Local-electrostatics-induced oxygen octahedral distortion in perovskite oxides and insight into the structure of Ruddlesden-Popper phases

Youngjae Hong1, Pilgyu Byeon1, Jumi Bak1, Yoon Heo1, Hye-Sung Kim1, Hyung Bin Bae2 & Sung-Yoon Chung1

As the physical properties of ABX3 perovskite-based oxides strongly depend on the geometry of oxygen octahedra containing transition-metal cations, precise identification of the distortion, tilt, and rotation of the octahedra is an essential step toward understanding the structure-property correlation. Here we discover an important electrostatic origin responsible for remarkable Jahn-Teller-type tetragonal distortion of oxygen octahedra during atomic-level direct observation of two-dimensional [AX] interleaved shear faults in five different perovskite-type materials, SrTiO3, BaCeO3, LaCoO3, LaNiO3, and CsPbBr3. When the [AX] sublayer has a net charge, for example [LaO]+ in LaCoO3 and LaNiO3, substantial tetragonal elongation of oxygen octahedra at the fault plane is observed and this screens the strong repulsion between the consecutive [LaO]+ layers. Moreover, our findings on the distortion induced by local charge are identified to be a general structural feature in lanthanide-based An−1BnX3n+1-type Ruddlesden-Popper (RP) oxides with charged [LnO]+ (Ln = La, Pr, Nd, Eu, and Gd) sublayers, among more than 80 RP oxides and halides with high symmetry. The present study thus demonstrates that the local uneven electrostatics is a crucial factor significantly affecting the crystal structure of complex oxides.

1Department of Materials Science and Engineering and KAIST Institute for the Nanocentury, Korea Advanced Institute of Science and Technology, Yuseong-gu, Daejeon 34141, Korea. 2KAIST Analysis Center, Korea Advanced Institute of Science and Technology, Yuseong-gu, Daejeon 34141, Korea.

✉email: sychung@kaist.ac.kr
The crystal structure of ABO$_3$-type perovskites is based on the geometric stability of packing for ions of different sizes and valence states. While the relatively large cation A is coordinated with 12 oxygen anions at the dodecahedral sites, the smaller cation B is located in the oxygen octahedra connected to each other by corner-sharing, agreeing well with Pauling’s first rule for ionic compounds. In particular, the distortion, including tilt and rotation, of oxygen octahedra is known to be a crucial factor governing the dielectric, magnetic, optical, and catalytic properties of perovskites. As a consequence, the direct identification and description of the geometry of oxygen octahedra in perovskites and derivative oxides have been key issues over the last decades.

In addition to the relative ionic size and the cation–anion bond strength, a significant electronic contribution to the structural distortion in oxides has been noted as the Jahn–Teller effect. The central argument of this effect is that the energy of d-orbital electrons of transition-metal ions can be significantly lowered by the splitting of the d-orbital levels. As is already known very well, high-spin Mn$^{3+}$ (3$d^4$) and low-spin Cu$^{2+}$ (3$d^9$) are typical Jahn–Teller active cations, demonstrating that the splitting of the degenerate $d_{x^2}$ and $d_{y^2}$ levels of the $e_g$ subshells and subsequent energy stabilization of electrons is achieved via substantial tetragonal distortion of oxygen octahedra. A key message of the Jahn–Teller effect is that the electronic structure of transition-metal ions should be taken into account to fully understand the crystal (or molecular) structure.

Recently, intriguing local distortion of oxygen octahedra was reported in heteroepitaxial LaNiO$_3$ thin films. Even though this nickelate has a pseudocubic structure, tetragonal elongation of [NiO$_6$] octahedra was identified to take place exclusively at the planes of Ruddlesden–Popper (RP) two-dimensional shear faults. Taking into account that the shear faults, consisting of consecutive [LaO]$^+$ layers, have electrostatically repulsive instability, the protrusion of negatively charged oxygen anions toward the shear plane was reasonably claimed to be energetically favorable for alleviating the strong repulsion between the [LaO]$^+$ layers. Although this previous study was confined to a single material, LaNiO$_3$, provided a physically sound explanation, further extensive work remains necessary in order to clarify and generalize the local-charge hypothesis as another important electrostatic origin responsible for the remarkable tetragonal distortion of oxygen octahedra in perovskite-derivative oxides.

To this end, in this work, we extended our atomic-scale direct observations of two-dimensional shear faults into five different perovskite-type materials, SrTiO$_3$, BaCeO$_3$, LaCoO$_3$, LaNiO$_3$, and CsPbBr$_3$. First, we observed substantial tetragonal distortion of [BO$_6$] octahedra at the fault plane, when an interleaved [AO]–[AO] shear fault has a local charge (for example, [LaO]$^+$ in LaCoO$_3$ and LaNiO$_3$). In stark contrast, no significant structural distortion was probed in the [BO$_6$] octahedra when the [AO] sublayer has no effective charge ([SrO] in SrTiO$_3$, [BaO] in BaCeO$_3$, and [CsBr] in CsPbBr$_3$). Furthermore, by examining a series of 87 $A_{n-1}B_2X_{3n+1}$-type Ruddlesden–Popper (RP) oxides and halides with high symmetry, we found that this local-charge-induced tetragonal elongation of oxygen octahedra is a striking common feature observed in most of the RP phases having the charged [AO] layer. Consequently, our findings on the electrostatic origin of the distortion appear to be a general phenomenon in oxides containing multiple cations.

