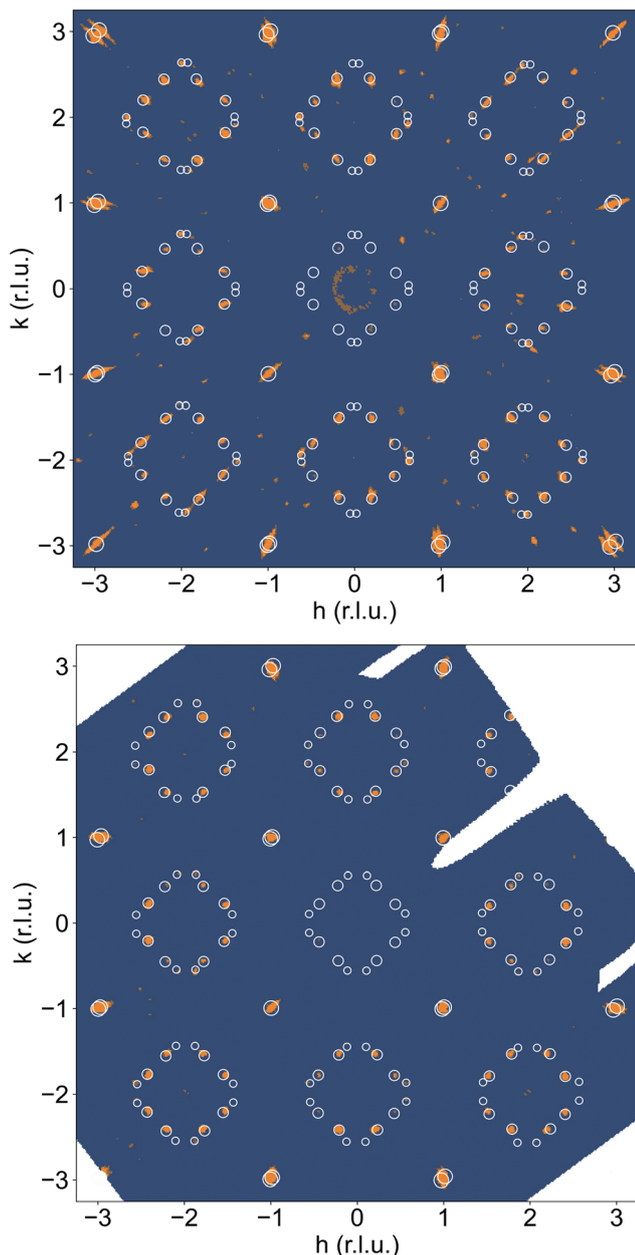
### Temperature dependent SXRPD studies

**Heating.** We investigated the high- $T$  phase diagram of  $\text{Nd}_2\text{NiO}_{4.23}$  by synchrotron powder diffraction measurements from RT up to 873 K in order to determine structural changes with increasing  $T$ . Fig. 3(c) shows a 2D colormap of the evolution of the satellite reflections in the  $Q$ -range of 2.4–3  $\text{\AA}^{-1}$ ; selected patterns are given in Fig. 3(d). This section doesn't show any basic reflections, except for the very weak  $(202/022)$  and is therefore ideally suited to follow up variations of the satellite positions and intensities. A total of 6 different phases can be identified in the investigated  $T$ -range, the monoclinic/tetragonal phase transition appearing at 800 K. We notice that all phase transitions are characterized by a sharp transition temperature, suggesting being all of 1<sup>st</sup> order. Together with well-defined

satellite profiles, it confirms fast-ordering kinetics as well as a very homogeneous definition of the oxygen stoichiometry throughout the sample. From the complex patterns, it is in principle very difficult to determine the changes in the modulation vectors, due to a non-reasonable high number of possible solutions for their indexation.

