Cube-shaped diffuse scattering and the ground state of BaMg$_{1/3}$Ta$_{2/3}$O$_3$

A. Cervellino,¹,²,³ S. N. Gvasaliya,²,³ B. Roessli,² G. M. Rotaru,² R. A. Cowley,⁴ S. G. Lushnikov,⁵ T. A. Shaplygin,⁵ A. Bossak,⁶ and D. Chernyshov⁷

¹Swiss Light Source, Paul Scherrer Institut, CH-5232 Villigen, Switzerland
²Laboratory for Neutron Scattering, PSI, CH-5232 Villigen, Switzerland
³Neutron Scattering and Magnetism Group, Institute for Solid State Physics, ETH Zürich, Zürich, Switzerland
⁴Clarendon Laboratory, Department of Physics, Oxford University, Parks Road, Oxford, UK
⁵Ioffe Physical Technical Institute, 26 Politekhnicheskaya, 194021 St. Petersburg, Russia
⁶European Synchrotron Radiation Facility, Bole Postal 220, F-38043 Grenoble Cedex, France
⁷Swiss-Norwegian Beam Lines at ESRF, 38043 Grenoble, France

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A quite unusual diffuse scattering phenomenology was observed in the single-crystal x-ray diffraction pattern of cubic perovskite BMT (BaMg$_{1/3}$Ta$_{2/3}$O$_3$). The intensity of the scattering is parametrized as a set of cubelike objects located at the centers of reciprocal space unit cells, resembling very broad and cubic-shaped (1/2,1/2,1/2) satellites. BMT belongs to perovskites of formula $AB'_{1/3}B''_{2/3}O_3$ ($A = $ Mg, $B' = $ Ta, $B'' = $ Mg). The cubes of the intensity can be attributed to the partial correlations of the occupancies of the $B$ site.

I. INTRODUCTION

An experiment using x-ray single-crystal diffraction at the Swiss-Norwegian Beam Lines (SNBL) of the European Synchrotron Radiation Facility (ESRF) on a single crystal of cubic perovskite BMT (BaMg$_{1/3}$Ta$_{2/3}$O$_3$) has shown an interesting diffuse scattering pattern, consisting of cube-shaped domains of nearly constant intensity, with the centers located on the (1/2,1/2,1/2) points of the reciprocal lattice and with an edge of ~1/3 of a reciprocal lattice unit (rlu). All the cubes have the same intensity level, apart from the usual slow modulations due to atomic and thermal factors.

BMT belongs to perovskites of formula $AB'_{1/3}B''_{2/3}O_3$ ($A = $ Ba, $B' = $ Ta, $B'' = $ Mg) where different species on the $B$ site can be ordered to various degrees. Detailed discussion of the properties of BMT can be found in Ref. 1. The scattering depends on the site occupancy and the ordering is directly related to the pair correlation function of the different atomic sites.

Lattice deformations and displacements, due to different atomic radii of Mg and Ta (see Ref. 3), are expected. So there is additional diffuse scattering originating from these correlated displacements. In fact streaklike diffuse scattering—relatively weaker—has been found at the foot of the cubic Bragg peaks; see Fig. 1. We shall address also this point in a future paper.

The simplest possibility for $B$-site ordering is the formation of subregions enriched in either Mg or Ta cations. This possibility, however, has been thoroughly ruled out as energetically unfavorable in $AB'_{1/3}B''_{2/3}O_3$ perovskites because of the large charges in the subregions. There are then different homogeneous ways of ordering the $B'$-$B''$ cations. These are (i) homogeneous random distribution of $B$ cations; (ii) correlated distribution with occupation numbers being a well-defined function of distance between two lattice nodes, but without periodicity; (iii) periodic long-range ordering of the Mg/Ta concentrations in different sublattices.