**Results and discussion**

**Observation of shear faults.** For fault observations, we prepared polycrystalline dense SrTiO$_3$, BaCeO$_3$, and LaCoO$_3$ sintered pellets, epitaxial LaNiO$_3$ thin films, and CsPbBr$_3$ nanocrystals (see the “Methods” section), all of which contain homologous stacking faults represented by two consecutive rocksalt [AX] layers (A = Sr, Ba, La, Cs; X = O, Br). Figure 1 shows a series of high-angle annular dark-field (HAADF) scanning transmission electron microscopy (STEM) images at atomic resolution, verifying the presence of rocksalt [AX] faults in all five perovskite crystals (see Supplementary Fig. 1 for lower magnification images). As the atomic number, Z, of Ba ($Z_{Ba} = 56$) and Ce ($Z_{Ce} = 58$) is very similar, individual columns are not readily discriminated in BaCeO$_3$, although the image contrast of the faults is sufficiently distinguishable at a lower magnification (Supplementary Fig. 1b). A chemical map obtained by electron energy-loss spectroscopy (EELS) is thus provided in Fig. 1b to confirm the [BaO]–[BaO] layers at the fault. In addition, note that the Z value of Cs ($Z_{Cs} = 55$) is much lower than that of Pb ($Z_{Pb} = 82$) in CsPbBr$_3$. (See Supplementary Fig. 2 for additional STEM images for the formation of homologous shear faults. Each image was acquired in **a** SrTiO$_3$, **b** BaCeO$_3$, **c** LaCoO$_3$, **d** LaNiO$_3$, and **e** CsPbBr$_3$. The yellow broken lines in each image indicate the fault planes, and CsPbBr$_3$ nanocrystals consisting of two consecutive [AX] sublayers, as schematically described in **a** together with an annular bright-field image. The atomic-scale chemical map shown in **b** was obtained by using EELS to verify the consecutive [BaO] sublayers.)
images of CsPbBr₃ nanocrystals). Consequently, the column intensity at the [CsBr] faults in a HAADF image should be distinctively lower, as shown in Fig. 1e, in contrast to the image feature at the faults observed in SrTiO₃, LaCoO₃, and LaNiO₃.

**Oxygen displacement and local electrostatics.** To visualize the position of light-element anions, we utilized the annular bright-field (ABF) mode in STEM[22–29]. All the ABF images acquired from the fault region in the five perovskites are provided in Fig. 2 (see Supplementary Fig. 3 for pairs of ABF and HAADF images of the bulk crystals as references). A readily recognizable structural feature from this set of atomic-scale observations is that oxygen octahedra at the faults are asymmetrically elongated along the z axis in LaCoO₃ and LaNiO₃ (Fig. 2b), while no distortion of anion octahedra is identified in the other three perovskites, SrTiO₃, BaCeO₃, and CsPbBr₃, as denoted by a white square diamond in each image in Fig. 2a. In particular, more than a 20% increment in the Co–O length from 1.9 to 2.3 Å in LaCoO₃ and a 15% increment in the Ni–O length from 1.9 to 2.2 Å in LaNiO₃ are presented in the magnified images in Fig. 2b.

These two perovskite groups revealing distinct octahedral aspects have their own local electrostatic characteristics from the viewpoint of a layer structure. As schematically depicted in Fig. 3a, no effective local charge is noted in each of the [AX] and [BX₃] sublayers in three perovskites, SrTiO₃, BaCeO₃, and CsPbBr₃. For example, because the valence states of Sr, Ti, and O in SrTiO₃ are 2+, 4+, and 2−, respectively, the sublayers, [SrO]⁺ and [TiO₂]²−, have no net charges[30, 31]. Therefore, no electrostatic perturbation is induced at the homologous [SrO]–[SrO] fault plane, indicating that the (pseudo)cubic bulk structure can be preserved at the fault region without showing elongation of oxygen octahedra. In the same manner, there is no local charge at the faults in BaCeO₃ ([BaO]²⁻) and CsPbBr₃ ([CsBr]¹⁰⁻).

In contrast, the La-based perovskite oxides, LaCoO₃ and LaNiO₃, contain local net charge at each of the sublayers, as described in Fig. 3b. As the valence states of La, Co (and Ni), and O are 3+, 3+, 3+, and 2−, respectively, the bulk structure of LaCoO₃ (and LaNiO₃) consists of positively charged [LaO]⁺ layers and negatively charged [CoO₂]²⁻ (and [NiO₂]²⁻) layers in an alternating manner along the z direction. Consequently, if two [LaO]⁺ layers are consecutively placed at the faults, electrostatically strong repulsion is inevitably induced, resulting in serious local instability (Fig. 3b). One of the easiest way to reduce the positive-charge repulsion between two [LaO]⁺ layers is that the negatively charged O²⁻ anions adjacent to the fault displace toward the fault plane and thereby screen the effective positive charge. Even if the displacement of oxygen anions is at an angstrom level, it appears to be fairly efficient to significantly reduce the repulsive instability at the fault plane.

**Theoretical calculations.** The displacement of oxygen anions was also verified by ab initio density functional theory (DFT) calculation. Figure 3c shows the atom position variation in a LaNiO₃ supercell containing a shear fault during the DFT calculation for geometry optimization. As shown in the supercell illustration of the first self-consistent field (SCF) iteration step, the initial configuration of oxygen atoms (light red) at the fault plane was set to be identical to the position of oxygen in the bulk. It is noted that the Ni–O bonds adjacent to the fault substantially elongate even after one SCF step, as can be seen in the second illustration, together with significant stabilization of the lattice energy (see Supplementary Fig. 4 for the plot of the lattice energy variation with SCF steps). The third illustration shows the finally optimized configuration of the supercell cell, revealing that the Ni–O bond length adjacent to the fault plane is 2.1 Å in the magnified illustration. This value after geometry optimization is in fairly good agreement with the experimentally observed Ni–O length, 2.2 Å, as directly compared in the ABF-STEM image. The DFT calculation thus confirms that elongation of the Ni–O bond length at the fault is an energetically stable configuration. At the same time, La³⁺ cations are also observed to displace along the z axis out of the fault plane in the series of supercell illustrations in order to facilitate the reduction of the repulsion at the fault.