We could anyhow characterize the modulation vector for the high- $T$  phase  $\text{Nd}_2\text{NiO}_{4.25}$ , related to its structural kinetic stability upon cooling to ambient conditions. We therefore heated a pristine  $\text{Nd}_2\text{NiO}_{4.23}$  single crystal from RT to 723 K, which, after oxygen uptake to  $\text{Nd}_2\text{NiO}_{4.25}$  was cooled down to RT within 24 h, and subsequently investigated by single crystal X-ray diffraction measurements. Fig. 4b shows the  $(hk1)$  plane for the as-obtained  $\text{Nd}_2\text{NiO}_{4.25}$ , which significantly differs from

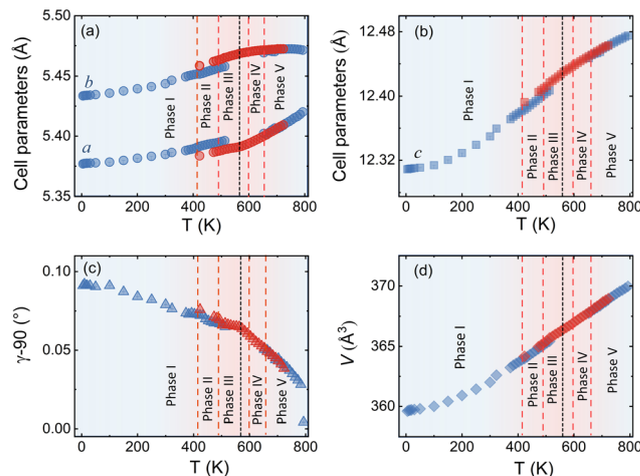




**Fig. 4** Section of reconstructed  $(hk1)$  planes obtained with single X-ray diffraction (STOE StadiVari, MoK $_{\alpha}$ ) measurements at 300 K on a NNO single crystal (a) which has been stored at ambient conditions (starting phase) for more than 4 years and (b) after the heating to 723 K in O $_2$ , and subsequent cooling to room temperature within 24 h. Simulated positions of the superstructure reflections are overlaid onto the experimental data with white circles. Both phases correspond to (a)  $\delta \approx 0.23$  and (b) 0.25, respectively, as revealed by precise TG measurements. The superstructure reflections in the starting phase correspond to a modulation vector  $\mathbf{Q}_n = 0.813(2)\mathbf{a}^* + 0.519(2)\mathbf{b}^*$  up to 3<sup>rd</sup> order, while those in the end phase correspond to a modulation vector  $\mathbf{Q}_n = 7/9\mathbf{a}^* + 5/9\mathbf{b}^*$  up to 2<sup>nd</sup> order. The simulations consider a relative intensity modification with the diameter of the circles being largest for the basic reflection while decreasing with the order of satellites.

Nd $_2$ NiO $_{4.23}$  (Fig. 4a) for both, the orthorhombic splitting of the twin domains, as well as for the modulation vector. Satellite

reflections up to the 2<sup>nd</sup> order are present, yielding a modulation vector of  $\mathbf{Q}_n = \pm 0.7778\mathbf{a}^* \pm 0.5556\mathbf{b}^*$ . Using this vector as starting value for the indexation of the SXRPD data obtained for phase (V), Rietveld refinements yielded  $\mathbf{Q}_n = \pm 0.7835(1)\mathbf{a}^* \pm 0.5524(1)\mathbf{b}^*$  at 703 K (see Fig. S4, ESI $^\dagger$ ). These values nearly coincide with the formation of a commensurate periodicity and thus  $\mathbf{Q}_n = \pm 7/9\mathbf{a}^* \pm 5/9\mathbf{b}^*$  with a corresponding unit cell of  $14\mathbf{a} \times 10\mathbf{b}$ . Beside the important thermal stability of this modification up to 800 K, this monoclinic phase (V) shows consequently translational periodicities of 94 Å. This is deduced from the length of the diagonal in the  $(a,b)$ -plane of the commensurate unit cell with  $14\mathbf{a} \times 10\mathbf{b}$ , the corresponding lattice parameters being  $a = 5.40653(2)$  Å,  $b = 5.47047(2)$  Å and  $c = 12.45542$  Å. Above 800 K, the symmetry changes from monoclinic to tetragonal, while no sharp satellite reflections are observed anymore. Some broad reflections, however, continue to exist even beyond the phase transition, indicating that structural correlations still exist, however, on a shorter length scale. The  $T$ -dependence of the monoclinic distortion is outlined in Fig. 3b as well as in Fig. 5c, together with all other lattice parameters, indicating this discontinuous transition to be of 1<sup>st</sup> order. The characteristics for all other phase transitions are different as they all preserve a monoclinic symmetry. Thereby the positions of all basic reflections, e.g.  $(202/022)$  in Fig. 3c, remain at the same angular position in the  $T$ -range from RT to 600 K (phases I–III), while satellite reflections change positions and intensities. The  $(448)$  and  $(\bar{4}48)$  reflections are clearly splitted up to 800 K, indicating the monoclinic symmetry. It shows that



