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Sub-extensive Entropies and Open Order in Perovskite Oxynitrides**

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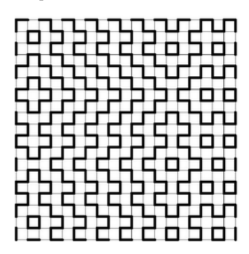
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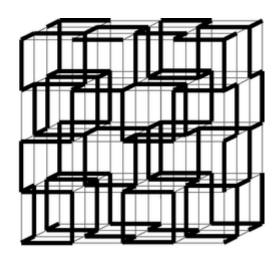
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Keywords:

Oxynitride perovskite; configurational entropy; ice rules

Abstract

Unusual subextensive configurational entropies that vary with particle size and tend to zero per atom in macroscopic samples are predicted for AMO_{3-z}N_z oxynitrides with perovskite type crystal structures. These materials are crystallographically disordered on the atomic scale, but local anion order produces chains of M–N–M bonds that undergo a 90° turn at each M cation, giving rise to subextensive entropies in materials such as SrTaO₂N, LaNbON₂, and EuWO_{1.5}N_{1.5}. A general Pauling ice-rules formula is used to calculate the extensive molar entropies for other cases such as SrMoO_{2.5}N_{0.5} and BaTaO₂N. The subextensive oxynitrides are usefully classified as showing an "open order", related to the correlated order of displacements in ferroelectric perovskites such as BaTiO₃. This raises the possibility that further open-ordered oxynitride or molecular structures may be accessible, and other states such as spins and charges may also show novel open orders.

Introduction

Configurational entropies provide useful information about atomic correlations and local order, and they may sometimes be predicted from simple statistical models as in Pauling's famous calculation of the residual entropy of ice. ^[1] Configurational entropies for crystalline materials containing atomic disorder are normally extensive, being proportional to the number of atoms. A recent study has revealed that AMO_{3-z}N_z perovskite oxynitrides contain chains of M–N–M bonds that undergo a 90° turn at each M cation, for which Pauling icerules predict zero configurational entropy despite the apparent structural disorder. ^[2] This unusual observation prompted the full entropy analysis presented here.

The ideal AMX₃ perovskite structure consists of small M cations at the vertices of a simple cubic cell bridged by anions X at the centers of all edges and a large A cation at the cell center. The lattice contains an infinite cubic network of vertex sharing MX₆ octahedra. Most perovskites are based on a single anion such as oxide or fluoride, but transition-metal oxynitride perovskites AMO_{3-z}N_z have been of recent interest for their optical and electronic properties.^[3, 4] Although full long-range anion order is not observed in these materials, a recent analysis of SrMO₂N (M = Nb, Ta) showed that well-defined *cis*-MO₄N₂ octahedra are present, resulting in disordered zigzag MN chains within two-dimensional perovskite layers.^[2] Perovskite-like layers of disordered zigzag chains are also expected in the K₂NiF₄-type oxynitrides Sr₂NbO₃N,^[5]Sr₂TaO₃N,^[6] and Ba₂TaO₃N.^[7]
The constraint that the chains must turn by 90° at each M site (with the same condition on zigzag MO chains in AMO_{3-z}N_z when z > 1.5, e.g. LaNbON₂)^[8] is not observed in other materials or in magnetic analogues of crystalline atomic materials such as spin ices. Here we explore the configurational entropies of the principal AMO_{3-z}N_z structural models that arise from this unusual structural constraint. Some structures have the unusual property of being subextensive, where the entropy per atom tends to zero in macroscopic samples,

while others have extensive entropies that are estimated by extending Pauling's ice model. We also report a new classification of structures based on their sets of long- and short-range correlation vectors, from which the unusual "open ordered" nature of the subextensive oxynitride structures is apparent.

