Materials Advances

PAPER

Cite this: *Mater. Adv.*, 2021, 2, 2398

Received 9th February 2021, Accepted 21st February 2021

DOI: 10.1039/d1ma00122a

rsc.li/materials-advances

Introduction

Oxynitride materials have attracted recent interest for use in applications such as pigments, dielectrics, and optoelectronic materials.1 The overlap between the band gaps of many oxynitride materials and the solar spectrum also makes them potential water splitting photocatalysts that can operate in the visible-light region.² Several niobium oxynitride perovskites, for instance, are promising candidate semiconductors for solar energy conversion owing to their wide visible light absorption.³⁻⁶

However, niobium oxynitrides are known to be more prone to defect formation during their synthesis via nitridation, compared to other perovskite oxynitrides (e.g. tantalum oxynitrides). This is a feature, rather than a flaw. Structural defects, such as zero-dimensional defects (e.g. vacancy, interstitial and

Anion ordering and vacancy defects in niobium perovskite oxynitrides†

Joshua J. Brown,^a Youxiang Shao,^b Zhuofeng Ke D^b and Alister J. Page D^{*a}

Niobium perovskite oxynitrides are emerging as promising semiconductor materials for solar energy conversion processes, due to their physical properties and amenability to defect engineering. However, defect engineering in mixed-anion semiconductors such as perovskite oxynitrides is generally hindered by the absence of long-range order in the crystal lattice, a phenomenon known as anion-ordering. We demonstrate how anion ordering influences the stability and mobility of point defects in two exemplar perovskite oxynitrides, BaNbO₂N and LaNbON₂. Accurate first-principles calculations show that fully cis -anion orderings in BaNbO₂N are more stable than fully trans-anion orderings, whereas anion orderings with mixed dimensionality may be more prevalent in the lower-symmetry LaNbON2. Anion ordering in LaNbON₂ is also influenced by a pronounced A-site coordination sphere effect not observed in BaNbO₂N, whereby local La- $(O,N)_{12}$ coordination environments give rise to alternating LaO and LaN layers in the bulk material. Anion order was predicted to effect the redistribution of electrons upon anion vacancy creation to the cation sublattice. Diffusion barriers for O^{2-} vacancies in trans-ordered BaNbO₂N were found to be lower than those for N^{3-} vacancies, suggesting that stabilising transordered phases of BaNbO₂N will yield more effective retention of nitrogen content in this material. The reverse is the case for LaNbON₂, with N^{3-} vacancy defects exhibiting more facile diffusion than O^{2-} vacancy defects. We believe these insights will aid the emergent understanding of defect engineering in mixed-anion perovskite oxynitride semiconductors, and specifically help facilitate strategies for stabilising their nitrogen content. PAPER
 **Example 18 Constrained and Society and Vacancy defects in niobium

Constants Mark 2021

Consider Adv. 2021

Consider Adv. 2021

Consider Adv. 2021

Consider Adv. 2021

Consider a conversion process due to their sh**

anti-site defects) and higher-dimensional defects (dislocations, grain boundaries, and volumetric defects) provide design parameters for modulating surface active sites, charge carrier mobilities, and band edge positions.^{7,8} Of these, zero-dimensional point defects have the strongest link to material non-stoichiometry.⁷ Niobium oxynitrides are therefore ideal candidates for defect engineering. For instance, the enhanced conductivity of $BaNbO₂N$ compared to $BaTaO₂N$ is suspected to arise from defect concentration in the former.⁹ Niobium oxynitrides also have larger absorption tails extending in the visible region of the spectrum, which is indicative of nitrogen vacancies and reduced B-site cations, compared to their tantalum counterparts.⁹

However, engineering structural defects in perovskite oxynitride photocatalysts, such as $AM(O,N)_3 (M = Ti, Zr, Hf, Ta, Nb)$, in a controlled manner is complicated by the lack of long-range structural order in their crystal lattices. Such disorder is otherwise known as 'anion ordering'; despite exhibiting local ordering of anions within individual $MO_{6-x}N_x$ octahedra, long-range structural order is lost due to the similarity between the $M^{n+}-O^{2-}$ and $M^{n+}-N^{3-}$ orbital overlaps, and hence bond lengths. The slightly larger $M^{n+}-N^{3-}$ bond length drives a predominantly cis-ordering in individual octahedra, giving rise to N^{3-} - M^{n+} - N^{3-} chains (or O^{2-} – M^{n+} – O^{2-} in the case of LaTaON₂ and LaNbON₂).¹⁰

 a School of Environmental & Life Sciences, The University of Newcastle,

Callaghan 2308, NSW, Australia. E-mail: alister.page@newcastle.edu.au

 b School of Materials Science & Engineering, PCFM Lab, Sun Yat-sen University, Guangzhou, 510275, P. R. China

[†] Electronic supplementary information (ESI) available: Relative energies of $LaNbO₂N$, BaNbON₂ anion orderings in the presence of vacancy defects, comparison of vacancy diffusion barrier heights as a function of supercell size. See DOI: 10.1039/d1ma00122a

Fig. 1 Anion ordering in M - O_4N_2 and M - O_2N_4 octahedra.

Anion ordering induces local distortions 11 in the crystal structure, influencing the ferroelectricity,^{12,13} bandgap^{14,15} and effective charge carrier mobility of these materials.¹⁶⁻¹⁸

Anion ordering in oxynitride perovskites can be categorised into by the regimes of cis- or trans-ordering as illustrated in Fig. 1, as well as by the dimensionality the ordering travels in i.e. 2D/3D for cis-ordering or 1D/2D for trans-ordering. Recent research has focused on characterising the relative stabilities of these different regimes. While cis-ordering is the more stable for d^0 oxynitrides, both regimes are sensitive to stoichiometry, strain, and interfacial effects. For instance, synthetic conditions and A-site cation substitution can yield fully 2D or 3D transorderings.¹⁶⁻²⁰ For instance, increased Sr^{2+} doping in CaTaO₂N induces partial trans-ordering, with cis-ordering becoming energetically prohibitive due to lattice strain.²¹ Similarly, Vonruti et al.²² have shown that $>$ 4% compressive strain stabilizes trans-ordering in bulk LaTiO₂N. Ninova et al.²³ have predicted that, for LaTiO₂N, surface layers can favour transordering due to the charge neutral stacking afforded by (LaN)- $(Tio₂)$ layers.

However, the impact of anion ordering on anion vacancy defect formation and anion vacancy diffusion remains largely unexplored in perovskite oxynitride photocatalysts.

Understanding this relationship is the first step towards controlling their defect engineering for enhanced photocatalysis. Here we address this issue by demonstrating how the stability and mobility of O^{2-} and N^{3-} vacancy point defects are influenced by local anion ordering in two representative niobium perovskite oxynitrides, $BaNbO₂N$ and $LaNbON₂$. Vacancy defect formation energies and migration barriers are investigated systematically as a function of both *cis*- and *trans*-ordering of N^{3-} - M^{n+} - N^{3-} (BaNbO₂N) and O²⁻-Mⁿ⁺-O²⁻ (LaNbON₂) chains using accurate first-principles calculations. Cis-anion orderings are in general more energetically favourable than trans-orderings in these materials. However, there are secondary ordering effects that arise from the stoichiometric distribution of anions in the A-site cation coordination sphere, most prevalent for LaNbON₂. We show below how both the stability and mobility of O^{2-} and N^{3-} vacancy defects are influenced by these two ordering effects.