**Fig. 5** Variation of lattice parameters of the average monoclinic unit cell extracted from SXRPD data of Nd $_2$ NiO $_{4.23}$  as a function of temperature. Data obtained during heating and cooling are given in blue and red symbols, respectively. Six different modulated phases which are distinguished during heating are also marked while the red dashed lines roughly indicate the transition temperatures. The black dotted line at ca. 550 K denotes the temperature at which the modulation vector of the excess oxygen ordering changes upon cooling. The missing data points in the temperature range of 723–853 K during cooling are due to a technical problem during data collection. Lattice parameters around the formation of phase IV are not given due to the presence of oxygen uptake in this region, accompanied by the formation of additional broadened reflections, rendering a uniform refinement difficult.



the integrity of the  $\text{Nd}_2\text{NiO}_{4+\delta}$  framework remains predominantly untouched, while several defined phases with different modulation vectors emerge. On a qualitative basis it becomes thus clear that distinct oxygen ordering scenarios get established, each showing a well-defined periodicity, following a topotactical rearrangement, essentially concerning the oxygen atoms. It also suggests that all phases are close energetically and can easily switch in between these different ordering arrangements.

The comparison of the reflection positions and their respective intensities suggest that phases (V) and (III) are pretty much identical, which becomes also clear from the refinement of the modulation vector being  $\mathbf{Q}_n = \pm 0.7778(1)\mathbf{a}^* \pm 0.5556(1)\mathbf{b}^*$  at 583 K as displayed in Fig. S5 (ESI<sup>†</sup>). Both phases are separated by region (IV), where additional diffraction peaks can be identified between 600 K and 650 K. Comparison with the TG results shows that in this region an oxygen uptake into the lattice occurs, which is also indicative for the slight changes in the scattering angles of the basic reflections. Phases (III) and (V) consequently have a different oxygen stoichiometry: while phase (III) still shows the initial stoichiometry corresponding to  $\text{Nd}_2\text{NiO}_{4.23}$ , phase (V) is supposed to have a stoichiometry with  $\delta$  close to 0.25. We note that upon heating, diffuse type additional reflections (see Fig. 3c) appear at  $T \geq 523$  K, indicating the development of a secondary phase, coexisting with phases (III) and (IV). Attempts to determine the corresponding lattice parameters of this secondary phase at  $T \approx 625$  K were so far unsuccessful, as it is probably composed as an intergrowth phase, showing a kind of layer stacking disorder, similar to the interstratified arrangements of graphite or silicate frameworks.

Phase (II) appearing at around 400 K is an interesting case, showing a discontinuous modification of the modulation vector, together with small but significant changes in the lattice parameters as shown in Fig. 3c and 5(a–d). Compared to the starting phase (I), the modulation vector for phase (II) considerably changes towards the values found for phases (III) and (V) which can be refined to  $\mathbf{Q}_n = \pm 0.7762(1)\mathbf{a}^* \pm 0.5623(1)\mathbf{b}^*$  at 433 K, as demonstrated in Fig. S6 (ESI<sup>†</sup>). In the absence of single crystal data, this phase does not permit an unambiguous assignment so far. The slight splitting of the reflection at  $\mathbf{Q} = 2.87 \text{ \AA}^{-1}$  could, however, only be achieved with a slight deviation of the  $\mathbf{b}^*$  component from the commensurate indexation ( $\pm 0.5556\mathbf{b}^*$ ) as observed for phases (III), (V) and (VI) as discussed outlined above.

**Cooling.** During cooling, the average monoclinic symmetry reversibly reappears below the phase transition temperature at  $T \approx 800$  K as revealed by laboratory temperature-dependent X-ray diffraction measurements (see Fig. S7, ESI<sup>†</sup>). Due to a technical problem during data collection, we could not measure the complete  $T$ -range and only patterns below 723 K became accessible.

A 2D colormap representing the temperature evolution of oxygen superstructure reflections during cooling in the  $\mathbf{Q}$  range of  $2.1\text{--}2.95 \text{ \AA}^{-1}$  is shown in Fig. S8 (ESI<sup>†</sup>). Two phases, corresponding to phase (V) and (VII) could clearly be distinguished

during cooling, showing again a sharp transition situated at around 550 K.