Results

(i) Pauling Entropies of Perovskite Oxynitrides

Configurational entropies of AMO_{3-z}N_z perovskites are estimated by extending Pauling's ice model to a general lattice of N M cations, each connected to n other M cations by either M-N-M or M-O-M bridges, so there are in total nN/2 bridges. The number of M-N-M bridges emanating from each M cation is denoted by x (x = 2z for AMO_{3-z}N_z perovskites) so that the total number of M-N-M bridges is xN/2 and the fractions of M-N-M and M-O-M bridges are f = x/n and (1 - f), respectively. In the absence of local bonding constraints, the number of ways of arranging the M-N-M bridges is

$$W = \frac{(nN/2)!}{(fnN/2)![(1 - f)nN/2]!}$$

A mean-field approximation for the fraction of allowable configurations is p^{N} , where

$$p = wf^{x}(1 - f)^{n-x} = wf^{fn}(1 - f)^{(1-f)^{n}}$$

is the probability of an M ion having its bonding constraints satisfied, and w is the number of ways of orienting M-N-M bridges to the M cation's n neighbors. Using Stirling's approximation ($\ln N! \approx N \ln N - N$), the entropy is

$$S = k_B \ln(Wp^N) \approx Nk_B \ln[wf^{fn/2}(1-f)^{(1-f)n/2}]$$

As a check, identifying M-N-M and M-O-M bridges with donor and acceptor hydrogen bonds in ice, there are w = 6 local configurations per molecule, n = 4 hydrogen bonds for each O atom, and a fraction $f = \frac{1}{2}$ of donor hydrogen bonds; this recovers Pauling's famous estimate $S = R \ln(\frac{3}{2})$ for hexagonal water ice, which is accurate to within 1% of values from more detailed calculations and experimental measurements.

The two anion distributions found in AMO₂N perovskites (and AMON₂ analogues) are represented by SrTaO₂N and BaTaO₂N. In SrTaO₂N types (including materials such as SrNbO₂N, EuTaO₂N, EuNbO₂N, and CaTaO₂N), [10] the zigzag M-N-M chains are confined to two-dimensional planes [2] Assuming that each possible structure is degenerate, ordered structures such as that in Fig. 1a are possible but statistically unlikely (and have not been observed experimentally), and it is overwhelmingly likely that any randomly generated structure will show disorder, such as that shown in Fig. 1b. This corresponds to a *cis* variant of the square-ice lattice, [11] where M–N/M–O bonds map on to short/long O–H bonds in ice, and only structures in which the

two short O–H bonds are adjacent (cis) to one another are allowed. This variant of the well-known ice model is little explored, although other variants have been devised for ferroelectric ordering in KH₂PO₄[14] and for antiferroelectric orders. The AMO₂N model for SrTaO₂N types has n = 4 bridges per M atom and a fraction of M-N-M connections $f = \frac{1}{2}$, but the cis constraint restricts the number of allowed local configurations to w = 4 (the number of edges of a square) and gives S = 0, as noted previously. Such structures that give S = 0 in the Pauling approximation have subextensive entropies, as described in section (ii).

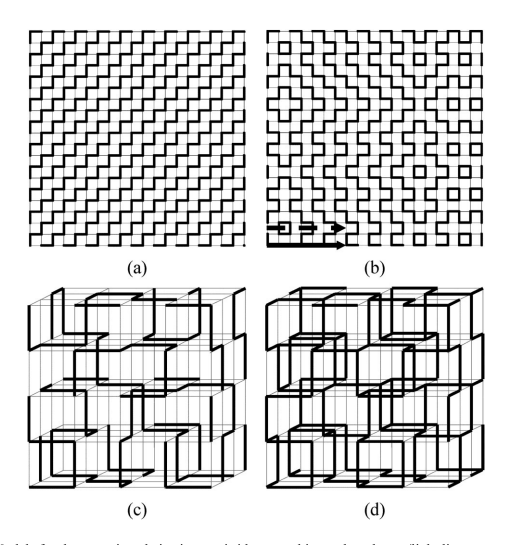


Figure 1. Models for the *cis*-anion chains in oxynitride perovskites, where heavy/light lines correspond to M-N-M/M-O-M connections: (a) long-range ordered and (b) disordered (but open-ordered) configurations of *cis*-MN chains confined to two-dimensional planes within AMO₂N materials such as SrTaO₂N. The solid and dashed arrows on part b, respectively, represent long-range ($\xi \to \infty$) and short-range ($\xi = 0$) anion ordering correlations parallel to the horizontal axis. This structure has subextensive configurational entropy. (c) Three-dimensionally disordered *cis*-MN chains in BaTaO₂N; the fraction of M-N-M connections is 1/3, and the structure has extensive configurational entropy. (d) Cross-linked chains in the AMO_{1.5}N_{1.5} perovskites where

three mutually *cis* M-N-M bridges meet at each M cation and define three-dimensional zigzag patterns on a cubic lattice. The entropy is subextensive, and the structure has three-dimensional open order.