Computational methodology

All calculations reported here employ the Perdew–Burke– Ernzerhof generalized gradient approximation (GGA) exchange– correlation functional, revised for solids $(PBESol)²⁴$ as implemented in the Vienna Ab initio Simulation Package (VASP) code^{25,26} using the projector augmented wave (PAW) method.²⁷ A Hubbard-U $correction²⁸$ has not been employed in this work, due to the non-trivial choice of selecting $a+U$ potential that is independent of the anion ordering and vacancy defect type. All calculations employ (core)/valence configurations of Nb: ([Ne]3s)/3p3d4s; O: ([He])/2s2p N: ([He])/2s2p and a plane wave energy cutoff of 450 eV. Crystal structures of BaNbO₂N and LaNbON₂ at standard conditions were obtained from the International Crystal Structure Database (ICSD crystal IDs 434770 and 247261, respectively²⁹), see Fig. 2. These structures are, respectively, of $P_{m\bar{3}m}$ and P_{nma} symmetry. However, these symmetries are only nominal; we show below that local anion orderings within each extended supercell induce differing degrees of three-dimensional anisotropy in these ideal lattice structures when they are fully relaxed. Monkhorst–Pack meshes consisting of $6 \times 6 \times 6$ *k*-points were used for the 40 atom $2 \times 2 \times 2$ BaNbO₂N supercell, while $4 \times 6 \times 4$ k-point meshes were used for 80 atom $2 \times 1 \times 2$ $LaNbON₂$ supercell. Vacancy migration barriers were preoptimised using the nudged elastic band (NEB) algorithm 30 to a force criterion of 0.1 eV \AA^{-1} , before being re-optimised the Climbing Image (CI-NEB) algorithm³¹ to a 0.05 eV $\rm \AA^{-1}$ criterion. Five images were used in both cases.³¹ Paper Martial Advances Article 2022

Access Article 2021. The Context Article is a considerable article is a considerable article. The common and the state of the state

The formation of a neutral oxygen vacancy in $BaNbO₂N$ and $LaNbON₂$ can be expressed in Kröger–Vink notation as,

$$
O^{\times} + 2M^{\times} \rightarrow 2M' + V_O^{\bullet \bullet} + \frac{1}{2}O_2 \tag{1}
$$

and for a neutral nitrogen vacancy,

$$
N^{\times} + 3M^{\times} \rightarrow 3M' + V_N^{\bullet \bullet \bullet} + \frac{1}{2}N_2 \tag{2}
$$

The oxidation state of the M cations in the vicinity of the defect will be determined by the localization of the electron density, which may vary between $2M' + V_O^{\bullet\bullet}$ and $2M' + V_O^{\bullet\times}$ for

Fig. 2 Unit cells of (a) $P_{m\bar{3}m}$ BaNbO₂N and (b) P_{nma} LaNbON₂

a neutral oxygen vacancy, and $3M' + V_N^{\bullet\bullet}$ and $3M + V_N^{\bullet\times}$ for a neutral nitrogen vacancy. As Nb is capable of exhibiting a range of oxidation states (+5 to -1), we reason that it will likely favour charged vacancies and reduced Nb cations in anion-deficient $BaNbO₂N$ and $LaNbON₂$. To this end we investigate the effect of anion ordering effect on electron localization and Nb oxidation via a DDEC6 charge decomposition analysis. $32-34$

For a neutral supercell, the anion vacancy formation energy, $E_{\text{f,vac}}$, can be obtained from the simplified expression:³⁵

$$
E_{\text{f,vac}} = E_{\text{def}} - E_{\text{pris}} + \frac{1}{2}\mu_i \tag{3}
$$

where E_{def} is the defective supercell, E_{pris} is pristine supercell and μ_i chemical potential for the missing anion. In this work we seek only to establish how anion ordering influences anion vacancy formation (and vice versa), and do not consider the additional effects induced by oxygen/nitrogen-rich synthesis conditions. To this end we only employ a single nitrogen and oxygen chemical potential,obtained from the fitted elemental-phase reference energies (FERE) ($\mu_{\rm O}$ = -4.76 eV and $\mu_{\rm N}$ = -8.51 eV).³⁶

Results and discussion

Anion ordering in Nb perovskite oxynitrides

BaNbO₂N. Fig. 3 illustrates each anion ordering considered here for BaNbO₂N. This anion ordering is defined in terms of N^{3-} –Nb⁵⁺–N^{3–} chains. The high symmetry of this structure

 $(P_{m3m}$ space group) enables an extensive search of anion orderings, including those that possess completely cis- or trans-ordered N^{3-} –Nb⁵⁺–N^{3–} chains, as well as those that possess a mixture of both cis- and trans-orderings. Fig. 4a depicts the relative energies of these distinct anion orderings, compared to the lowest energy structure of BaNbO₂N. The latter is the fully *cis-*C1 ordering, which has previously been found to be the most stable ordering in LaTiO₂N,¹⁴ CaTaO₂N¹⁷ and SrTaO₂N,¹⁵ Fig. 4a shows that several fully cis-orderings considered here lie within 0.3 eV of C1, with the parallel ring structure C8 being the only exception. By comparison, local structures with fully trans-orderings (T1 and T2) are >2 eV less stable than the C1 ordering. Of these, the lowest energy fully trans configuration is T2, a 2D ordering comprised of stacked orthogonal layers of $trans-N^{3-}-Nb^{5+}-N^{3-}$ chains. Interestingly, Fig. 4a shows that the partial *trans-anion* orderings considered here (Q and H, which denote quarter and half trans-orderings, respectively) are all significantly lower in energy than the fully trans-configurations. Indeed, several quarter trans-orderings, such as Q1 and Q4, are predicted to be within 0.2 eV of the lowest energy C1 ordering. This suggests that distortion caused by trans-chains can be stabilized to a point by adjacent cis-chains. These results suggest that stabilizing the formation of a fully *trans*-ordered phase in BaNbO₂N will be significantly more challenging than conversion from *cis*- to partially/fully trans-ordered phases, or forming mixed-phases comprising both cis- and trans-ordered regions. Of the ring & loop configurations considered here (i.e. C7, C8, C9, Q2, H2), Materials Advances

a neutral congest Week on 2021. As beyonds Contribution and 13.9 – 11.09.2024 15:21:27. The material consistents Article is likely and the common and the stress Complete Common and the stress Article i

Fig. 3 $\,$ N^{3–}Nb⁵⁺–N^{3–} anion orderings in a 2 \times 2 \times 2 BaNbO₂N supercell (Ba²⁺ and O^{2–} atoms not shown for clarity). N^{3–} and Nb⁵⁺ are dark and light blue atoms, respectively.