We also carried out another set of DFT calculations to compare the density of states (DOS) of the Ni 3d and O 2p orbitals in LaNiO₃ with and without a fault. As demonstrated in Supplementary Fig. 5, the overall substantial degree of overlap between the Ni 3d and O 2p states in addition to the metallic behavior with no bandgap does not significantly vary between the two DOS plots. However, as indicated by arrows, the noticeable increases of O 2p and Ni 3d states at the bottom, in the middle, and at the top of the band in the DOS plot are key changes of the...
by the oxygen displacements toward the fault plane, as denoted by small arrows in the illustrations. Good agreement of the Ni\[NiO2\] DFT calculation and the direct ABF-STEM observation is verified. LaCoO3 and LaNiO3 is noted to contain strong electrostatic repulsion at the fault.

Electronic structure that result from the z axis elongation of [NiO2]\[NiO2\] octahedra at the fault plane. When six oxygen ligands around a metal cation are located in an octahedral symmetry, the metal 3d states are discreetly split as two degenerate t2g and e\[e\] levels. In contrast, if this symmetry is broken either by oxygen octahedral elongation or via other perturbation\[27, 28, 32, 33\], the degeneracy of the two levels is removed, resulting in a wide range of energy levels of the metal 3d states in addition to the O 2p states for hybridization.

EELS analyses. Atomic-column resolved EELS analyses on the RP shear faults were performed to examine the variation of energy-loss spectra in addition to the oxidation state of transition metals. As the Ni-L3 edge seriously overlaps with the La-M4 edge, we selected LaCoO3 and SrTiO3 samples containing a shear fault instead of LaNiO3 for the EELS analyses. Figure 4 shows a comparison of Co-L and O-K edges acquired from the bulk and the fault in LaCoO3. First, no noticeable changes of the peak intensity and position in the Co-L2,3 edges from the bulk and the fault were identified, directly indicating the oxidation state of Co3+ is invariant at the fault. Second, the three major peaks in the O-K edge are represented for the hybridizations of O 2p with Co 3d (peak A), La 5d (peak B) and Co 4s/4p (peak C), respectively.\[34–37\]. The intensity of peak A obtained from the fault substantially diminishes, as denoted by a red arrow in the plot, while the intensity of the other peaks does not significantly vary between the bulk and the fault. This reduction of the pre-peak in the O-K edge appears to consistently correlate with the elongation of the Co–O bond length at the faults. Because the bond distance between Co and O becomes longer, the effect of oxygen ligands on Co should be reduced, and thereby the degree of hybridization of O 2p with Co 3d orbitals decreases. Consequently, as demonstrated in a perovskite cobaltate\[35\] and layered lithium transition-metal oxides\[38, 39\], which contain oxygen vacancies, the

**Fig. 3 Description of sublayer polarity in perovskites and geometry optimization of atom configuration during DFT calculation.** The effective charges of [AX] and [BX2] sublayers in the five perovskites are indicated. a No polarity of sublayers is present in SrTiO3, BaCeO3, and CsPbBr3. b In contrast, [LaO]+ in LaCoO3 and LaNiO3 is noted to contain strong electrostatic repulsion at the fault. c Geometry optimization with SCF steps at the fault in LaNiO3 is achieved by the oxygen displacements toward the fault plane, as denoted by small arrows in the illustrations. Good agreement of the Ni–O elongation obtained by the DFT calculation and the direct ABF-STEM observation is verified.
lower intensity of peak A likely stems from the comparatively weak orbital interaction. Indeed, such a peak reduction was not observable in the O-K edge obtained from the fault in SrTiO₃, where no elongation of Ti–O bond length is identified at the faults (see Supplementary Fig. 6 for details on the EELS analysis of the fault in SrTiO₃).

**Local structures of RP oxides.** In addition to directly demonstrating the distortion of oxygen octahedra at the faults in perovskite oxides, the findings in this work have further notable implications for the in-depth understanding of the structures of a variety of RP oxides and halides, which exhibit a very wide range of physical properties encompassing dielectric and unusual ferroic, multiferroic, electrocatalytic, and optical behavior. As can be easily recognized from their general formula, Aₙ₊₁BₓXₙ₋₁ or equivalently AX[ABX₃]ₙ, the presence of consecutive [AX] sublayers is a common structural feature irrespective of the value of n. In particular, it is noted that every anion octahedron in layer-structured A₂BX₄ (n = 1) (or AX[ABX₃]) is in contact with two shear planes, as denoted by yellow broken lines in the atomic illustration of Sr₂TiO₄ in Fig. 5a. If the [AX] layer has an effective local charge, as aforementioned in [LaO]⁺, charge-induced electrostatic repulsion would considerably accumulate over the entire crystal lattice of A₂BX₄. Consequently, substantial elongation of [BX₆] octahedra should take place to significantly reduce the repulsive instability in the consecutive [AX] layers.

We examined more than 80 RP-type oxides and halides with high symmetry from the open database of inorganic materials (the Materials Project) and the Inorganic Crystal Structure Database (ICSD) to scrutinize whether the local net charge of [AX] sublayers exerts a general influence on the tetragonal distortion of anion octahedra in RP phases. Note that most of the RP oxides and halides under our examination in this work were confined to centrosymmetric phases with a single composition at each of the A and B sites in order to exclude any other complexity originating from the polarity induced by either cation off-centering from the second-order Jahn–Teller effect, octahedral rotations, or cation ordering. Taking this into consideration, structure comparisons were made largely between RP phases with a space group, H4mmm.

Supplementary Tables 1–3 list all the structural information regarding the ratios of x axis and z axis B–X bond lengths, Lₓ/Lₚ, in 87 RP oxides and halides (see the detailed illustration in Supplementary Table 1 for the definition of Lₓ/Lₚ). Ca-, Sr-, and Ba-based RP oxides with no local charge in Supplementary Table 1, K- and Cs-based RP halides in Supplementary Table S2, and lanthanide-based RP oxides in Supplementary Table 3. The series of tables and supplementary figures demonstrates the notable and exclusive effect of the [AX] local charge on the z axis elongation of octahedra (Lₓ/Lₚ > 1.04) in the cases of lanthanide-based (A = La, Pr, Nd, Eu, and Gd) RP oxides. We set the criterion of 4% elongation by rounding down the minimum value, 1.048 (which corresponds to 4.8% elongation in the case of EuCOO₃ in Sr₂SnO₄, as listed in Supplementary Table 1. Because these RP oxides have no effective charge in [SrO] sublayers, no distortion of oxygen octahedra is identified in each structure. Magnified ABF images directly provide quantitative information on the Ti–O and Sn–O bond lengths.