BaTaO₂N does not show deviations from cubic symmetry that would reflect confinement of the N atoms to two-dimensional planes, ^[16] so the zigzag TaN chains are assumed to propagate in all three dimensions (Fig. 1c). Two *cis* M-N-M bridges link each metal ion on a cubic lattice son = 6 and $f = \frac{1}{3}$. The number of ways for an M cation to be linked by two M-N-M bridges in a *cis*conformation is w = 12 (the number of edges of an octahedron). These parameters give an extensive (nonzero) configurational entropy of $S = 2R \ln(\frac{4}{3}) \approx 0.58R$ per mole (where R = 8.314 J K⁻¹ mol⁻¹ is the molar gas constant). Hence, the molar configurational entropies of AMO₂N materials are predicted to vary from zero in the two-dimensional SrTaO₂N limit, to 0.58R in the disordered cubic BaTaO₂N structure. Intermediate situations might be realized by quenching SrTaO₂N from high temperatures where some propagation of TaN chains between planes was observed, ^[2] so that the order is not purely two-dimensional.

In the proposed model for AMO_{1.5}N_{1.5} perovskites such as magnetoresistive EuWO_{1.5}N_{1.5}, [17] and the pigment $(La_{0.5}Ca_{0.5})TaO_{1.5}N_{1.5}$, [18] each M cation is bonded to three nitrogen atoms in a mutual *cis* conformation—also known as the *fac* (facial) configuration of an octahedron—so that the M-N-M bridges define three-dimensional zigzag patterns on a cubic lattice (Fig. 1d). Each M cation has n = 6 neighbors, the fraction of M-N-M bridges is $f = \frac{1}{2}$, the number of ways for M to be linked by x = 3 M-N-M *cis* bridges in a *fac* conformation is w = 8 (the number of faces of an octahedron), and so the Pauling estimate of the entropy is S = 0.

In oxynitride perovskites with z = 0.5, such as SrMoO_{2.5}N_{0.5},^[19] each M cation has one N neighbor. If these are distributed equally in all three dimensions, then there are w = 6 local configurations (the number of vertices of an octahedron) for the connection of an M cation to its n = 6 neighbors; the fraction of M-N-M bridges is $f = \frac{1}{6}$ and the molar configurational entropy is $S = (R/2) \ln(3125/1296) \approx 0.44R$. If the N atoms were confined to planes as in SrMO₂N (M = Nb, Ta), then the parameters would change to n = w = 4 and $f = \frac{1}{4}$, and the molar configurational entropy would be $S = (R/2) \ln(27/16) \approx 0.26R$. Confinement of N atoms to one of the three cubic axes gives n = w = 2 and $f = \frac{1}{2}$, giving another case where S = 0. Detailed neutron studies of anion distributions in AMO_{2.5}N_{0.5} perovskites have not been reported, so there is no experimental evidence for two-or one-dimensional confinements of the N atoms.