The modulation vector above 550 K, describing the superstructure peaks of phase (V), was refined to be  $\mathbf{Q}_n = \pm 0.7835(1)\mathbf{a}^* \pm 0.5524(1)\mathbf{b}^*$  at 623 K (see Fig. S9, ESI<sup>†</sup>) *i.e.* pretty identical with the modulation vector determined for phase (V) during heating. Noticeable changes are found below this temperature in the evolution of lattice parameters as represented in Fig. 5. Below 550 K, the modulation vector remains almost the same down to 300 K, and it was refined to  $\mathbf{Q}_n = \pm 0.7734(1)\mathbf{a}^* \pm 0.5569(1)\mathbf{b}^*$  at 423 K (see Fig. S10, ESI<sup>†</sup>).

Thus, the modulation vector at room temperature after cooling is found to be slightly different from the one before heating, which is justified by the difference of the oxygen content ( $\Delta\delta \approx 0.02$ ) between starting and end phase.

### Single crystal neutron diffraction and maximum entropy analysis

In view of the different phase transitions described above, we were also interested in the respective evolution of thermal displacement parameters, at least on an average scale, especially for the apical and equatorial oxygen atoms, and their correlation with the ordering of excess oxygen atoms. For this purpose, high-resolution single crystal neutron diffraction measurements were carried out, taking advantage of the increased sensitivity of neutrons for low- $Z$  atoms, *i.e.*, oxygen. Due to the lack of suitable 2D neutron area detectors, this study only considers the basis reflections, *i.e.*, reflections fulfilling the conditions for  $F$ -centering in the whole  $T$ -range. The second reason making a data collection of satellite reflections problematic is related to the superposition of the satellite intensities as resulting from the formation of pseudo-merohedral twin domains, causing an important overlay of different twin domains not allowing to obtain unique structure factors for the superstructure reflections.

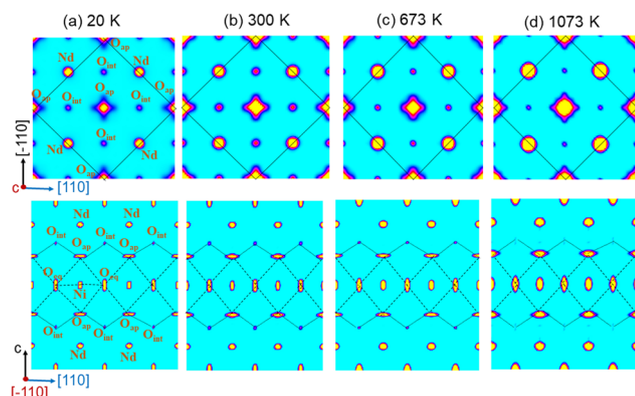
The superposition of intensities from the pseudo-merohedral twin domains also causes a systematic overlay of all main Bragg reflections (see Fig. S2, ESI<sup>†</sup>). We consequently averaged all overlaid integrated intensities of ( $hkl$ ) and ( $kh$ l) type Bragg reflections, resulting into a refinement by least squares methods in the tetragonal  $F4/mmm$  space group using JANA2006, despite the real symmetry being monoclinic. The results of all refinements are summarized Table 1.

At 300 K, the oxygen over-stoichiometry  $\delta$  in the crystal structure is refined to be 0.238(17), in agreement with the value obtained from TGA analysis of the polycrystalline sample. The reason for the lower occupancies for the  $\text{O}_{\text{ap}}$  site with respect to a full site occupancy, is related to strong and anharmonic displacements of all  $\text{O}_{\text{ap}}$  atoms, as discussed in ref. 19 and which do not allow to cover all  $O$ -scattering density within a harmonic description. Fig. 6 illustrates the 2D nuclear density distributions for different sections, obtained with the MEM algorithm at various temperatures. The nuclear density data were obtained taking into consideration the phased structure factors obtained for the refinements given in Table 1. We note that only 1/8 of all available interstitial sites are effectively occupied.