Fig. 4 (a) PBEsol energies of BaNbO₂N anion orderings relative to the lowest energy configuration for the $2 \times 2 \times 2$ supercell. Note the different scales for the cis- and trans-anion orderings. Deviation of lattice vectors from cubic symmetry for (b) cis-orderings and (c) Quarter, half and full trans-orderings in BaNbO₂N

Fig. 4a indicates that these orderings generally have higher energies than their counterparts. This suggests potentially unfavourable strain is produced when the $\mathrm{N}^{3-}\mathrm{-Nb}^{5+}\mathrm{-N}^{3-}$ chains 'loop back' upon themselves too tightly.

Fig. 4 also considers the relative stability of anion orderings in BaNbO₂N from the point of view of dimensionality, by highlighting which orderings are 2D and which are 3D. In general, cis-orderings are considered to be 2D, in that they consist predominantly of $N^{3-}-Nb^{5+}-N^{3-}$ chains aligned parallel with each other. Introducing locally *trans*-ordered $N^{3-}-Nb^{5+} N^{3-}$ chains, either partially (Q configurations) or completely (T configurations) naturally increases the dimensionality of the ordering when 'twists' and 'loops' are present in the $N^{3-}-Nb^{5+} N^{3-}$ sublattice (see Fig. 3). Nevertheless, Fig. 4a shows that there is no clear relationship between dimensionality and stability here. For instance, the 2D fully cis-ordered structures are lowest in energy (e.g. C1, C2, C3) whereas the 2D halftrans-ordered structures (H1, H2, H3, H4) are less stable by

up to \sim 1.2 eV. Similarly, of the 3D orderings studied, the fully cis-C7, C10 and C11 and the quarter trans-Q1 and Q4 orderings have mostly very similar energies. Notably, the C10, Q1 and Q4 orderings are only ~ 0.13 eV higher in energy than the most stable C1 configuration. On the other hand, the dimensionality of the ordering has a clear relationship with the degree of lattice anisotropy, measured here via the deviation of each lattice constant from its value for a pristine $P_{m\bar{3}m}$ unit cell for each ordering (*i.e.* we define this value as the average of the a, b and c lattice constants in each case). Fig. 4b and c show that the 2D orderings considered here clearly induce greater anisotropy in the crystal lattice compared to 3D orderings, as anticipated. Indeed, lattice constant deviations correlate with the predominant alignment of $N^{3-}-Nb^{5+}-N^{3-}$ chains in the 2 \times 2 \times 2 supercell. For instance, for the most stable C1 ordering, the N^{3-} -Nb⁵⁺-N³⁻ chains are aligned along the a and c lattice vectors (Fig. 3). This alignment causes equivalent expansion in a and c of ~ 0.03 Å, while the lattice is compressed in b approximately twofold, by \sim -0.07 A (Fig. 4b). On the other hand, C7 yields an essentially isotropic $P_{m\bar{3}m}$ structure, and Fig. 3 shows that alignment of the N^{3-} -Nb⁵⁺-N³⁻ chains are balanced exactly in all three directions. These results suggest that inducing lattice strain in $BaNbO₂N$ will yield selectivity over anion ordering which is in line with recent studies for other oxynitride perovskites.^{16,21,22} We return to a discussion of how these orderings influence thermokinetic parameters of vacancy defects below.

Beyond dimensionality, Fig. 4b and c show that these $BaNbO₂N$ anion orderings fall into three distinct groups. The first of these comprise orderings that minimally distort the cubic symmetry of the ideal BaNbO₂N lattice, *i.e.* $a \approx b \approx c$. This isotropic group includes both high energy fully *trans*orderings, as well as relatively stable C7 and Q2 orderings, and feature motifs in which $N^{3-}-Nb^{5+}-N^{3-}$ chains form a closed loop within a 2 \times 2 \times 2 extension of the BaNbO₂N lattice. The second group consists of orderings that yield anisotropic compression/expansion in a single dimension (*i.e.* two lattice constants essentially equal but distinct from the third), such as C1, C2, C3, H1, H3, and H4. The third group are those orderings that yield complete anisotropy in the sense that all three BaNbO₂N lattice constants are unique (e.g. C3, C4 Q3, Q4, Q5). In the second group, the C1, C2 & C3 orderings (the three most stable orderings overall) are compressed in one axis and expanded in the other two. Interestingly, the half transorderings in this group (H1, H3, H4) all show the opposite trend, i.e. compression along two axes, and expansion in the third. These results suggest that *cis*-orderings in perovskite oxynitrides may be stabilised generally via in-plane lattice expansion, while trans-ordering may be stabilised via in-plane lattice compression. Indeed, Oka et al^{21} demonstrated the formation of a metastable *trans*-type anion orderings in CaTaO₂N via Sr^{2+} A-site doping, in which N^{3-} preferentially resided in axial sites in TaO₄N₂ octahedra due to lattice strain induced by the A-site dopant. Similarly, Vonruti et al ²² predicted the stabilization of parallel *trans*-ordering of N^{3-} anions (akin to T2) in LaTiO₂N when in-plane compressive strains greater than 4% were applied to the lattice. While an exhaustive strain engineering study is beyond the scope of this work, results presented here indicate that intermediate partial trans-orderings potentially impede the stabilization of fully trans-orderings via compressive in-plane strain, and so are likely an important consideration in strain engineering of perovskite oxynitrides more generally.

 LaNbON_2 . Fig. 5a illustrates the anion orderings considered here for LaNbON₂. Here $NbO₂N₄$ octahedra mean that anion ordering is best examined via O^{2-} –Nb $^{5+}$ – O^{2-} chains, as opposed to $N^{3-}-Nb^{5+}-N^{3-}$ chains. The lower orthorhombic symmetry here prevents an exhaustive study of anion orderings such as that presented above for the cubic BaNbO₂N. Here we consider a set of chain structures that can provide an adequate representation of fully *cis*- or fully *trans*-anion orders in LaNbON₂, and are commensurate with representation anion order sets considered in other DFT studies.^{15,17,22} Due to the lower symmetry of LaNbON₂, we have also considered the chains in perpendicular orientations. The total orderings considered was ultimately limited to those configurations that could be constructed in the unit cell and subsequently extended into the $2 \times 1 \times 2$ supercell. Partial *trans*-orderings (akin to the Q and H structures considered for $BaNbO₂N$) are not possible in the orthorhombic LaNbON₂ unit cell without violating the $NbO₂N₄$ octahedra selection rule, and so have not been considered.