**Fig. 5** STEM images showing the structure of Sr-based RP oxides. Atomic-column-resolved images were acquired in a Sr₂TiO₄ and Sr₃Ti₂O₇ and b Sr₂SnO₄ as listed in Supplementary Table 1. Because these RP oxides have no effective charge in [SrO] sublayers, no distortion of oxygen octahedra is identified in each structure. Magnified ABF images directly provide quantitative information on the Ti–O and Sn–O bond lengths.
Supplementary Table 3), to three decimal places. As denoted by the red-brown background in Supplementary Table 3, all the Ln-based RP oxides exhibit more than 4% substantial elongation of z axis B–O bond lengths, showing strong tetragonal distortion of octahedra (see Supplementary Figs. 46–60 for their crystal structures).

**Direct observations in RP oxides.** To experimentally visualize the local-charge-induced elongation of oxygen octahedra in addition to referencing the structural database, we carried out direct atomic-column-resolved ABF-STEM observations in a variety of RP oxides selected from Supplementary Tables 1 and 3. Figure 5 exemplifies ABF images and their enlargements of three Sr-based RP oxides, SrTiO₃, Sr₂TiO₇, and SrSnO₃ (see Supplementary Figs. 61 for low-magnification STEM images), with no local charge at the fault plane of [SrO]₁₀−[SrO]₁₀ consecutive layers, as listed in Supplementary Table 1. The nearly identical x axis and z axis Ti–O (and Sn–O) bond lengths (Lₓ and Lₚ) in each oxide agree well with the information provided in the table, verifying the ratio Lₓ/Lₚ = 1.

Figure 6 shows the actual structures and their atomic-scale ABF images of five lanthanide-based RP nicklelates, La₃NiO₄, La₃NiO₅O, La₄NiO₅O, Nd₃NiO₄, and Pr₂NiO₄, selected from Supplementary Table 3, with an effective local charge of the sublayers, [LaO]⁺, [NdO]⁺, and [PrO]⁺, as schematically illustrated in the left-hand column (see Supplementary Figs. 62 for low-magnification STEM images). In particular, Fig. 6b, c specifically exemplify the tetragonal elongation of every oxygen octahedra in Nd₃NiO₄ and Pr₂NiO₄ in addition to La₃NiO₅O in Fig. 6a. The Ni–O bond lengths in each RP nicklelate can be directly acquired from the ABF images in the right-hand column. As denoted by yellow arrows in the enlargements, the substantial elongation (more than 15%) of z axis Ni–O bonds toward the fault planes is readily recognized. This structural feature is consistently confirmed in La-based RP cobaltates and cuprates, as demonstrated in Fig. 7. Moreover, Cu²⁺ is a well-known strong Jahn–Teller cation. As a consequence, more prominent z axis elongation (>27%) of [CuO₆] octahedra in La₂CuO₄ (Fig. 7b) than that of [CoO₆] in La₃CoO₅ and [NiO₆] in La₄NiO₅O is identified by a combination of the [LaO]⁺ local charge and the Jahn–Teller effect. To clarify that this >27% strong elongation of [CuO₆] octahedra is a common aspect in Pr₂CuO₄ and Nd₂CuO₄ as well (Supplementary Table 3), an additional structure comparison is provided in Supplementary Fig. 63.

We summarize several representative A₂BX₄ (n = 1) oxides and fluorides among the RP phases with the space group I4/mmm in Supplementary Fig. 64 to highlight the impact of the net charge of the sublayers on the significant elongation of oxygen octahedra. As schematically described in Supplementary Fig. 64a, [SrO] in Sr₃TiO₄ [KF] in K₂NiF₃, and [CF] in Cs₂CuF₄ have no effective charge. The six B–X bond lengths in each of {TiO₆}, {NiF₆}, and {CuF₆} octahedra are thus nearly identical (<4% variation), showing no substantial geometric distortion. In contrast, as the [LaO]⁺ sublayers in all the La-based RP oxides are positively charged, the protrusion of oxygen anions toward the shear planes is identified and this reduces the repulsive electrostatic instability. Even though B-site cations are the same (Ni²⁺ in K₂NiF₃ and La₂NiO₄ and Cu²⁺ in Cs₂CuF₄ and La₂CuO₄, respectively), the distinct elongation of [NiO₆] in La₂NiO₄ and [CuO₆] in La₂CuO₄ in contrast to [NiF₆] in K₂NiF₃ and [CuF₆] in Cs₂CuF₄ clarifies the crucial influence of the electrostatic charge on the tetragonal distortion of oxygen octahedra in RP oxides. Similarly, because the other lanthanide cations, Ln = Pr, Nd, Gd, at the A sites in RP oxides are trivalent, the [LnO]⁺ sublayers are also positively charged, as in the case of the La-based RP oxides. As specifically exemplified in Supplementary Fig. 64c for Pr- and Nd-based cobaltates, therefore, substantial z axis octahedral elongation is consistently identified.

When the structure of A-site solid solution (A′A″)₃B₂O₆, (where the valence states of A′ and A″ are 3+ and 2+, respectively) is scrutinized, the effect of electrostatic charge on the elongation of oxygen octahedra is further supported. Figure 8 shows the STEM images of (A′A″)₃B₂O₆-type (La,Sr)₂AlO₄, (La,Sr)₂GaO₄, and (Nd,Ca)₂AlO₄ (see Supplementary Fig. 65 for composition verification of A-site solid solutions). As schematically depicted in the illustrations in the left-hand column, the A-site sublayers of these RP oxides encompass [SrO]₁₀−[SrO]₁₀ with no local charge in addition to charged [LaO]⁺ (or [NdO]⁺) as a solid solution. Consequently, the positive-charge electrostatics between the consecutive shear layers should diminish, resulting in comparatively weak interlayer repulsion. Considerably strong elongation of oxygen octahedra for screening the effective positive charge at the fault is thus no longer necessary. Indeed, as denoted in each of the magnified ABF images in the right-hand column, the length of z axis Al–O and Ga–O bonds does not exceed 2.2 Å, in contrast to a value of 2.3 Å in some nicklelates (Fig. 6) and cobaltates (Fig. 7a) and even 2.5 Å in cuprates (Fig. 7b). Therefore, this series of direct observations in Fig. 8 provides consistent evidence for our findings on the electrostatics-induced distortion of oxygen octahedral in RP oxides.