**Table 1** Structural parameters obtained from single crystal neutron diffraction data of  $\text{Nd}_2\text{NiO}_{4.23}$  compound in  $F4/mmm$  space group. Thermal parameters  $U_{ij}$  are given in  $\text{\AA}^2$ . Single crystal data at 20 K and 300 K were recorded on Trics/ZEBRA ( $\lambda = 1.178(1) \text{\AA}$ ) at SINQ, PSI while 673 K and 1073 K data were measured on D9 ( $\lambda = 0.837(1) \text{\AA}$ ) at ILL

Atoms	20 K	300 K	673 K	1073 K
Ni (0 0 0)				
Occ.	1	1	1	1
$U_{11} = U_{22}$	0.0025(6)	0.0041(5)	0.0086(4)	0.0118(3)
$U_{33}$	0.0064(7)	0.0141(7)	0.0238(5)	0.0327(6)
Nd (00z)				
Occ.	2	2	2	2
$z$	0.3598(12)	0.3596(11)	0.35933(8)	0.36016(9)
$U_{11} = U_{22}$	0.0080(5)	0.0121(5)	0.0209(5)	0.0269(4)
$U_{33}$				
$O_{\text{ap}}$ (00z)				
Occ.	1.70(2)	1.74(2)	1.66(3)	1.82(2)
$z$	0.1728(2)	0.1735(3)	0.1735(2)	0.1759(2)
$U_{11} = U_{22}$	0.058(2)	0.060(2)	0.077(3)	0.085(3)
$U_{33}$	0.0016(16)	0.0076(15)	0.0120(10)	0.0190(10)
$O_{\text{eq}}$ (0.25,0.25,0)				
Occ.	2.007(15)	1.985(14)	2.027(19)	2.026(15)
$U_{11} = U_{22}$	0.0044(8)	0.0065(7)	0.0165(7)	0.0216(5)
$U_{33}$	0.0490(18)	0.0520(16)	0.0635(16)	0.0648(16)
$U_{12}$	0.0004(7)	-0.0014(6)	-0.0061(5)	-0.0087(5)
$O_{\text{int}}$ (1/4 1/4 1/4)				
Occ.	0.238(17)	0.252(16)	0.128(14)	0.112(17)
$U_{\text{iso}}$	0.033(9)	0.036(8)	0.040(6)	0.046(16)
$R_{\text{p}}$ (%)	5.65	5.81	5.63	4.35
$R_{\text{wp}}$ (%)	8.15	7.6	8.29	6.30



**Fig. 6** Maximum entropy analysis of single crystal neutron diffraction data of  $\text{Nd}_2\text{NiO}_{4.23}$  obtained at (a) 20 K, (b) 300 K, (c) 673 K, and (d) 1073 K. (Upper row) (110)– $(\bar{1}10)$  projection of 2D nuclear density maps particularly showing  $O_{\text{ap}}$  displacements toward the nearest  $O_{\text{int}}$  site. The black continuous line outlines the  $F$ -centred average chemical unit cell. (Lower row) (110)/(001) projection of 2D nuclear density maps displaying the  $O_{\text{ap}}-O_{\text{int}}-O_{\text{ap}}$  oxygen diffusion pathway along the [110] direction. Single crystal data at 20 K and 300 K were recorded on ZEBRA/TriCS ( $\lambda = 1.178(1) \text{\AA}$ ) at SINQ, PSI while 673 K and 1073 K data were measured on D9 ( $\lambda = 0.837(1) \text{\AA}$ ) at ILL.

The  $O_{\text{ap}}$  shows a  $U_{12}$  contribution in  $F4/mmm$ , contrary to the standard setting in  $I4/mmm$ . This is related to the change of the point group symmetry, being  $(m.2m)$  and  $(mmm)$  for the respective positions in  $(xx0)$  in  $F4/mmm$ , and  $(\frac{1}{2}, 0, 0)$  in  $I4/mmm$ . For the  $(xx0)$ -position in  $F4/mmm$  the  $x$  value is then constrained to exactly  $\frac{1}{4}$ . A strong anharmonic contribution of the  $O_{\text{ap}}$  displacement is observed specifically along [110] with