Situation (i) would give an almost wave-vector-independent diffuse scattering, but this is not observed. $B$-site ordering of type (iii) has been observed in BMT, with simultaneous lowering of the cubic symmetry. In fact, BMT may exist in an ordered trigonal structure ($PTrm1$) modification and it has been shown that this modification mainly entails perfect periodic ordering of the $B$ site. The trigonal lattice is essentially a threefold superlattice of the cubic one, with extremely small geometric distortions, realized by stacking periodically a sequence of one Mg plane and two Ta planes orthogonally to one of the cubic $(1,1,1)$ directions. The atomic sites are also very close to those of the underlying cubic sublattice, with only small atomic displacements consistent with the reduced symmetry. It is noteworthy that the trigonal BMT superstructure—usually cast unto four equally populated domain orientations with the threefold axis along either of the cubic $(111)$ directions, so that the average symmetry is conserved—can be considered as a commensurate distortion of cubic BMT. In this framework, the modulation contributes eight additional $(±1/3,±1/3,±1/3)$ satellite Bragg peaks to the diffraction pattern, and these coincide with the vertices of the diffuse scattering cubes. Another partial $B$-site ordering type of class (iii) has been observed in other $AB'_{1/3}B''_{2/3}O_3$ cubic perovskites (PMT, with $A = $ Pb, $B' = $ Ta, $B'' = $ Mg).

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partial ordering energetically favored by certain features of the Fermi surface.\textsuperscript{9,10} Similar effects should not occur in BMT, which is an excellent dielectric.

We stress here that the observed cubes of diffuse intensity are not ideal, as careful inspection (see Fig. 1) shows that “cubes” have a minimum in the center about 10% lower than at the periphery and the corresponding intensity distribution does not have a flat top. We see in particular that intensity tends to increase near the vertices [the \((\pm 1/3, \pm 1/3, \pm 1/3)\) satellites of the trigonal superstructure] and decrease in the center [the \((1/2,1/2,1/2)\) satellite of the hypothetical partly ordered NaCl-type superstructure], indicating an embryo of transition to the trigonal phase. We must therefore consider the phase with idealized cubes as a limiting situation that may not be perfectly realized in our sample. As we argue in the following that the phase with ideal diffuse cubes is not the ground state at room temperature but only a metastable phase, kinetically hindered from transforming to the trigonal ground state, this is perfectly possible, as a germ of transition may have occurred due to finite quenching speed. However, in this framework, the limiting phase with ideal diffuse cubes is a key to understanding the \(B\)-site ordering phenomena in BMT, and therefore we will deal mainly with this phase.

A. A few remarks

As we are discussing the multiple phases of BMT, we should comment on an important point about phase stability. In the thermodynamic sense, a stable structure observed in an experiment corresponds to a minimum of the free energy at the conditions of crystal formation. At ambient conditions the Mg-Ta distribution is quenched and experimentally observed configurations may not necessarily correspond to a true ground state. A disordered or incompletely ordered state can be stabilized at high temperature by a strong entropy term and could be a metastable state at lower \(T\). In fact, at room temperature the diffusion of Mg and Ta is practically negligible, and this in turn decreases the free energy, \(G \approx U - TS\), to the advantage of the stability of the (statistically defined) structure. We intend to show that the \(B\)-site order in BMT, which gives rise to the cubes of

![FIG. 1. (Color online) The diffuse scattering observed for sections of reciprocal space for BMT. Top left: The (100)-(001) section through the origin of the reciprocal space, showing weak diffuse streaks (here magnified) of displacive origin. Top right: The (110)-through the origin of the reciprocal space, showing weak diffuse emissions of reciprocal space for BMT. Top left: The (100)-(001) section largely concentrated in the range from \((h+1/3,k+1/3,l+1/3)\) to \((h+2/3,k+2/3,l+2/3)\).](image-url)
the diffuse intensity, belongs to this case, based on the fact that pair (electrostatic) interactions between the $B$ cations are sufficient to explain the phase diagram of the different BMT phases which derive from the ordering of $B$ sites.

The pair (two-body) and the set of the higher order (many-body) correlation functions can fully describe any structure in an unambiguous way, including stochastically defined ones.2,15 Considering higher order correlation functions is necessary, however, only when they make important contributions to the internal energy. This is sometimes the case, for instance, for structures that show displacive diffuse scattering.16–18 In any case, higher correlation functions are also a way of describing a structure in the crystal-chemical sense. This is especially convenient when the structure itself cannot be easily represented because it is only statistically defined. However, simpler ways of presenting the crystal-chemical information are also possible; see Sec. III G.

Our approach is as follows. The pair correlation function that is related to the diffuse cubes can be simply analyzed in terms of the $B$-site chemical order as a site concentration pair correlation function.2 This turns out to have a simple analytical form that shows a rather peculiar power-law decrease with distance that is observed in only a few cases for rather exotic systems.