The expected evolution of configurational entropy with composition in AMO_{3-z}N_z perovskites is summarized in Fig. 2. The striking feature is the suppression of a measurable entropy near z = 1.5 due to the formation of subextensive states, and at z = 1 and z = 2 when the minority anions are confined to planes, as observed in SrTaO₂N, LaNbON₂, etc. This is contrary to normal expectations for a disordered system such as AMO_{3-z}N_z, where the maximum extensive entropy is expected to occur at the z = 1.5 midpoint. The EuWO_{3-z}N_z series, which has a wide composition range 1.5 < z < 2.2 reflecting variable cation oxidation states, is thus predicted to show two configurational entropy minima at z = 1.5 and 2.0.^[17]

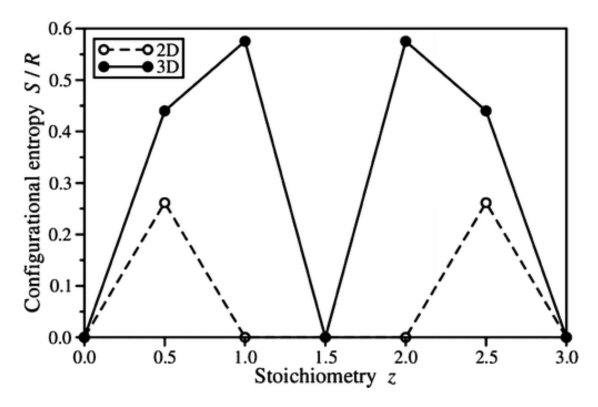


Figure 2. Plot of predicted molar configurational entropy against composition for $AMO_{3-z}N_z$ perovskites. Data are shown per mole of M cations, with the minority anions (N for z < 1.5, O for z > 1.5) constrained in *cis* conformations to two dimensions (2D—open symbols, dashed lines) or to three dimensions (3D—filled symbols, solid lines).

(ii) Subextensive Perovskite Oxynitrides

Several of the above models are predicted to have zero configurational entropy, but it is unclear in the Pauling approach whether a small finite entropy would be obtained from a more detailed treatment. An exact result is obtained by noting that the bonding constraints for structures such as those in Fig. 1b and d generate strictly alternating M-N-M and M-O-M bridges along the rows of the lattice. Hence, there are only two possible sequences for each row. In the SrTaO₂N model where the MN chains are confined to layers as in Fig. 1b, the number of configurations of an $L \times L \times L$ lattice of L layers each containing 2L rows is $W = 2^{2L2}$. For $N = L^3$ M cations, the configurational entropy is $S = k_B \ln W = 2N^{2/3}k_B \ln 2$, where k_B is Boltzmann's constant. This type of entropy is "subextensive", as the exponent of the number of atoms N is less than one, so that the entropy per atom becomes vanishingly small for large N, in asymptotic agreement with the Pauling result. Particles containing N formula units have a configurational entropy of $S = (2R \ln 2)/N^{1/3}$ per mole of SrTaO₂N, showing that the entropy of such a subextensive material is strongly dependent on the particle size. A single crystal containing one mole ($N = N_A$, the Avogadro number) of SrTaO₂N has $S \approx 10^{-7}R$, which would not be measurable, but a powder of 40 nm nanoparticles, each containing $N \approx 10^6$ formula units, has a significant configurational entropy of $S \approx 0.1R$ per mole of SrTaO₂N.

The configurational entropy of the AMO_{1.5}N_{1.5} perovskite model is similarly subextensive. Here the M-N-M bridges define three-dimensional zigzag patterns on a cubic lattice, and these bonding constraints imply alternating M-N-M and M-O-M bridges along any row of the cubic lattice. Following the previous argument, the exact number of configurations is $W = 2^{3L2}$ for $3L^2$ rows in an $L \times L \times L$ cubic lattice and so $S = 3N^{2/3}k_B$ ln 2, which is again subextensive. Hence, the molar configurational entropy of EuWO_{1.5}N_{1.5}, like that of SrTaO₂N, is predicted to be strongly dependent on the particle size. Confinement of N atoms to one of the three cubic axes in AMO_{2.5}N_{0.5} perovskites would also create subextensive states, although no experimental evidence for this partial anion order has been reported.