Fig. 5b displays the relative energies of the different anion orderings for the $2 \times 1 \times 2$ LaNbON₂ cell. It is clear that *cis*orderings are preferred over trans-orderings, consistent with results for BaNbO₂N presented in Fig. 4a. For LaNbON₂ all *trans*-orderings considered here (LT1, LT2 & LT3) are $>$ 2 eV less stable than the most stable cis-ordering, LC9. The latter ordering features 'zigzag' O $^{2-}$ –Nb $^{5+}$ –O $^{2-}$ chains running through the $2 \times 1 \times 2$ supercell alternating between the (101) and (101) planes and perpendicular to the (011) and (011) directions. This zigzag motif is shared with the other stable anion orderings, LC1, LC2, and LC10, the only difference being the alignment of the motif within the crystal structure. For instance, in the LC10 ordering, the same zigzag O^{2-} –Nb⁵⁺– O^{2-} chains are present in the $2 \times 1 \times 2$ supercell as those observed in the LC9 ordering. However, they are perpendicular to the (110) and (110) lattice directions, and as a result are ~ 0.2 eV higher in energy. Similarly, the LC1 and LC2 orderings also feature the zigzag O^{2–}–Nb⁵⁺–O^{2–} chain motif, but perpendicular to the (101) and (101) lattice directions, respectively. Both orderings are only \sim 0.15 eV less stable than the most favourable ordering, LC9. The other cis-orderings considered here consist of different structural motifs and are less stable than the zigzag motif. These include the LC3, LC4, LC5 and LC6 orderings, all of which consist of $O^{2-}Nb^{5+}-O^{2-}$ chains that create a repeating square-tooth pattern in the $2 \times 1 \times 2$ LaNbON₂ supercell and are \sim 0.3–0.6 eV less stable than LC9, and the ring-like LC7 and LC8 motifs, which are both \sim 0.5 eV less stable than LC9.

Fig. 5b shows that the dimensionality of the ordering is also apparently unrelated to its stability. For instance, LC9 and LC10 are both 3D orderings, whereas all their *cis*-orderings of O^{2-} - $Nb⁵⁺-O²⁻$ chains in LaNbON₂ are 2D. This is different from what was found in Fig. 4 for BaNbO₂N, and suggests that controlling for 2D or 3D more ordering may not be possible

Fig. 5 (a) Representative $O^{2-}Nb^{5+}-O^{2-}$ anion orderings in a 2 x 1 x 2 LaNbON₂ supercell (La³⁺ and N³⁻ not shown for clarity). O^{2-} and Nb⁵⁺ are red and light blue atoms, respectively. (b) PBEsol energies of LaNbON₂ anion orderings shown in (a), relative to the lowest energy ordering LC9.

in LaNbON₂. Though we note that 2D-3D ordering transition temperatures have been observed in the comparable structure LaTaON $_2$.²⁰

Further, comparison of Fig. 5b with equivalent results for $BaNbO₂N$ (Fig. 3 and 4a), shows striking similarities between the relative stabilities of comparable O^{2-} –Nb $^{5+}$ – O^{2-} orderings in LaNbON₂ and N^3 --Nb⁵⁺-N³⁻ orderings in BaNbO₂N. For instance, cis-orderings featuring zigzag chain motifs are the most stable in both materials, and fully trans-anion orderings are significantly higher in energy (by \sim 1 eV). Further, the ringlike and loop-like motifs are also energetically similar in both cases. For instance, LC3, LC4, LC5 and LC6 in $LaNbON₂$ and C4, C5 and C6 in BaNbO₂N both feature repeating square tooth O^{2–}–Nb⁵⁺–O^{2–} and N^{3–}–Nb⁵⁺–N^{3–} motifs and have comparable relative energies (\sim 0.3–0.4 eV). Similarly, the loop motifs in LC7 and LC8 for LaNbON₂ and C8 and C9 in BaNbO₂N are structurally similar and \sim 0.5–0.6 eV higher in energy than the most favourable ordering.

Nevertheless, the La^{3+} and Ba^{2+} A-site cations also drive secondary ordering effects that differ between $BaNbO₂N$ and LaNbON₂. Comparing the La- $(O,N)_{12}$ coordination spheres between different orderings for LaNbON_2 , the lowest energy orderings (LC1–LC4 and LC9–10) all possess stoichiometric $La-O₄N₈$ coordination spheres, whereas the LC5–LC8 orderings feature a mix of La-O₆N₆ and La-O₂N₁₀ coordination spheres. This mix creates alternating LaO and LaN layers between NbON^{3–} sheets aligned perpendicular to the *b*-axis in the 2 \times 1×2 LaNbON₂ supercell. There is a clear energetic difference between these two groups, with LC5–LC8 being ~ 0.3 eV higher on average than LC1–LC4 and LC9–10. This A-site effect is notably absent in the BaNbO₂N orderings, shown in Fig. 4. While the most stable zigzag orderings in $BaNbO₂N$ do indeed create an even distribution of $Ba-O_8N_4$ coordination spheres, other orderings like C6 & C8 have energies 0.15–0.30 eV higher than orderings with comparable Ba- $(O,N)_{12}$ coordination sphere distributions. Thus, the $N^{3-}-Nb^{5+}-N^{3-}$ chain ordering in the more densely packed $BaNbO₂N$ supercell effectively masks the effect of the Ba- $(O,N)_{12}$ coordination sphere on the stability of the material.

Vacancy defect formation energies in Nb perovskite oxynitrides

We now consider the impact of anion order in $BaNbO₂N$ and LaNbON₂ on the stability of vacancy defects in both O^{2-} and N^{3-} lattice positions. This is achieved via predicting the vacancy defect formation energies, $E_{f,vo}$ and $E_{f,VN}$ (eqn (3)) at the 12 distinct anion positions in the respective supercells of these materials, for each specific anion ordering considered in the preceding discussion. These distinct positions 1–12 for the $2 \times 2 \times 2$ BaNbO₂N and $2 \times 1 \times 2$ LaNbON₂ supercells are depicted in Fig. 6a and b.

BaNbO₂N. $E_{f,vo}$ and $E_{f,VN}$ in BaNbO₂N for the 22 individual anion orderings introduced previously are shown in Fig. 6c, and grouped according to their cis-/trans-character. This figure shows that nitrogen vacancy defects in $BaNbO₂N$ have a smaller variation in formation energies in the presence of fully cis- (2.24–2.50 eV) or fully trans-ordering (1.14–1.46 eV). By contrast,

Fig. 6 (a and b) Depiction of the 12 unique anion positions in the centroid of the (a) $2 \times 2 \times 2$ BaNbO₂N and (b) $2 \times 1 \times 2$ LaNbON₂ supercells. Light and dark blue spheres as in Fig. 3, red, green and yellow spheres represent oxygen, barium and lanthanum atoms respectively. (c and d) PBEsol vacancy formation energies V_N and V_O in positions 1–12 as a function of anion ordering for the (c) $2 \times 2 \times 2$ BaNbO₂N and (d) $2 \times 1 \times 2$ LaNbON₂ supercells; red indicates an O^{2-} vacancy defect and blue indicates an N^{3-} vacancy defect at that position. The colour depth indicates the magnitude of the formation energies V_N and V_O (blue and red, respectively) in eV, as indicated by the scale bar.

the range of $E_{f,VN}$ formation energies found for the quarter and half trans anion orderings (i.e. Q1-Q5 and H1-H4) is larger, 1.35–2.77 eV, presumably due to the greater structural anisotropy in these orderings, which gives rise to a greater number of distinct vacancy sites. Fig. 6c also shows that V_N sites in local cis- or local trans-anion sites of the partial trans-orderings are \sim 0.2 eV higher in energy than if the whole ordering was cis or trans.