Monte Carlo simulations were performed for a large scale in order to find the $B$-site configurations that give the same pair correlation as the diffuse intensity. Successfully determined configurations were then analyzed in terms of a long-range order parameter showing that these partially ordered configurations that are compatible with the observed scattering phenomena are not distinguishable from completely disordered systems. Next, we consider that BMT is an ionic compound and that the electrostatic energy is the main term in the crystal Hamiltonian. The electrostatic energy of a rigid system depends only on pair interactions and the electrostatic energies and the entropy terms for the different BMT phases were calculated. These gave a good correspondence with the observed phase diagram, confirming that the cubic phase must be a high-temperature ground state of BMT that is metastable at room temperature.

II. EXPERIMENTAL

The single crystals of BMT were synthesized according to the procedure described by Galasso and Pinto.19 The powder of BMT was prepared according to two-stage synthesis described in Ref. 20 and is essentially the same as used in the neutron diffraction study.21

The experiments were performed on x-ray scattering instruments at the ESRF and SLS. At the ESRF the diffuse scattering from BMT was measured at the Swiss-Norwegian Beam Line (SNBL) and the XMAS beamline, and at the SLS the measurements were made at the beamline X04SA. More details of the measurements and the experimental results for BMT are given in Ref. 1. In this paper we concentrate on the unusual diffuse scattering observed from BMT and typical results are shown in Fig. 1. Similar results were obtained from the SLS diffractometer X06SA, but from the XMAS data the scattering contained sharp peaks at the $(H + 1/3, K + 1/3, L + 1/3)$ positions showing the structure was trigonal. A BMT powder sample was also studied at the SLS on the beamline X04SA. It was revealed that the trigonal satellites were significantly broader than the intense peaks of the cubic phase showing that the domain size was about 20 nm for the trigonal structure. These results support the suggestion that our powder and the surfaces of relatively large BMT crystals consist to some extent of trigonal-structured domains while the bulk of crystals is cubic and shows the diffuse scattering of Fig. 1.

This paper concentrates on the discussion and modeling of the unusual cube-shaped distribution of scattering and on how the different structures of BMT can exist. It is suggested that the structures are unable to relax when rapidly cooled from high temperature.

BaMg$_{1/3}$Ta$_{2/3}$O$_3$ is cubic with lattice parameter $a = 0.404$ nm. The $A$ cation (Ba) is located within the unit cell in $(1/2,1/2,1/2)$, the mixed $B$ site is in $(0,0,0)$, and the three oxygens occupy the $(0,1/2,1/2), (1/2,0,1/2), (1/2,1/2,0)$ positions.

III. IDEALIZED MODEL OF DIFFUSE INTENSITY

In this section a simple mathematical model of the diffuse intensity is described. The model is simple because each atom is assumed to be a point scatterer, with element $X_X$ having a constant scattering length $b_X$. This means neglecting the atomic form factors and Debye-Waller factors, whose effect is just a smooth decay of the intensity at large transferred momentum values.

A. Idealized sharp cubes

The scattering from BMT has diffuse scattering cubes centered at $q = (h,k,l) + (\frac{1}{2},\frac{1}{2},\frac{1}{2})$ with half side 1/6 (all in rlu). A simple mathematical description of their shape can be given as follows. In one dimension, define a box function

$$B_{w}(x) = \begin{cases} 1 & \text{if } |x| < w, \\ 0 & \text{otherwise.} \end{cases}$$

Then a periodic sequence of such boxes centered on the half integers is

$$\Phi_{w}(x) = \sum_{n=-\infty}^{+\infty} B_{w}(x - n - 1/2).$$

Multiplying three such terms (one for each coordinate) and adding a constant background (that we will show to be necessary) we obtain a model of the x-ray diffuse intensity:

$$I_{\text{diff}}(q) = I_{B} + I_{0} \Phi_{1/6}(q),$$

where

$$\Phi_{1/6}(q) \equiv \begin{cases} 1 & \text{within the cubes} \\ 0 & \text{outside} \end{cases}$$

$$= \prod_{a=1}^{3} \sum_{h_a=-\infty}^{+\infty} B_{1/6}(q_a - h_a - 1/2).$$
Here $I_B$ is the level of a constant background, $I_0$ is the intensity within the cubes above the background, and $w = 1/6$ is the half-width of the observed cubes in reciprocal lattice units. Taking the Fourier transform of the periodic array of cubes of x-ray diffuse intensity $I_{\text{dif}}(q)$ [Eqs. (2) and (3)], we evaluate the pair correlation function as

$$G_{\text{dif}}(r) = \int \frac{d^3q}{3\pi^2} e^{-2\pi i q \cdot r} I_{\text{dif}}(q).$$