(iii) Open Order

The unusual nature of the anion ordering in oxynitride perovskites is illustrated by the subextensive configurational entropies predicted above. Subextensive phases form an intermediate category of translationally ordered matter, between perfectly ordered crystals such as β' -CuZn (β' -brass) and quasicrystals, which ideally have zero configurational entropy, and disordered crystals, such as the random alloy β -CuZn (β -brass), which have extensive configurational entropies. The configurational entropies of subextensive materials are practically zero in the macroscopic limit, but they appear crystallographically disordered. Disordered packings of well-ordered chains or layers (e.g., random stacking of close-packed slabs is observed in many intercalation compounds such as the battery cathode material LiCoO₂) also have subextensive entropies, but the unusual nature of the oxynitrides is apparent from the crystal directions in which atoms are correlated. The model for oxynitride planes in SrTaO₂N (Fig. 1b) shows that M-N-M and M-O-M bridges alternate perfectly along vectors in the vertical and horizontal directions but order over only a few unit cell lengths in other in-plane directions defined by sums or differences of the vertical and horizontal vectors. This is expressed below using correlation lengths ξ , where the probability that atoms at two sites separated by distance d satisfies a particular ordering rule is proportional to $\exp(-d/\xi)$.

Perfect crystals of linear dimension La (where a is the cubic lattice spacing) have long-range order with $\xi \gg La$ (written here as $\xi \to \infty$) in all lattice directions, whereas disordered crystals have zero or short-range correlations $\xi \ll La$ (written as $\xi = 0$) in all directions. The correlation lengths in all possible [UVW] directions (for vector $U\mathbf{a} + V\mathbf{b} + W\mathbf{c}$, where \mathbf{a} , \mathbf{b} , and \mathbf{c} are the unit cell vectors and U, V, and W are integers) fall into $\xi \to \infty$ and $\xi = 0$ sets. For perfectly ordered crystals such as β' -CuZn, all [UVW] vectors are in the $\xi \to \infty$ set and the $\xi = 0$ set is empty. In mathematical terminology, these sets are "closed" under addition or subtraction operations, as all sums or differences of [UVW] vectors fall into the same $\xi \to \infty$ set as the [UVW] vectors themselves, and the null $\xi = 0$ set is also closed under these operations. The two sets are also closed under addition or subtraction operations for disordered structures such as that of β -CuZn, as all [UVW] vectors fall in the $\xi = 0$ set.

A full classification of the subextensive states for two-state (Ising) orderings (e.g., of O/N at anion sites in the oxynitride models in section (ii)) along the crystal axes is shown in Table 1. Perfectly ordered alternating chains parallel to the axes can be formed in up to c = 3 directions. Atomic planes defined by two of these chain-ordered axes may be fully ordered, so the maximum possible number of ordered planes p is ${}^{c}C_{2}$. For c = 2 axes, only the plane defined by these axes may be ordered; for example, only the xy plane for chains parallel to the x and yaxes, but for c = 3 directions, up to three planes (xy, xz, and yz) may be ordered. The p = 3 condition implies the full long-range order of a perfect crystal.

$c\mu$	$S/k_{\rm B} \ln 2$	ordered ($\xi \to \infty$) set closure	disordered ($\xi = 0$) set closure	atomic and displacive examples
		closed (null)	closed	β-CuZn
		closed	open	$VO(H_2AsO_4)_2$
		open	closed	SrTaO ₂ N
		closed	open	LiCoO ₂
		open	closed	$EuWO_{1.5}N_{1.5}$, C - $BaTiO_3$
	$N^{2/3} + N^{1/3}$	open	closed	T - $BaTiO_3$
		open	closed	O-BaTiO₃
33	30	closed	closed (null)	β'-CuZn, <i>R-BaTiO</i> ₃

Table 1. Types of Translationally-Ordered Structures, Classified by the Number of Orthogonal Axes along Which Long-Range Ordered ($\xi \to \infty$) Chains Are Oriented (c), and the Number of Long-Range Ordered Planes Containing These Chains (p). The configurational entropy in each case is shown for Ising-like systems of N atoms, each having two possible states (e.g. oxide/nitride site occupation). The open or closed nature of the sets of [UVW] lattice vectors with correlation length $\xi \to \infty$ or $\xi = 0$ is indicated (see text). The c = p = 0 (top row) and c = p = 3 (bottom row) cases correspond to normal disordered and ordered crystals with extensive and zero configurational entropies, respectively. The intermediate cases have subextensive entropies, and the unusual oxynitrides also have open sets of $\xi \to \infty$ vectors ("open order"). Examples of atomic and displacive orderings are shown in roman and italic text, respectively.