In conclusion, we have observed the formation of [AX]−[AX] interleaved shear faults as common planar defects in five ABX₂ perovskite-type phases, SrTiO₃, BaCeO₃, LaCoO₃, LaNiO₃, and CsPbBr₃. As each fault generated in LaCoO₃ and LaNiO₃ consists of two positively charged [LaO]⁺ sublayers in contrast to [AX] with no local charge in SrTiO₃, BaCeO₃, and CsPbBr₃, strong electrostatic repulsion can be established at the fault plane. We identified that this charge-induced instability could remarkably diminish via the asymmetric tetragonal distortion of oxygen octahedra by the protonation of oxygen anions toward the fault plane and thereby effectively screen the repulsion. Furthermore, via direct atomic-column-resolved visualization by ABF imaging in 14 representative RP oxides, our atomic-level findings regarding the tetragonal distortion of oxygen octahedra at the shear faults provide crucial insight to consistently explain the octahedron elongation commonly observed in lanthanide-based RP oxides among more than 80 RP-type oxides and halides. In addition to the electronic structure of metal cations on the basis of the Jahn–Teller effect, the present study suggests that the local electrostatic instability is a significant cause of the tetragonal distortion of oxygen octahedra in complex oxides.

**Methods**

**Preparation of polycrystalline samples.** To prepare polycrystalline perovskite and RP phase samples, powders were first synthesized via a conventional solid-state reaction method using SrCO₃ (99.9%, Sigma Aldrich) and TiO₂ (99.9%, Sigma Aldrich) for Sr₃TiO₇, SrTiO₃, Sr₃TiO₇, and SrCO₃ and SnO₂ (99.9%, Sigma Aldrich) for SrSnO₃; BaCO₃ (99.9%, Sigma Aldrich) and Co₃O₄ (99.9%, Sigma Aldrich) for Pr₂NiO₄; BaCO₃ (99.9%, Sigma Aldrich) and Co₃O₄ (99.9%, Sigma Aldrich) for BaCoO₂; La(OH)₃ (99.9%, Sigma Aldrich) and NIO (99.9%, Sigma Aldrich) for LaNiO₃ and LaNiO₅O; La(OH)₂ and Co₂O₃ (99.5%, Sigma Aldrich) for LaCoO₃; La(OH)₂ and Cu₂O (99.9%, Sigma Aldrich) for LaCuO₃; Nd₂NiO₄ and Pr(NO₃)₆·6H₂O (99.9%, Sigma Aldrich) and Ni(CH₃COO)₂·4H₂O (99.9%, Sigma Aldrich) for Pr₂NiO₄. Each mixture of the starting materials with 5%-excess A-site cations for perovskites and stoichiometric composition for RP phases was ball-milled in high-purity ethyl alcohol for 24 h with a zirconia jar and balls. After being dried, the slurries were calcined in air at 1100 °C for 10 h for SrTiO₃, at 1200 °C for 10 h for Sr₃TiO₇, and at 1200 °C for 10 h for Sr₃TiO₇ and Sr₃TiO₄. SrSnO₃ at 1050 °C for 10 h for SrSnO₃; at 1050 °C for 10 h for BaCeO₃; at 1050 °C for 10 h for BaCeO₃; at 1200 °C in an O₂ flow atmosphere for 24 h for LaCoO₃; at 1200 °C in an O₂ flow atmosphere for 24 h for LaCoO₃; at 1200 °C in an O₂ flow atmosphere for 24 h for SrSnO₃. The calcined samples were ground to reduce the grain size to below 2 µm. 10% Gd was doped in BaCeO₃ to improve its sinterability and a dense microstructure. After grinding, the powders were cold isostatically pressed in a stainless steel die to form pellets, which were then sintered at 1450 °C for SrTiO₃ and SrSnO₃ and at 1300 °C for BaCeO₃, LaCoO₃, and Pr₂NiO₄. The sintered samples were ground to 2 µm and re-pressed to form pellets for the STEM measurements.
powders were ball-milled again to finally obtain fine particles. X-ray diffractometry (D/MAX-2500, Rigaku) with Cu-Kα radiation confirmed that single-phase powders were synthesized. Each of the powders was slightly pressed into disks of 10 mm diameter and isostatically pressed under 200 MPa. The pellets were sintered in air at 1400 °C for 10 h for SrTiO3, Sr2TiO4, and Sr3Ti2O7; at 1550 °C for 12 h for Sr2SrO; at 1400 °C for 5 h for BaCeO3 and LaCoO3; at 1300 °C for 2 h for Nd2CuO4; at 1400 °C for 2 h for Nd2NiO4 and Pr2NiO4; and at 1350 °C for 2 h for La3Ni2O7 in an O2-flow atmosphere to obtain dense polycrystalline microstructure.

Thin-film fabrication. Epitaxial LaNiO3, La3Ni2O7, and La4Ni3O10, Nd2NiO4, and Pr2NiO4, as listed in Supplementary Table 3. Strong local repulsion between [LnO]+ sublayers is illustrated in the left-hand column. Significant elongation of the Ni–O bonds at the shear planes is denoted by yellow arrows in the ABF images, providing quantitative information on the bond lengths. Red arrows in each image indicate substantial displacement of oxygen anions.