respect to the  $F$ -symmetry cell, *i.e.* towards the nearest interstitial lattice site as evidenced Fig. 6a–d. As indicated in the  $ab$ -projection of the rock salt layer in the range of  $0.14 \leq z \leq 0.28$ , strong and defined  $O_{\text{ap}}$  displacements are also observed at 20 K, indicating a significant static contribution to the  $O_{\text{ap}}$  shifts. This static component can most probably be attributed to the incommensurate structural modulations, representing all displacements of the  $O_{\text{ap}}$  atoms from their average positions. The scattering densities of  $O_{\text{ap}}$  atoms are strongly enhanced at 300 K along [110] indicating a shift of about  $2 \text{\AA}$  from the average position, corresponding to a tilt of *ca.*  $25^\circ$  for the  $\text{NiO}_6$  octahedra. It presents the average deviation from the equilibrium position of  $O_{\text{ap}}$  in (00z), taking into account a dynamical tilting along [110], as discussed in ref. 19. They are essentially related to the presence of interstitial oxygen atoms in  $(\frac{1}{4}, \frac{1}{4}, \frac{1}{4})$ , pushing away the adjacent apical oxygen atoms, related to steric constraints. This is the main reason also why dynamically activated diffusion takes place for Ruddlesden Popper type oxides only for  $\delta > 0$ . The strongly enhanced displacements of the  $O_{\text{ap}}$  atoms in the  $(a,b)$ -plane are already reflected in the anisotropic displacement factors in Table 1, where  $U_{11}$  (eq. to  $U_{22}$ ) presents at least four times to the value of  $U_{33}$ . The anharmonic description which implicitly results from the MaxEnt analysis then clearly indicates the large components to be along [110], which cannot be analyzed from a harmonic refinement in  $F4/mmm$ , as the main displacement components are defined with respect to the principal axes.

The strong enhancement of scattering density at already 300 K shows an increasing dynamical origin with temperature, in parallel to contributions from incommensurate structural modulations. The displacements of the  $O_{\text{ap}}$  atoms thus allow a symmetric increase of the  $O_{\text{int}}-(O_{\text{ap}})_4$  tetrahedra with an  $O_{\text{int}}-O_{\text{ap}}$  distance of about  $2.8 \text{\AA}$ , compared to  $2.15 \text{\AA}$  for the radius of an empty tetrahedron. These findings are fully equivalent to what is reported for the homologous  $\text{Pr}_2\text{NiO}_{4+\delta}$ , indicating an oxygen diffusion pathway between apical and vacant interstitial sites in the [110]-direction inside the rock salt layer. Such a dynamical activation is also in agreement with a phonon-assisted diffusion mechanism, which becomes of importance in view of the diffusion pathway along [110],<sup>7,13,15,19,42</sup> showing a sequence of filled and empty  $(O_{\text{ap}})_4$  tetrahedra, of strongly different size (see Fig. 1c). The oxygen diffusion mechanism close to room temperature therefore involves important local structural distortions that are far from a rigid body description but resembling along the diffusion pathway to a successively in- and exhaling mechanism of the involved tetrahedra. At higher  $T$  (673 K), the incommensurate structural modulations vary, becoming different to those at room temperature, showing a significantly broader delocalization of the  $O_{\text{ap}}$  atoms within the  $ab$ -plane and likely due to enhanced thermal contributions. However, no double-well potential, as reported for the homologous  $\text{Pr}_2\text{NiO}_{4+\delta}$ ,<sup>19</sup> is observed for  $\text{Nd}_2\text{NiO}_{4+\delta}$ , related to the shift of the tetragonal phase transition towards higher temperatures.

A significantly different scenario is, however, observed for  $O_{\text{ap}}$  atoms at 1073 K, *i.e.* above the monoclinic/tetragonal phase



transition, clearly indicating a more isotropic nuclear scattering density distribution in the *ab*-plane. At this temperature all satellite reflections related to oxygen ordering are vanished, indicating a disordered and thus uncorrelated  $O_{\text{int}}$  scenario.

## Discussion

While oxygen can be reversibly intercalated into  $\text{Nd}_2\text{NiO}_{4+\delta}$  at room temperature, our findings evidence for the first time an easy switching of long-range oxygen ordering at the sub-mesoscopic length scale with fast ordering kinetics. This directly implies the presence of extremely mobile interstitial oxygen atoms, resulting into the formation of long-range oxygen ordering, organized as 3D modulated structures, manifested by the observation of a series of discontinuous, *i.e.* 1<sup>st</sup> order phase transitions, in a *T*-range from RT to 800 K. The incommensurate modulation vector for  $\text{Nd}_2\text{NiO}_{4.23}$  evolves from  $\mathbf{Q}_n = \pm 0.8126\mathbf{a}^* \pm 0.5185\mathbf{b}^*$  at RT to  $\mathbf{Q}_n = \pm 0.7762\mathbf{a}^* \pm 0.5623\mathbf{b}^*$  at 433 K, towards  $\mathbf{Q}_n = \pm 0.7778\mathbf{a}^* \pm 0.5556\mathbf{b}^*$  at 583 K, the latter corresponding into a commensurate indexation with  $\mathbf{Q}_n = \pm 7/9\mathbf{a}^* \pm 5/9\mathbf{b}^*$ . However, this lock-in transition towards a commensurate indexation is not the final state and further evolves, along with an oxygen uptake, towards  $\text{Nd}_2\text{NiO}_{4.25}$  with  $\mathbf{Q}_n = \pm 0.7835\mathbf{a}^* \pm 0.5524\mathbf{b}^*$  at 703 K.