The $E_{f,vo}$ formation energies vary quite significantly across the fully cis- (4.87–6.08 eV) and partial trans- (4.87–5.98 eV) orderings, compared to fully trans- (5.37–5.60 eV). This indicating that

neutral oxygen vacancy formation energies are more sensitive to N^{3-} –Nb $^{5+}$ –N $^{3-}$ chain disorder, whereas nitrogen vacancy formation energies depend simply on whether the site is in a cis- or *trans*-chain. N³⁻ anions in *trans*-N³⁻-Nb⁵⁺-N³⁻ chains, whether they be partially or fully *trans*-, have a reduced $E_{f,VN}$ formation energy.

The question as to whether anion vacancies have the potential to affect local anion order in BaNbO₂N is considered in Fig. $S1$ (ESI†), which shows the relative energies of these anion orderings in the presence of V_{Ω} and V_{N} defects. Compared to the defect-free relative energies in Fig. 4a, it can be seen in Fig. S1a (ESI†) that in several cases anion vacancies likely influence the anion ordering. For instance, the quarter-trans orderings (Q1–5) become energetically indistinguishable from the cis (C1-11) orderings in the presence of a neutral oxygen vacancy defect. Additionally, the relative orderings in the presence of a neutral nitrogen vacancy defect (Fig. S1b, ESI†) are significantly different from the defect-free orderings. While the cis C1-11 orderings are effectively equivalent in energy with and without a V_N defect, the Q1-5 orderings are lowered by \sim 0.5 eV, as the defect disrupts an $N^{3-}-Nb^{5+}-N^{3-}$ chain, thus making them the most favourable ordering.

LaNbON₂. $E_{f,vo}$ and $E_{f,VN}$ in LaNbON₂ for the 12 individual anion orderings introduced previously are shown in Fig. 6d, and grouped according to their cis-/trans-character. It is clear from this figure that N^{3-} anion vacancies form more easily in fully trans-ordered structures (LT1, LT2, LT3) than the cis-ordered LC structures. The LC5, LC6, LC7, and LC8 orderings are exceptions here, exhibiting $E_{f,vo}$ values between only 4.82-5.54 eV (the lowest values for all orderings considered here). We note that these are the orderings that yield alternating LaO/LaN layers in LaNbON₂. $E_{f,NN}$ values are notably higher than $E_{f,NO}$ for the LC orderings. This means that LaNbON_2 domains rich in these more stable cis-anion orderings are more likely to exhibit O^{2-} anion vacancies than N^{3-} vacancies. The same trend was noted above in our discussion of $BaNbO₂N$ vacancy defects, indicating that the relative stabilities of O^{2-} and N^{3-} point vacancy defects in AMO₂N & AMON₂ perovskite oxynitrides are potentially independent of the A-site cation and crystal symmetry. $E_{f,VN}$ formation energies predicted for LaNbON₂ (1.74 eV range) here are more sensitive to the local anion ordering, and thus exhibit a wider range than observed for $BaNbO₂N$

(1.36 eV range). This is attributed to N^{3-} being the majority anion LaNbON₂.

Finally, we consider whether or not V_O and V_N defects in $LaNboN₂$ have the potential to influence the relative stabilities of these anion orderings. Fig. S2 (ESI†) shows the relative energies of the anion orderings in the defective LaNbON_2 structures. Compared to the defect-free relative energies in Fig. 5a and Fig. S2a (ESI†) the presence of V_{Ω} does not appreciably influence the relative stability of many of the orderings. However, the LC5–9 orderings become the most stable orderings, as they can be lowered by up to ~ 0.2 eV. Fig. S2b (ESI†) shows that, by comparison, V_N defects are much more influential on the stabilities of these anion orderings, with all LC orderings being lowered by up to \sim 0.5 eV. Nevertheless, the relative stabilities for all LC orderings become virtually indistinguishable in the presence of a V_N defect.

Vacancy defects and charge redistribution

Nb is capable of exhibiting multiple low oxidation states. While our calculations enforce charge neutrality on the entire supercell, the region surrounding a vacancy defect is itself not necessarily guaranteed to be charge neutral due to electron density delocalisation. In turn, the oxidation states of Nb cations are not necessarily uniform throughout the supercell, since, despite the formal vacancy charge (*i.e.* $V_O^{\bullet\bullet}$ and $V_N^{\bullet\bullet\bullet}$) electron density delocalisation around the vacancy may lead to charge redistribution on neighbouring atoms. We examine this possibility in BaNbO₂N and LaNbON₂ via the DDEC6 charge decomposition analysis presented in Table 1, which summarizes the partial atomic charges and charge redistribution in these materials following the formation of V_{O} and V_{N} defects. We limit our discussion to only the most thermodynamically stable cis/trans anion orderings for each material, established in the preceding discussion (i.e. C1 and T2 in BaNbO₂N, and LC9 and LT2 for L aNbON₂). Due to the size of the supercells used in this work, we have not verified these PBEsol results with higherlevel exchange correlation functionals, such as HSE06.³⁷ We point out, however, that charges based on PBEsol have been shown to be in close agreement with higher level functionals for closely related transition metal oxide perovskites.^{38,39} Materials Advances

National Society. Notices Articles Arti

> For BaNbO2N, Table 1 shows that charge redistribution, due to delocalisation of the electron density around the defect site, leads to the 'true' charge of $V_O^{\bullet\bullet}$ and $V_N^{\bullet\bullet\bullet}$ defects being only

Table 1 DDEC6 average partial charges and charge redistribution (e) in BaNbO₂N and LaNbON₂ due to 'neutral' V_O and V_N vacancy defects using PBEsol. The redistributed charge of an atom is defined as the difference between its DDEC6 charges in the pristine and defective supercell; negative values indicate reduction in the presence of the defect. All values are averaged over all O^{2-}/N^{3-} positions in each respective supercell. 'Adjacent' Nb cations are those that are coordinated to the anion removed to form the defect, 'other' Nb cations are all that are not

		BaNbO ₂ N				LaNbON ₂			
		C ₁		T2		LC9		LT ₂	
		V_{Ω}	$\rm V_N$	V_{Ω}	$\rm V_N$	V_{Ω}	V_{N}	V_{O}	V_{N}
Partial Charge	Adjacent Nb Other Nb	1.89 2.21	1.73 2.20	1.93 2.20	1.84 2.17	1.78 2.08	1.64 2.08	1.64 2.09	1.89 2.08
Charge Redistribution	Ba/La Nb Total	-0.25 -1.26 -1.51	-0.30 -1.65 -1.95	-0.28 -1.15 -1.43	-0.32 -1.47 -1.79	-0.27 -0.79 -1.06	-0.35 -1.11 -1.46	-0.46 -1.71 -2.17	-0.34 -1.36 -1.70