**Fig. 6 STEM images showing the structure of lanthanide-based RP nickelates.** Atomic-column-resolved images were acquired in a La2NiO4, La3Ni2O7, and La4Ni3O10, b Nd2NiO4, and c Pr2NiO4, as listed in Supplementary Table 3. Strong local repulsion between [LnO]+ sublayers is illustrated in the left-hand column. Significant elongation of the Ni–O bonds at the shear planes is denoted by yellow arrows in the ABF images, providing quantitative information on the bond lengths. Red arrows in each image indicate substantial displacement of oxygen anions.
by adding 3 mol% more of the La nitrate for LaNiO₃ films. For complete dissolution of the source materials, refluxing was carried out at 80 °C for 1 h. Each of the precursor solutions was deposited on (001) LaAlO₃ and (001) SrTiO₃ (Shinkosha, Japan) single-crystal substrates by a spin-coating method at 5000 rpm for 10 s. The wet films were dried at 150 °C for 10 min on a hot plate, and then annealed at 800 °C for 1 h in the air for LaNiO₃ films, at 900 °C for 1 h in the air for La₂NiO₄ films, and at 1000 °C for 1 h in N₂-flow atmosphere for La₂CoO₄, and La₂CuO₄ films for crystallization. X-ray diffractometry (X'Pert-PRO MRD, PANalytical) and STEM observation verified the epitaxy of grown films. Commercially available (001) single-crystal substrates (MTI corporation) were purchased for structure observation in La₃SrAlO₄, La₃SrGaO₄, and NdGaAlO₄.

**Fig. 7 STEM images showing the structure of La-based RP oxides.** Atomic-column-resolved images were acquired in a La₂CoO₄ and La₂CoO₄O₂ and b La₂CuO₄ as listed in Supplementary Table 3. Strong local repulsion between [LaO]⁺ sublayers is illustrated in the left-hand column. Significant elongation of the Co–O and Cu–O bonds at the shear planes is denoted by yellow arrows in the ABF images, providing quantitative information on the bond lengths. As Cu²⁺ is known as a strong Jahn–Teller cation, {CuO₆} octahedra in La₂CuO₄ show more prominent z axis elongation. Red arrows in each image indicate substantial displacement of oxygen anions.

**Nanocrystal synthesis.** CaPbBr₂ nanocrystals were synthesized by a microwave-assisted method²⁰. A solvent mixture was prepared using 1-octadecene (5 mL), oleylamine (0.5 mL), and oleic acid (0.5 mL). This mixture was degassed by evacuation and stirred in a round bottom flask at 170 °C on a hot plate for 30 min. After the degassing process, the mixture was transferred into a 10-mL vial and followed by adding and dispersing Cs₂CO₃ (0.0217 g, 99.995%, Sigma Aldrich) and PbBr₂ (0.0738 g, 99.999%, Sigma Aldrich) with a vortex mixer. This vial was placed into a microwave oven and subsequent microwave irradiation was carried out at 700 W for 5 min. The 5-min irradiation together with a 1-min intermission was further repeated eight more times, resulting in irradiation for 45 min in total. To separate residual solvents and a solid product, the mixture after the microwave irradiation was centrifuged at 1398×g for 5 min. The solid product was dispersed in toluene and further purified by centrifugation at 5590×g for 5 min so as to finally acquire a toluene solution containing homogeneously dispersed CaPbBr₂ nanocrystals.

**STEM, EELS, and specimen preparation.** Specimens for STEM observation of sintered polycrystalline samples were prepared by mechanical grinding to a thickness of 80 µm, dimpling to a thickness of <10 µm, and finally ion-beam thinning by using a precision ion polishing system (PIPS, Gatan Inc.) for electron transparency. STEM specimens for LaNiO₃ and NdNiO₃ thin films were prepared by lift out via ion-beam milling in a focused ion-beam system (Quanta 3D FEG and Helios G4 UX, ThermoFisher Scientific). Protective amorphous carbon and thin Pt layers were applied over the region of interest before milling. To minimize the sidewall damage and sufficiently thin the specimen for electron transparency, final milling was carried out at a voltage of ~2 kV. HAADF- and ABF-STEM images were acquired with a transmission electron microscope (Titan cубс g2 60–300, ThermoFisher Scientific) at 200 kV or 300 kV with a spherical aberration (Cs) corretcor (CEOS GmbH). The optimum size of the electron probe was ~1 Å with a convergence semiangle of 53.6 (or 70.0) mrad. The collection semiangle of the STEM detectors was set to 40.0–200 mrad for HAADF imaging and 9.61–53.6 (or 10.1–19.1) mrad for ABF imaging. EELS analysis was performed with a Gatan Image Filter (GIF Quantum 965, Gatan Inc.). Electron energy-loss spectra for the Ba-M₄ (781 eV) and Ce-M₃ (883 eV) edges were acquired for atomic-column spectrum imaging in BaCoO₃ and with a dispersion of 0.25 eV per channel and a collection aperture of 5 mm in diameter. The O-K, Ti-L₂,3, and Co-L₂,3 edges were also collected in SrTiO₃ and LaCoO₃ samples containing shear faults to examine whether the oxidation states of Ti and Co and the orbital hybridization with oxygen vary at the fault plane.

**DFT calculations.** Ab initio DFT calculations for DOS variation in LaNiO₃ were carried out using the spin-polarized local density approximation (LDA) functional for exchange-correlation, along with the ultrasoft pseudopotentials for ionic cores, as implemented in the CASTEP code (Biovia Inc.). To consider the electron localization near Ni ions, the LDA + U method with the Hubbard U parameter (U = 4.0 eV for Ni 3d states) was employed. A low-spin (t¹₂g⁴e₅g) for d⁵ Ni³⁺ configuration was assumed for the spin-polarized calculations. The plane-wave basis set for the kinetic energy cutoff was 500 eV. Relaxation of the internal coordinates for each atom was performed using the Broyden–Fletcher–Goldfarb–Shanno (BFGS) algorithm with convergence tolerances of 0.1 eV/Å for the maximum ionic force, 5 × 10⁻⁶ eV/atom for the total energy, and 0.005 Å for the maximum ionic displacement.
(La,Sr)2GaO4, and repulsion between the layers diminishes. Therefore, the elongation of the Al–O bonds toward the shear planes is not remarkable, showing a bond length of ≤2.2 Å. Magnified ABF images directly provide quantitative information on the bond lengths. Red arrows in each image indicate a slight displacement of oxygen anions.