Structures with subextensive entropy have $\xi \to \infty$ in some directions within the crystal and $\xi = 0$ in others, and one of the two sets of [UVW] vectors is not closed (hence "open") under addition or subtraction operations. Disordered stackings of well ordered chains or planes have an open $\xi = 0$ set. For instance, when c = 2 and p = 1, long-range ordered layers stacked randomly in the z direction have all [UV0] vectors in the $\xi \to \infty$ set, so this is closed, but the $\xi = 0$ set is open, e.g. because the subtraction of [101] from [201] gives [100] in the $\xi \to \infty$ set. However, the subextensive oxynitrides fall into a separate class where the $\xi \to \infty$ set is open; for example, for xy-plane oxynitride layers such as those in Fig. 1b with c = 2 and p = 0, there are $\xi \to \infty$ vectors parallel to [100] and [010] but not to [110] or $[\overline{110}]$. $\xi = 0$ vectors also run parallel to [100] and [010], as shown in Fig. 1b, and in all other directions, so the $\xi = 0$ set contains all [UVW] vectors and is thus closed.

The above set closure conditions are usefully shortened to "closed/open order/disorder", where "closed/open" refers to whether or not a set of [UVW] vectors is closed under addition or subtraction operations, and "order/disorder" refers to the $\xi \to \infty/\xi = 0$ sets. Hence, perfectly ordered crystals and random alloys show both closed order and closed disorder. Subextensive states have open order or disorder; those based on random stackings of well-ordered chains or layers have open disorder, but the oxynitride structures are distinctive from other types of translationally ordered structure by having open order.

Although subextensive atomic orders are rare in simple solids, analogous displacive orders are better-established and provide useful comparisons. For example, the formation of VO^{2+} vanadyl groups within chains of corner-linked VO_6 octahedra gives rise to randomly oriented $O^{--}V^{-}O^{$

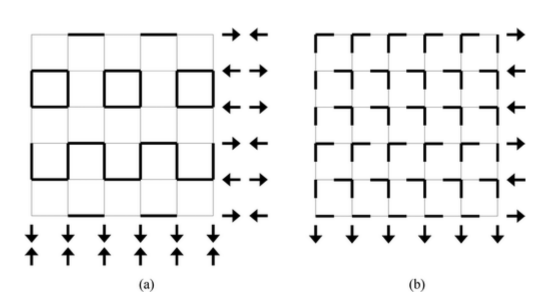


Figure 3. Illustrations of the structural analogy between correlated anions in AMO_{1.5}N_{1.5} and atomic displacements in ferroelectric perovskites: (a) an oxynitride perovskite plane showing M-N-M/M-O-M connections as heavy/light lines; (b) an analogous ferroelectric plane in BaTiO₃, obtained by replacing N-M-O connections from left to right and top to bottom in part a with O- - -Ti-O connections in part b. Short Ti-O/long Ti- --O bonds are shown as heavy/light lines. Both structures have subextensive configurational

entropies. The correlated displacive order in part b gives rise to a net electrical polarization for each plane as displayed by the arrows, whereas in part a the chains are antiferroelectric but the relative order of dipoles is analogous. The drawn configurations have long-range order between the vertical chains but zero or short-range correlations between successive horizontal chains. Part b illustrates the order in the intermediate orthorhombic and tetragonal phases of BaTiO₃, where chain polarizations are long-range ordered parallel to two and one of the three perovskite axes, respectively.