65–75% of the expected formal values. For instance, for the C1 ordering the total charge redistribution in the $BaNbO₂N$ lattice around a V_O defect is -1.51 e (rather than exactly -2), while for a V_N defect this figure is -1.95 e (rather than exactly -3 e). Charge redistribution is slightly smaller for the T2 ordering (1.43 and 1.79 e, respectively), indicating that anion vacancies near trans- N^{3-} -Nb⁵⁺-N³⁻ anion configurations in BaNbO₂N lead to less charge redistribution away from the vacancy site in the cation sublattice. Table 1 also shows that, following the formation of a neutral oxygen or nitrogen vacancy defect in $BaNbO₂N$, the oxidation state of a Nb cation depends on its vicinity to the defect. For instance, the average partial charge of adjacent Nb cations is \sim 0.5 e lower than those not coordinated to the vacancy. This is true for both the C1 and T2 anion orderings, although the effect is again marginally smaller for the latter (\sim 0.3 e), likely for the same reason noted above.

Very similar trends are observed in Table 1 regarding partial charges of adjacent and uncoordinated Nb cations in LaNbON₂. For instance, for the lowest energy LC9 ordering, Nb cations adjacent to a V_O defect are more reduced by \sim 0.3 e, compared to Nb cations further away. Similarly, for a V_N defect, the difference is \sim 0.4 e (adjacent Nb cations being more reduced). Nb partial charges in the LT2 ordering are analogous by comparison. As for BaNbO₂N, Table 1 shows that the DDEC6 charges obtained using PBEsol are notably lower than the expected formal values. For instance, in the LC9 ordering, the LaNbON₂ lattice is reduced by only 1.06 e (V_O) and 1.46 e (V_N), respectively. However, a marked increase in charge distribution is observed for an oxygen vacancy defect in the LT2 ordering. Here, the LaNbON₂ lattice is reduced by 2.17 e, which is comparable to the expected formal charge $(i.e. -2)$. This is attributed to the effect of the ordering itself. While the pristine

LT2 and LC9 structures exhibit essentially equivalent average Nb partial charges (2.14 compared to 2.10, not shown in Table 1), the introduction of the V_O defect disrupts a *trans*-O²⁻-Nb⁵⁺-O²⁻ chain in the LT2 structure, but leaves the chain in the cis LC9 ordering unperturbed. This reduces the Nb sublattice by \sim 1 e (-1.71 e) more in LT2, compared to LC9 ordering (-1.06 e) .

Vacancy diffusion barriers in Nb perovskite oxynitrides

With the thermochemical stability of $V_N^{\bullet\bullet}$ and $V_O^{\bullet\bullet}$ vacancy defects in BaNbO₂N and LaNbON₂ now established, we conclude our discussion by considering the impact of anion ordering on vacancy diffusion kinetics in these materials. Fig. 7 presents diffusion pathways of $V_N^{\bullet\bullet\bullet}$ and $V_O^{\bullet\bullet\bullet}$ vacancy defects in BaNbO₂N and LaNbON₂ elucidated using the CI-NEB method. Due to the expense of this approach, we consider defect diffusion in only the most stable *cis*- and *trans*-anion orderings (*i.e.* the most relevant), and only those diffusion pathways corresponding to vacancies moving between adjacent positions in the crystal lattice $(i.e.$ the distinct positions defined in Fig. 6a and b). For BaNbO₂N we assess the accuracy of diffusion barriers obtained with the 2 \times 2×2 supercell by comparing them with barriers obtained with a $3 \times 3 \times 3$ supercell in Fig. S3 and Table S3 (ESI†). This comparison indicates that shorter diffusion pathways (rotation about a Nb ion) are significantly less sensitive to supercell size than longer pathways (shifting to other Nb ion coordination spheres), as anticipated. Importantly, Table S3 (ESI†) indicates that the trends in vacancy diffusion barriers found using the 2 \times 2×2 supercell are reproduced with the $3 \times 3 \times 3$ supercell. The longer diffusion pathways in LaNbON₂ are orientated to the a/c plane in the $2 \times 1 \times 2$ orthorhombic supercell, and so it is likely they are already converged with respect to the a/c dimensions of the supercell. Paper

80-75% of the capcted formal values. Por instance, for the C1 172 and LO structures entries the structure
of certain the capital common and the structure of the structure of the structure of the structure of the str

Fig. 7 V_N & V_O diffusion pathways and associated PBEsol diffusion barriers (eV) for the most stable cis- and trans-anion orderings in (a and b) BaNbO₂N and (c and d) LaNbON₂. (a) BaNbO₂N, C1 ordering, (b) BaNbO₂N, T2 ordering, (c) LaNbON₂, LC9 ordering, (d) LaNbON₂, LT1 ordering. All diffusion pathways were calculated in the complete $3 \times 3 \times 3$ and $2 \times 1 \times 2$ supercells, respectively, however only the core region is depicted here for clarity. Anion positions are defined in Fig. 5a and b for BaNbO₂N and LaNbON₂, respectively.

BaNbO₂N. Diffusion pathways for $V_N^{\bullet\bullet}$ and $V_O^{\bullet\bullet}$ vacancies in the most stable C1 and T2 orderings of $BaNbO₂N$ and the corresponding energy barriers are presented in Fig. 7a and b. There are two distinct pathways for each vacancy type; the defect can rotate positions about a Nb^{5+} cation (O1 \rightarrow O2, $N7 \rightarrow N12$ in C1 (Fig. 7a); O5 \rightarrow O2 in T2 (Fig. 7b)) or the defect can migrate to a non-adjacent position, shifting to a different Ta⁵⁺ cation but rotating about the Ba²⁺ cation (O1 \rightarrow O4, $N7 \rightarrow N5$ in C1 (Fig. 7a); O5 \rightarrow O7, N1 \rightarrow N4 in T2 (Fig. 7b)). The latter of these pathways is obviously the longest and so is anticipated to correlate with the highest diffusion barrier. Fig. 7a and b show that this is the case; $V_O^{\bullet\bullet}$ and $V_N^{\bullet\bullet\bullet}$ migration between non-adjacent sites is impeded by barriers ranging from 5.33 eV to 6.70 eV while rotation of the defect to adjacent positions around the $Nb⁵⁺$ cation corresponds to a barrier of only 1.13 eV to 1.53 eV. Fig. 7a and b also shows that, in general, $V_O^{\bullet\bullet}$ diffusion are barriers around \sim 25% lower than the analogous diffusion pathways for $V_N^{\bullet\bullet\bullet}$. $V_N^{\bullet\bullet\bullet}$ diffusion in the less stable T2 ordering is further impeded by the fact that it can only occur between adjacent NbO_4N_2 octahedra, and not *via* rotation around an individual $Nb⁵⁺$ cation, which would correspond to a lower barrier. Thus, despite being more stable in BaNbO₂N, O $^{2-}$ vacancy defects are expected to undergo more facile diffusion, most notably in trans-ordered phases, impeding control over long range ordering in this material. We note also that these findings are consistent with our recent study of TaON, in which it was shown that diffusion barriers for $V_O^{\bullet\bullet}$ migration are potentially low enough to displace nitrogen content from the oxynitride.⁴⁰ Thus, it is suggested that stabilising trans-ordered phases of BaNbO₂N is likely to yield more effective retention of nitrogen content in this material, even if long-range order cannot be controlled. **Materials Advances**
 Excels the most stable C1 and 12 orderings of the 2021. The articles of a sample of the access are the common stable C1 and 12 ordering are none stable C1 and 12. This are not stable and the same i