**Data availability**
The data sets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Received: 7 December 2020; Accepted: 3 September 2021; Published online: 20 September 2021

**References**
1. Pauling, L. The principles determining the structure of complex ionic crystals. *J. Am. Chem. Soc.* **51**, 1010–1026 (1929).
2. Glazer, A. M. The classification of tilted octahedra in perovskites. *Acta Crystallogr.* **B 28**, 3384–3392 (1972).
3. Woodward, P. M. Octahedral tilting in perovskites. I. Geometrical considerations. *Acta Crystallogr.* **B 53**, 32–43 (1997).
4. Glazer, A. M. Inversion symmetry breaking by oxygen octahedral rotations in the Ruddlesden–Popper NaRTiOx family. *Phys. Rev. Lett.* **112**, 187602 (2014).
5. Pitcher, M. J. et al. Tilt engineering of spontaneous polarization and magnetization above 300 K in a bulk layered perovskite. *Science* **347**, 420–424 (2015).
6. Kim, T. H. et al. Polar metals by geometric design. *Nature* **533**, 68–72 (2016).
7. Lu, X.-Z. & Rondinelli, J. M. Epitaxial-strain-induced polar-to-nonpolar transitions in layered oxides. *Nat. Mater.* **15**, 951–953 (2016).
8. Zhu, T. et al. Cation exchange as a mechanism to engineer polarity in layered perovskites. *Chem. Mater.* **30**, 8915–8924 (2018).
9. Yoshida, S. et al. Hybrid improper ferroelectricity in (Sr,Ca)3Sn3O9 and beyond: universal relationship between ferroelectric transition temperature and tolerance factor in \( n = 2 \) Ruddlesden–Popper phases. *J. Am. Chem. Soc.* **140**, 15690–15700 (2018).
10. Zhang, Y., Wang, J. & Ghosez, P. Unraveling the suppression of oxygen octahedral rotations in \( A \cdot B \cdot O \) Ruddlesden–Popper compounds: engineering multiferroicity and beyond. *Phys. Rev. Lett.* **125**, 157601 (2020).
11. Bersuker, I. B. *The Jahn–Teller Effect* (Cambridge University Press, New York, 2006).
12. Halcrow, M. A. Jahn–Teller distortion in transition metal compounds, and their importance in functional molecular and inorganic materials. *Chem. Soc. Rev.* **42**, 1784–1795 (2013).
13. Bak, J., Bae, H. B., Kim, J., Oh, J. & Chung, S.-Y. Formation of two-dimensional homologous faults and oxygen electrocatalytic activities in a perovskite nickelate. *Nano Lett.* **17**, 3126–3132 (2017).
14. Wang, Z. et al. Dimensionality-driven insulator–metal transition in A-site excess non-stoichiometric perovskites. *Nat. Commun.* **1**, 106 (2010).
15. Arredondo, M. et al. Chemistry of Ruddlesden–Popper planar faults at a ferroelectric–ferromagnet perovskite interface. *J. Appl. Phys.* **109**, 084101 (2011).
16. Third, A. S. et al. Atomic structure and electrical activity of grain boundaries and Ruddlesden–Popper faults in cesium lead bromide perovskite. *Adv. Mater.* **31**, 1805047 (2019).
17. Guo, Q., Farokhipoor, S., Magín, C., Rivadulla, F. & Noheda, B. Tunable resistivity exponents in the metallic phase of epitaxial nickelates. *Nat. Commun.* **11**, 2949 (2020).
18. Bak, J., Bae, H. B., Oh, C., Son, J. & Chung, S.-Y. Effect of lattice strain on the formation of Ruddlesden–Popper faults in heteroepitaxial LaNiO3 for oxygen evolution electrocatalysis. *J. Phys. Chem. Lett.* **11**, 7253–7260 (2020).
19. Choi, S.-Y., Chung, S.-Y., Yamamoto, T. & Ikuhara, Y. Direct determination of dopant site selectivity in ordered perovskite CaCu$_3$Ti$_4$O$_{12}$ polycrystals by aberration-corrected STEM. 
Adv. Mater. 21, 885–889 (2009).

20. Chung, S.-Y., Choi, S.-Y., Kim, T. H. & Lee, S. Surface-orientation-dependent distribution of subsurface cation-exchange defects in olivine-phosphate nanocrystals. 
ACS Nano 9, 850–859 (2015).

21. Yoon, H.-I. et al. Probing dopant segregation in distinct cation sites at perovskite oxide polycrystal interfaces. 
Nat. Commun. 8, 1417 (2017).

22. Ishikawa, R. et al. Direct imaging of hydrogen-atom columns in a crystal by annular bright-field electron microscopy. 
Nat. Mater. 10, 278–281 (2011).

23. Findlay, S. D., Azuma, S., Shibata, N., Okunishi, E. & Ikuhara, Y. Direct oxygen imaging within a ceramic interface with some observations upon the dark contrast at the grain boundary. 
Ultramicroscopy 111, 285–289 (2011).

24. Kim, H.-S., Jung, A.; Choi, S.-Y.; Jung, W. & Chung, S.-Y. Vacancy-induced electronic structure variation of acceptors and correlation with proton conduction in perovskite oxides. 
Angew. Chem. Int. Ed. 55, 13499–13503 (2016).

25. Kim, J. H. et al. Atomic-scale observation of oxygen substitution and its correlation with hole-transport barriers in Cu$_3$ZnSnSe$_6$ thin-film solar cells. 
Adv. Energy Mater. 6, 1501902 (2016).

26. Byeon, P. et al. Atomic-scale observation of LiFePO$_4$ and LiCoO$_2$ dissolution behavior in aqueous solutions. 
Adv. Funct. Mater. 28, 1804564 (2018).