BaTiO₃ undergoes a series of transitions between phases of different symmetry on heating: rhombohedral (R) \rightarrow orthorhombic (O) \rightarrow tetragonal (T) \rightarrow cubic (C). The first three structures are ferroelectric while the cubic phase is paraelectric, and they have been described using an order–disorder model of dipole chains.^[21] In the limit that the dipole correlations within chains parallel to the three cubic axes (c = 3 in Table 1) are of longrange, as proposed in a recent computational study, ^[22] the sequence of phase transitions corresponds to decreasing the number of ordered planes: $p = 3 \rightarrow 2 \rightarrow 1 \rightarrow 0$ on heating. The p < 3 phases have subextensive configurational entropies, and the measured BaTiO₃ transition entropies are found to be consistently small (0.02R-0.06R); ^[23] the recent study suggested that changes in the phonon spectrum are the main contribution. ^[22] This relationship raises the question of whether ordered phases of AMO_{1.5}N_{1.5} analogous to the three ferroelectric phases of BaTiO₃ can be formed (see Fig. 3). Achieving long-range order of anions is a future challenge for perovskite oxynitride chemistry—careful high-temperature annealing studies will be needed to investigate this possibility.

Discussion

Translationally ordered matter falls into three classes on the basis of the exponent s for the variation of configurational entropy with number of atoms, $S \sim N^s$. Classical ordered and disordered crystals respectively have s = 0 and 1, and the intermediate class of subextensive matter described above has 0 < s < 1. Subextensive phases represent the most highly correlated states of matter that are possible without adopting a full long-range ordered crystal structure. Within the subextensive category, open atomic order is unusual, as it requires atoms to participate in intersecting well-ordered chains or layers without generating long-range order in intermediate directions. The two-dimensional AMO₂N (Fig. 1b) and three-dimensional AMO_{1.5}N_{1.5} (Fig. 1d) structures are canonical examples of atomically open-ordered arrangements. In the latter model, each M atom lies at the intersection of three perfectly ordered N-M-O-M-N-M-O-M-N chains parallel to the cubic axes, but no long-range order arises in any other direction. Open-ordered structures arise spontaneously in these oxynitrides due to the strong preference for *cis*-N-M-N bonding. We note that molecular chemistry can be used to synthesize larger units that possess the same geometric characteristics, such as *cis* isomers of

square planar MX_2Y_2 complexes, or fac- MX_3Y_3 octahedra, and open-ordered packings may result when either X...Y or X...X and Y...Y intermolecular interactions dominate.

Direct magnetic analogues of the atomic and displacive open orders on simple cubic lattices are not known and present a challenge for the design of new magnetic systems. Open spin orders on frustrated lattices have been reported; for example, antiferromagnetic Ising models (AFMIMs) on a honeycomb lattice^[24] and on an elastic triangular lattice^[25] have subextensive entropies which scale as $N^{1/2}$ per layer (equivalent to the $N^{2/3}$ scaling for the SrTaO₂N-type structure in Results section (ii), where stacks of layers were considered).

Throughout this work it has been assumed that the energies of all configurations are equal, so configurational contributions to the free energy are insufficient to drive an ordering transition on cooling. A recent study of the elastic AFMIM identified an "order by disorder" mechanism by which the degeneracy of the ground states may be lifted, leading to a stabilization of partially disordered zigzag arrangements of ferromagnetically aligned spins stabilized by the phonon contribution to the free energy. [24] In the unfrustrated perovskites displaying open anion or displacive orders, Coulombic forces are likely to drive the system to a full long-range closed order, as observed in the low-temperature rhombohedral form of BaTiO₃.

Many other atomic variables such as ionic charges in mixed-valent materials and orbital states (and their consequent Jahn–Teller displacements) can show order/disorder phenomena on periodic lattices. Open charge orders could be of interest in the context of charge fluctuation mechanisms for superconductivity in doped copper oxides. Arrangements of larger objects such as colloidal particles in optical crystals^[26] can also show classical ordered and disordered states, and again it is intriguing to consider whether open orders could be achieved on the mesoscale and what the resulting properties might be.

Conclusions

 $AMO_{3-z}N_z$ perovskite oxynitride structures provide interesting examples of ice-type disorder on square or cubic lattices. The local cis-coordination rule gives rise to predicted subextensive entropies in materials such as $SrTaO_2N$, $LaNbON_2$, and $EuWO_{1.5}N_{1.5}$. The subextensive structures are usefully classified as showing an "open order", related to the correlated local order of displacements in ferroelectric perovskites such as $BaTiO_3$.

The authors declare no competing financial interest.

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