LaNbON₂. Diffusion pathways for $V_N^{\bullet\bullet}$ and $V_O^{\bullet\bullet}$ vacancies in the lowest energy cis - and trans-ordered structures of LaNbON₂, LC9, and LT1, and the corresponding energy barriers are presented in Fig. 7c and d. In the LC9 *cis*-ordering, Fig. 7c, O^{2-} vacancies can only diffuse *via* the O10 \rightarrow O12 pathway, while N^{3–} vacancies have two possibilities for migration via rotation around a Nb⁵⁺ cation (N6 \rightarrow N3) and between non-adjacent (N6 \rightarrow N7) sites that cross the cavity between La^{3+} ions. In the latter case the defect is shifting to a separate $NbO₂N₄$ octahedron while rotating is coordinating about the local La- $(O,N)_{12}$ coordinate sphere. In the less stable LT1 trans-ordering, Fig. 7d, again O^{2-} vacancies can only migrate via a single long pathway (O6 \rightarrow O7) since rotation about an individual Nb⁵⁺ cation to another O^{2-} site isn't possible, whereas N^{3-} vacancies can migrate via the N12 \rightarrow N10 pathway and along the longer non-adjacent $N10 \rightarrow N3$ pathway.

Consistent with BaNbO₂N, the V^{**} diffusion barriers are seen to be sensitive to whether the diffusion path is between adjacent anion positions or non-adjacent positions with O10 \rightarrow O12 in LC9 predicted to have a barrier height of 0.71 eV, while O6 \rightarrow O7 in LT2 is 3.12 eV. However, the difference in the V^{***} diffusion barriers for adjacent and non-adjacent paths is greatly reduced compared to BaNbO₂N. In LC9 the adjacent pathway height is 1.56 eV with the non-adjacent path barrier only 0.29 eV higher, while in LT2 the adjacent $V_N^{\bullet\bullet\bullet}$ diffusion is now 0.72, with the non-adjacent barrier merely 0.82 eV.

Interestingly, the kinetic stability of N^{3-} vacancy defects in $LaNbON₂$ is significantly reduced from that observed in BaNbO₂N. In particular, in the less stable LT1 trans-ordering, N^{3-} vacancies diffuse more easily than O^{2-} vacancies. The origin of this reduction is a cooperative effect in which the migrating $V_N^{\bullet\bullet\bullet}$ vacancy partially displaces an adjacent O^{2-} anion. The additional space created by this displacement gives rise to an asymmetric path, in which the $V_N^{\bullet\bullet\bullet}$ vacancy does not pass directly between adjacent La^{3+} cations along the diffusion pathway. In doing so, the migrating $V_N^{\bullet\bullet\bullet}$ vacancy partially displaces an adjacent O^{2-} anion and so reduces the diffusion barrier.

Conclusions

The impact of anion ordering on defect engineering in perovskite oxynitride photocatalysts remains largely unexplored, despite it being the necessary first step towards controlling defects in these materials for improved photocatalytic performance. Using accurate first-principles calculations, we have presented an extensive investigation of *cis*- and *trans*-anion orderings in two exemplar perovskite oxynitrides, $BaNbO₂N$ and $LaNbON₂$, and examined how these orderings influence (1) overall stability, (2) the stability of N^{3-} and O^{2-} point defects and (3) the mobility of these defects through the bulk material. Analysis of partial charges with DDEC6 confirmed that the anion ordering influenced the degree of e^- redistribution to adjacent and outer Nb cations from the creation of an anion vacancy.

For BaNbO₂N, our results show clearly that fully *cis*-anion orderings are more stable than fully trans-anion orderings. The influence of these orderings on the cubic lattice vectors however demonstrates differing degrees of anisotropy. This ultimately has implications for strain engineering in this oxynitride; specifically, it may be difficult to stabilize the fully *trans*-orderings in BaNbO₂N, over half-trans or fully-cis orderings, via in-plane compression of the crystal lattice. In LaNbON_2 , 2D & 3D orderings have comparable energies, meaning that mixed dimensionality in anion ordering may be more prevalent in this material. However, our results show the presence of a secondary 'A-site coordination sphere effect' influencing the stability of different orderings in LaNbON₂, where the stoichiometric distribution of O^{2-} and N^{3-} in La- $(O,N)_{12}$ gives rise to obvious differences in the relative energies of different fully cis-anion orderings. This effect is less obvious in BaNbO₂N, which we contend is due to the more densely packed lattice having a higher degree of symmetry, which masks any Ba- $(O,N)_{12}$ coordination sphere on the overall stability of the material. Vacancy defect formation energies for BaNbO₂N indicate that domains rich in cis-anion orderings (the most favourable) may preferentially stabilise O^{2-} anion vacancies less uniformly in their structure, whereas N^{3-} anion vacancies are less likely. On the other hand, O^{2-} defect formation energies in $LaNbON₂$ are more sensitive to the local anion ordering, than its overall cis- or trans-character. This is attributed to the effect of the La^{3+} A-site cation, which for particular orderings considered here gives rise to alternating layers of LaO and LaN that reduce significantly O^{2-} vacancy defect formation energy.

For BaNbO₂N, V^{**} diffusion barriers were found to be lower than $V_N^{\bullet\bullet\bullet}$, most notably in *trans*-ordered phases. It is predicted that this will impede control over long range ordering in BaNbO₂N. Stabilising trans-ordered phases of this material is thus predicted to yield more effective retention of nitrogen content in its structure, irrespective of long-range anion order. In this sense, $BaNbO₂N$ bears some resemblance to TaON, for which $V_O^{\bullet\bullet}$ migration barriers are potentially low enough to displace nitrogen content from the oxynitride.⁴⁰ The reverse is observed true for $V_N^{\bullet\bullet\bullet}$ in LaNbON₂, *i.e.* N³⁻ vacancy defects exhibit lower diffusion barriers than O^{2-} vacancy defects. V_N diffusion *via* both adjacent and non-adjacent pathways in LaNbON_2 benefits from the lower orthorhombic symmetry. In part this is due to the disruption the migrating N^{3-} defect has on the local coordination environment; when rotating around a local La $^{3+}$ cation the N^{3-} anion vacancy partially displaces Paper Materials Advances (1999) and the set of Paper Matches Article of the computer of the property of the National Creative Computer Computer Computer Computer Computer Computer Commons Article is licensed under the National Creative Commons Articles. The Commons Ar

Conflicts of interest

an adjacent O^{2-} ion to lower the diffusion barrier.