Thus, the  $O_{\text{ap}}$  oxygen atoms not only show a much more isotropic delocalization and consequently a loss of correlations with the excess oxygen atoms. Therefore, they are supposed to be highly mobile in this temperature range as observed for  $\text{Pr}_2\text{NiO}_{4+\delta}$  in the disordered tetragonal phase.

Such a series of complex reorganisation of the oxygen ordering is completely unexpected and suggests all phases to stay on a quite similar energetical landscape. This also shows that oxygen is sufficiently mobile to build up new but structurally well-defined ordering scenarios, all on a large-scale. More importantly, they all involve a rapid reorganization of the ordering schemes, thus implying fast ordering kinetics. On the other hand, long-range correlations are indicative for cooperative diffusion mechanisms, suggesting that lattice instabilities, *i.e.*, the apical oxygen atoms following octahedra tilting, may result into a 1D shallow diffusion pathway along [110], directly involving alternating vacant and filled interstitial lattice sites as shown in Fig. 1c. In this way incommensurate periodicities either created by structural distortions or modulated distribution of the  $O_{\text{int}}$  atoms, may then contribute into a soliton-like diffusion mechanism, following non-linear structural as well as anharmonic contributions of dynamic origin.

The unusually high displacement factors found for the apical oxygen atoms by single crystal neutron diffraction, thereby suggest a strong deviation of all  $O_{\text{ap}}$  from their equilibrium positions due to the structural modulation.

In analogy to  $\text{Pr}_2\text{NiO}_{4+\delta}$ , the changes of the modulation vector of as grown single crystals observed with time while exposing the sample at ambient conditions, clearly indicate structural changes, which, together with the thermogravimetric studies (Fig. 2b) demonstrate that oxygen diffusion proceeds at RT, linked to long-range oxygen ordering.

Apart from the high  $O_{\text{int}}$  mobility at already moderate temperatures, the thermal stability of oxygen ordering up to 800 K appears to be surprising. It confirms the idea of a dynamically decoupled  $\text{Nd}_2\text{O}_2$  rock salt layer, separated by infinite  $\text{Ni}_2\text{O}_2$  layers as discussed for the homologous  $\text{Pr}_2\text{NiO}_{4+\delta}$  phase.<sup>19</sup> While at the same time  $O_{\text{int}}$  atoms are easily mobile inside the rock salt layer, they strongly distort locally the  $O_{\text{int}}(O_{\text{ap}})_4$  tetrahedra, which consequently becomes the origin of establishing a long-range organisation of all occupied and empty tetrahedra. It thus induces naturally a lattice dynamical component to trigger the oxygen diffusion along the [110] direction of the 1D tetrahedral chains.

One may suppose that the 1D character of the diffusion pathway together with the oxygen ordering is stabilising the monoclinic symmetry in oxygen rich  $\text{Nd}_2\text{NiO}_{4+\delta}$  phases, which transform directly into a tetragonal high-*T* parent phase showing  $O_{\text{int}}$  disorder. It is remarkable that no orthorhombic intermediate phase is identified during this phase sequence. The presence of  $O_{\text{int}}$  atoms thus induces strong structural correlations which, however, still allow a fine tuning with temperature as encountered here, indicated by different ordering schemes. Similar correlations have already been shown to induce electronic correlations for  $\text{Nd}_2\text{NiO}_{4.1}$  resulting into identical 3D modulation vectors for oxygen as well as spin ordering.