Note that as $I_{\text{dif}}$ is periodic its Fourier transform is necessarily a weighted set of Dirac deltas on the direct lattice.\(^{35}\)

$$G_{\text{dif}}(r) = \eta^2 \sum_M S_M \delta(r - M),$$

where $\eta^2$ is a positive constant related to the actual scattering lengths [see Sec. III C, Eq. (10)] and $S_M$ are the correlation coefficients. In fact, substituting $I_{\text{dif}}$ from Eqs. (2) and (3), we obtain

$$G_{\text{dif}}(r) = I_B \delta(r) + \frac{I_0}{2\pi} \sum_M \delta(r - M) \prod_{a=1}^3 \left[ (-1)^m \sin \left( \frac{\pi m_\alpha}{m_\alpha} \right) \right],$$

whereas the correlation coefficients are given by

$$S_M = \frac{I_0}{2\pi \eta^2} \prod_{a=1}^3 \left[ (-1)^m \sin \left( \frac{\pi m_\alpha}{m_\alpha} \right) \right],$$

$$M \neq (0,0,0);$$

$$S_{(0,0,0)} = \frac{27I_B + I_0}{27\eta^2}. \quad (6)$$

Here $M = (m_1, m_2, m_3)$ are direct lattice nodes, where the density (and the pair correlation function) are different from zero.

The distance $r = [m_1^2 + m_2^2 + m_3^2]^{1/2}$, so that $G_{\text{dif}}(r) \sim r^{-\alpha}$, with $1 \leq \alpha \leq 3$, depending on the direction. This is a very rare case of power-law correlation decay of structural origin. In quite exotic systems, including spin glasses and nematic crystals, such correlations have been described\(^{22,23}\) and named as quasi-long-range order (QLRO). We shall name QLRCO (as quasi-long-range cation order) the kind of order exhibited by the BMT sample. Two parameters are often used to characterize the degree and extent of the structural chemical order. The first is the long-range order parameter of Refs. 24 and 25; see Appendix A3. The value of the long-range order parameter $X = 1$ characterizes periodically ordered structures, as the type (iii) above, while zero is the value for randomly ordered structures, as type (i). Conversely, the short-range order parameters of Ref. 26 characterize the local degrees of order—supposedly when long-range order is absent. We obtain (see Secs. III G1 and III G2) that both long- and short-range parameters indicate a substantial degree of order, which is an unusual occurrence.

**B. Smoothed cubes**

The model of the diffuse intensity described above can be improved in one significant aspect. The cubes as observed are not perfect, as there are intensity fluctuations within them. Their edges, especially, are not very sharply defined. This effect can be included by convoluting $\Phi(q)$ with a point-spread function (PSF) $D(q)$. Accordingly, the pair correlation function $G_{\text{dif}}(r)$ is multiplied by the Fourier transform of the PSF, $\tilde{D}(r)$, as shown in Fig. 2. The latter will be named here the correlation decay function (CDF). This will give a stronger decay law for the pair correlation function. This is just what happens with every known diffraction phenomenon—as far as infinite crystals do not exist, even the most perfect standard crystal will show Bragg peaks convoluted by a suitable PSF and pair correlations damped by the corresponding CDF.

We will not, in the following, burden the notation by including a PSF, but we will calculate some of its effects in Sec. IV A. Different possible PSFs are discussed in Appendix A1. Here, as an example, we consider a Lorentzian PSF along the cubic axes (Fig. 2):

$$D(q) = (\pi k_0)^{-3} \prod_{a=1}^3 \left[ 1 + (q_a/k_0)^2 \right]^{-1}. \quad (7)$$

Its Fourier transform is

$$\tilde{D}(r) = \prod_{a=1}^3 e^{-2\pi i q_a k_0}. \quad (8)$$

The correlation decay length is $L = 1/(2\pi k_0)$. From our data, taking $k_0$ as the rise of the half-width of the cube edges, we can coarsely estimate that $L$ is about 10 unit cells. This is well below the coherent domain size; however, it remains a considerable distance when dealing with diffuse scattering.