There are no conflicts to declare.

Acknowledgements

This research was undertaken with the assistance of resources provided at the NCI National Facility systems at the Australian National University, through the National Computational Merit Allocation Scheme supported by the Australian Government. AJP acknowledges support from the Australian Research Council (INTERSECT, LE170100032). JJB acknowledges support from an Australian Government RTP Scholarship.

Notes and references

- 1 T. Takata and K. Domen, Dalton Trans., 2017, 46, 10529–10544.
- 2 M. Ahmed and G. Xinxin, Inorg. Chem. Front., 2016, 3, 578–590.
- 3 B. Siritanaratkul, K. Maeda, T. Hisatomi and K. Domen, ChemSusChem, 2011, 4, 74–78.
- 4 D. Y. Wan, Y. L. Zhao, Y. Cai, T. C. Asmara, Z. Huang, J. Q. Chen, J. Hong, S. M. Yin, C. T. Nelson, M. R. Motapothula, B. X. Yan, D. Xiang, X. Chi, H. Zheng, W. Chen, R. Xu, Ariando, A. Rusydi, A. M. Minor, M. B. H. Breese, M. Sherburne, M. Asta, Q. H. Xu and T. Venkatesan, Nat. Commun., 2017, 8, 15070.
- 5 X. Xu, C. Randorn, P. Efstathiou and J. T. S. Irvine, Nat. Mater., 2012, 11, 595–598.
- 6 K. Maeda, M. Higashi, B. Siritanaratkul, R. Abe and K. Domen, J. Am. Chem. Soc., 2011, 133, 12334–12337.
- 7 J. Liu, Z. Wei and W. Shangguan, ChemCatChem, 2019, 11, 6177–6189.
- 8 S. Bai, N. Zhang, C. Gao and Y. Xiong, Nano Energy, 2018, 53, 296–336.
- 9 Y.-I. Kim, P. M. Woodward, K. Z. Baba-Kishi and C. W. Tai, Chem. Mater., 2004, 16, 1267–1276.
- 10 M. Yang, J. Oró-Solé, J. A. Rodgers, A. B. Jorge, A. Fuertes and J. P. Attfield, Nat. Chem., 2010, 3, 47.
- 11 A. Fuertes, J. Mater. Chem., 2012, 22, 3293–3299.
- 12 D. Oka, Y. Hirose, H. Kamisaka, T. Fukumura, K. Sasa, S. Ishii, H. Matsuzaki, Y. Sato, Y. Ikuhara and T. Hasegawa, Sci. Rep., 2014, 4, 4987.
- 13 Y. Hinuma, H. Moriwake, Y.-R. Zhang, T. Motohashi, S. Kikkawa and I. Tanaka, Chem. Mater., 2012, 24, 4343–4349.
- 14 N. Vonrüti and U. Aschauer, *Phys. Rev. Mater.*, 2018, 2, 105401.
- 15 A. Ziani, C. Le Paven, L. Le Gendre, F. Marlec, R. Benzerga, F. Tessier, F. Cheviré, M. N. Hedhili, A. T. Garcia-Esparza, S. Melissen, P. Sautet, T. Le Bahers and K. Takanabe, Chem. Mater., 2017, 29, 3989–3998.
- 16 H. Johnston, A. P. Black, P. Kayser, J. Oró-Solé, D. A. Keen, A. Fuertes and J. P. Attfield, Chem. Commun., 2018, 54, 5245–5247.
- 17 A. Kubo, G. Giorgi and K. Yamashita, Chem. Mater., 2017, 29, 539–545.
- 18 P. J. Camp, A. Fuertes and J. P. Attfield, J. Am. Chem. Soc., 2012, 134, 6762–6766.
- 19 S. G. Ebbinghaus, H.-P. Abicht, R. Dronskowski, T. Müller, A. Reller and A. Weidenkaff, Prog. Solid State Chem., 2009, 37, 173–205.
- 20 L. Clark, J. Oro´-Sole´, K. S. Knight, A. Fuertes and J. P. Attfield, Chem. Mater., 2013, 25, 5004–5011.
- 21 D. Oka, Y. Hirose, F. Matsui, H. Kamisaka, T. Oguchi, N. Maejima, H. Nishikawa, T. Muro, K. Hayashi and T. Hasegawa, ACS Nano, 2017, 11, 3860–3866.
- 22 N. Vonrüti and U. Aschauer, *Phys. Rev. Lett.*, 2018, 120, 046001.
- 23 S. Ninova and U. Aschauer, J. Mater. Chem. A, 2017, 5, 11040–11046.
- 24 J. P. Perdew, A. Ruzsinszky, G. I. Csonka, O. A. Vydrov, G. E. Scuseria, L. A. Constantin, X. Zhou and K. Burke, Phys. Rev. Lett., 2008, 100, 136406.
- 25 G. Kresse and J. Furthmüller, Comput. Mater. Sci., 1996, 6, 15-50.
- 26 G. Kresse and J. Furthmüller, Phys. Rev. B: Condens. Matter Mater. Phys., 1996, 54, 11169–11186.
- 27 G. Kresse and D. Joubert, Phys. Rev. B: Condens. Matter Mater. Phys., 1999, 59, 1758–1775.
- 28 B. Himmetoglu, A. Floris, S. de Gironcoli and M. Cococcioni, Int. J. Quantum Chem., 2014, 114, 14–49.
- 29 M. Hellenbrandt, Cryst. Rev., 2004, 10, 17–22.
- 30 G. Henkelman and H. Jónsson, J. Chem. Phys., 2000, 113, 9978–9985.
- 31 G. Henkelman, B. P. Uberuaga and H. Jónsson, J. Chem. Phys., 2000, 113, 9901–9904.
- 32 N. G. Limas and T. A. Manz, RSC Adv., 2018, 8, 2678–2707.
- 33 T. A. Manz and N. G. Limas, RSC Adv., 2016, 6, 47771–47801.
- 34 N. G. Limas and T. A. Manz, RSC Adv., 2016, 6, 45727–45747.
-
- 35 M. Pavone, A. M. Ritzmann and E. A. Carter, Energy Environ. Sci., 2011, 4, 4933–4937.
- 36 V. Stevanović, S. Lany, X. Zhang and A. Zunger, Phys. Rev. B: Condens. Matter Mater. Phys., 2012, 85, 115104.
- 37 J. Heyd, G. E. Scuseria and M. Ernzerhof, J. Chem. Phys., 2003, 118, 8207–8215.
- 38 R. Wahl, D. Vogtenhuber and G. Kresse, Phys. Rev. B: Condens. Matter Mater. Phys., 2008, 78, 104116.
- 39 K. K. Ghose, A. Bayon and A. J. Page, J. Phys. Chem. C, 2020, 124, 27055–27063.
- 40 J. J. Bown and A. J. Page, J. Mater. Chem. A, 2019, 7, 13029–13035.