The strong anisotropic displacements as determined from the scattering density maps shown in Fig. 6 are strongest at lower temperatures, becoming more isotropic with increasing *T*. It becomes clear that at high *T* practically all lattice modes are sufficiently activated to obtain an isotropic diffusion behaviour, that can be described by the Arrhenius Ansatz. Given the anisotropic displacements at lower *T* to be at least partially of dynamic origin, this lattice dynamics could consequently promote an oxygen diffusion mechanism in the direction of these displacements. Similarly, anisotropic displacements have been found for bcc metals close to their bcc/hcp phase transition above 1000 °C, *e.g.* Ti or Hf, together with a significant increase of their self-diffusion coefficient by several orders of magnitude.<sup>43–45</sup> The same type of phase transition and respective increase of the self-diffusion coefficient is also observed below 2 K for solid He, excluding that the increase for self-diffusion, is not limited by the temperature.<sup>46,47</sup> For all these bcc/hcp phase transitions the increase of self-diffusion is essentially coupled to phonon softening. The dynamically activated anharmonic lattice dynamics of the  $\text{RE}_2\text{O}_2$  rock salt layers of oxygen-doped Ruddlesden–Popper type oxides, is supposed to have a very similar impact on the self-diffusion coefficients, here given as the oxygen mobility. Then, phonon softening can be easily promoted due to long-range oxygen ordering, which may result in lattice instabilities leaving the apical oxygen atoms on a shallow potential.

## Conclusions

We have evidenced a series of complex order/order phase transitions generation for  $\text{Nd}_2\text{NiO}_{4+\delta}$ , resulting into well-defined



oxygen ordering scenarios up to the sub-mesoscale, together with fast ordering kinetics. The fast rearrangement for different oxygen ordering schemes thereby directly involves a rapid oxygen mobility in a very moderate temperature range. These findings are new and suggest interpreting related oxygen diffusion mechanisms to be strongly correlated and/or cooperative. Such a mechanism is supposed to get amplified by phonon softening, *e.g.* engendered as a consequence of lattice instabilities, based on long-range oxygen ordering. Thereby, the oxygen-doped Ruddlesden–Popper phases present a special scenario, as the tetrahedral  $O_{\text{int}}(O_{\text{ap}})_4$  chains inside the  $\text{RE}_2\text{O}_2$  rock salt layer appear as an interface, dynamically decoupled from the rest of the structure. Such type of directional diffusion mechanism can in principle be understood as an anisotropic Arrhenius behaviour, in which an essential part of the lattice energy is projected along one direction, yielding a shallow, 1D diffusion pathway with low activation energies along the [110]-direction. Oxygen diffusion might then be understood as the result of a constructive lattice dynamics which is in phase with the diffusing species similar to a phason, taking profit of the aperiodicity of the underlying lattice and associated lattice dynamics.

We could also show that ordering kinetics play an important role to stabilize ordering schemes, *e.g.* during the cooling of  $\text{Nd}_2\text{NiO}_{4.25}$  from 723 K to ambient, while conserving the high- $T$  modulation vector. It offers an interesting concept to obtain phases with identical  $O$ -stoichiometry but different modulation vectors. If such a behaviour could be enlarged for other oxygen-doped oxides, it would offer the possibility to probe the influence of structural *vs.* electronic ordering phenomena at the same valence state configuration of a given transition metal atom, enabling to correlate charge- and spin ordering and related dynamics for different modulation vectors at invariant hole concentration. This is a unique opportunity for oxygen-doped phases, and which is not possible in case of Sr-doping, which leads to a statistical distribution of all A-cations. The fact that oxygen is mobile down to ambient conditions allowing to achieve different ordering schemes is therefore a challenging and so far mainly unexplored opportunity to probe correlations between structural and electronic ordering.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge the beam times used on TriCS/ZEBRA at PSI/SINQ and D9 at ILL as well as laboratory equipment and support from LDM/PSI, LBK/PSI, and the “Plateforme d’Analyse et de Caractérisation” of the ICG Montpellier. The authors gratefully acknowledge the financial support from the Swiss National Science Foundation (SNF) through grant 200021L\_157131 and the French National Research Agency (ANR) through grant 14-CE36-0006-01 of the SECTOR project as well as the grant ANR-19-CE05-0041-01 of the ExODiff project. The beamtime

allocation on D9 (DIR-152) is accessible under DOI: [10.5291/ILL-DATA.DIR-152](https://doi.org/10.5291/ILL-DATA.DIR-152).

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