27. Heo, Y. et al. Symmetry-broken atomic configurations at grain boundaries and oxygen evolution electrocatalysis in perovskite oxides. 
Adv. Energy Mater. 8, 1802481 (2018).

28. Bak, I., Bae, H. B. & Chung, S.-Y. Atomic-scale perturbation of oxygen octahedra via surface ion exchange in perovskite nickelates boosts water oxidation. 
Nat. Commun. 10, 2713 (2019).

29. Byeon, P. et al. Atomic-scale unvieling multiphase evolution during hydrates Zn-ion insertion in vanadium oxide. 
Nat. Commun. 12, 4599 (2021).

30. Ohtomo, A. & Hwang, H. Y. A high-mobility electron gas at the LaAlO$_3$/ SrTiO$_3$ superinterface. 
Nature 427, 423–426 (2004).

31. Nakagawa, N., Hwang, H. Y. & Muller, D. A. Why some interfaces cannot be sharp. 
Nat. Mater. 5, 204–209 (2006).

32. Wang, L. et al. Strain Effect on oxygen evolution reaction activity of epitaxial NdNiO$_3$ thin films. 
ACS Appl. Mater. Interfaces 11, 12941–12947 (2019).

33. Bak, J., Heo, Y., Yun, T. G. & Chung, S.-Y. Atomic-level manipulation in oxide and alloys for electrocatalysis of oxygen evolution and reduction. 
ACS Nano 14, 14323–14354 (2020).

34. Shah, A. B. et al. Probing interfacial electronic structures in atomic layer LaMnO$_3$ and SrTiO$_3$ superlattices. 
Adv. Mater. 22, 1156–1160 (2010).

35. Gazquez, J. et al. Atomic-resolution imaging of spin-state superlattices in nanopockets within cobaltite thin films. 
Nano Lett. 11, 973–976 (2011).

36. Zhu, G.-Z., Radtke, G. & Botton, G. A. Bonding and structure of a reconstructed (001) surface of SrTiO$_3$ from TEM. 
Nature 490, 384–387 (2012).

37. Song, C. W., Suh, H., Bak, J., Bae, H. B. & Chung, S.-Y. Dissolution-induced surface roughening and oxygen evolution electrocatalysis of alkali-earth iridates in acid. 
Chem. 5, 3243–3259 (2019).

38. Shakla, A. K. et al. Unravelling structural ambiguities in lithium- and manganese-rich transition metal oxides. 
Nat. Commun. 6, 8711 (2015).

39. Li, L. et al. Hidden subsurface reconstruction and its atomic origins in layered oxide cathodes. 
Nano Lett. 20, 2756–2762 (2020).

40. Lee, C.-H. et al. Exploiting dimensionality and defect mitigation to create tunable microwave dielectrics. 
Nature 502, 532–536 (2013).

41. Lee, Y.-L. et al. Kinetics of oxygen surface exchange on epitaxial Ruddlesden–Popper phases and correlations to first-principles descriptors. 
J. Phys. Chem. Chem. Phys. 7, 244–249 (2016).

42. Gu, X.-K. & Nikolla, E. Design of Ruddlesden–Popper oxides with optimal surface oxygen exchange properties for oxygen reduction and evolution. 
ACS Catal. 7, 5912–5920 (2017).

43. Meyer, T. L. et al. Strain control of oxygen kinetics in the Ruddlesden–Popper oxide La$_{1.5}$Sr$_{2.5}$CuO$_4$. 
Nat. Commun. 9, 92 (2018).

44. Zhu, Y. et al. Unusual synergistic effect in layered Ruddlesden–Popper oxides enables ultrafast hydrogen evolution. 
Nat. Commun. 10, 149 (2019).

45. Liu, S., Sun, C., Chen, J., Xiao, J. & Luo, J.-L. A high-performance Ruddlesden–Popper perovskite for bifunctional oxygen electrocatalysis. 
ACS Catal. 10, 13437–13444 (2020).

46. Cohen, R. E. Origin of ferroelectricity in perovskite oxides. 
Nature 358, 136–138 (1992).

47. Bersuker, I. B. Pseudo-Jahn–Teller effect—a two-state paradigm in formation, deformation, and transformation of molecular systems and solids. 
Chem. Rev. 113, 1351–1390 (2013).

48. Akamatsu, H. et al. A-site cation size effect on oxygen octahedral rotations in acentric Ruddlesden–Popper alkali rare-earth titanates. 
Phys. Rev. Mater. 3, 065001 (2019).

49. Balachandran, P. V., Puggioni, D. & Rondinelli, J. M. Crystal-chemistry guidelines for noncentrosymmetric A$_2$BO$_4$ Ruddlesden–Popper oxides. 
Inorg. Chem. 53, 336–348 (2014).

50. Long, Z., Ren, H., Sun, J., Ouyang, J. & Na, N. High-throughput and tunable synthesis of colloidal CsPbX$_3$ perovskite nanocrystals in a heterogeneous system by microwave irradiation. 
Chem. Commun. 53, 9914–9917 (2017).

Acknowledgements
This work was supported by the National Research Foundation of Korea (NRF), grant numbers 201M3H4A101079300 and 2021R1A2C2006072. S.-Y.C. was also financially supported by the Samsung Research Funding Centre of Samsung Electronics, grant no. SRFC-MA1602-52.

Author contributions
S.-Y.C. conceived and initiated the project, carried out the DFT calculations, assisted the STEM analyses, and wrote the paper. Y.Hong carried out all the syntheses and specimen preparation, performed the STEM image analyses, and collected the structure data of oxides. P.B. and H.B.B. performed the STEM, EELS and EDS analyses and contributed to the acquisition of image data. J.B., Y.Heo and H.-S.K. supported the experiments and specimen preparation. All the authors discussed the results and commented on the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-25889-6.

Correspondence and requests for materials should be addressed to Sung-Yoon Chung.

Peer review information Nature Communications thanks Rohan Mishra and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.