As the PSF $D(q)$ is convoluted with the scattered intensity, the CDF $\tilde{D}(r)$ is multiplied by the pair correlation function, which is the Fourier transform of the intensity. Its effect is then to limit the spatial extension of the pair correlation function. This is a familiar issue in crystallography. Crystal size can be estimated from the broadening of the Bragg peaks, due to the limited extension of the crystal—and, in turn, of the pair correlation function.

We wish to remark here that the “cube” scattering is quite different from Bragg scattering with respect to its response to a limitation of correlation length. In fact, convoluting the PSF with a Bragg peak gives the PSF itself. This is the reason for the familiar behavior of Bragg peaks, namely, that their width

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**FIG. 2.** (Color online) Effect of a Lorentzian PSF on the cube shape in one dimension. Black, the sharp cube profile; blue, the PSF; red, the smoothed cube profile.
is inversely proportional to the size of the correlated domain (a good estimate of this is $L$). The diffuse cubes, however, show a negligible change in width, second-order or more in $1/L$. This scattering form is therefore fundamentally different from Bragg scattering.

**C. B-site order and diffuse cubes**

To explain such diffuse scattering in terms of B-site cation order we split the correlation density $\rho(r)$ as

$$\rho(r) = \rho_A(r) + \Delta \rho(r),$$

with $\rho_A$ the periodic average and $\Delta \rho$ a nonperiodic zero-average fluctuation. If the $B$ site is the only one that varies, all of Ba and O partial densities are included in $\rho_A$. As already mentioned, we simplify the picture by considering pointlike atoms. Then $\Delta \rho(r)$ is zero everywhere except on the $B$ sites, and for $B = Ta$, $\Delta \rho(r) = \eta$; for $B = Mg$, $\Delta \rho(r) = -2\eta$. If $b_{Ba}$ and $b_{O}$ are the scattering lengths of Mg and Ta, respectively, then

$$\eta = (b_{Ta} - b_{O})/3.$$ 

As $\Delta \rho$ assumes only two values ($\eta$ on Ta sites and $-2\eta$ on Mg sites) we can consider (see Ref. 2) the site concentration, which is 1 if site $M$ is occupied by Mg and 0 otherwise. Similarly we take $c_M^a$ as the Ta local concentration, which is 1 if $M$ occupied by Ta and 0 otherwise. As we have no vacancies, $c_M^a = 1 - c_M$. Averaging over all sites, we have

$$\langle c \rangle = 1/3 \quad \text{and} \quad \langle c' \rangle = 2/3.$$ 

Furthermore, as $c$ and $c'$ can be only 0 or 1,

$$\langle c^2 \rangle = \langle c \rangle = 1/3 \quad \text{and} \quad \langle c'^2 \rangle = \langle c' \rangle = 2/3,$$ 

so the rms fluctuation of the concentration is given by

$$\delta c = \sqrt{\langle c^2 \rangle - \langle c \rangle^2} = \sqrt{2}/3.$$

The fluctuation of the density is given by

$$\Delta \rho(r) = \eta \sum_M (3c_M^a c_M^b) \delta(r - M)$$

$$= \eta \sum_M (1 - 3c_M) \delta(r - M).$$

**D. Correlation function**

In order to study the diffuse scattering from BMT, we need to evaluate the pair correlation function of $\Delta \rho$:

$$G_{\text{diff}}(r) = \frac{1}{\Omega} \int_{\mathbb{R}^3} d^3r' \Delta \rho(r') \Delta \rho(r' + r)$$

$$= \frac{\eta^2}{\Omega} \sum_M S_M \delta(r - M).$$

Here

$$\Omega = N v_c = N$$

is the crystal volume, $N$ the number of lattice nodes within it, $v_c$ the unit cell volume (taken as 1 hereafter, hence the last equality). We shall equate this to the pair correlation function derived from the data; see Eq. (4). The rightmost form is because $\Delta \rho$ is a weighted Dirac lattice, and this property is invariant upon self-convolution. This is the same form as Eq. (4), so clearly now we have to calculate the $S_M$ pair correlation coefficients for a given fluctuation density [Eq. (11)] and to compare them with those obtained from the Fourier transform of the data [Eqs. (5) and (6)].

Substituting Eq. (11) into Eq. (12) gives

$$G_{\text{diff}}(r) = \frac{\eta^2}{\Omega} \sum_M (1 - 3c_M)(1 - 3c_{M + L}) \delta(r - M),$$

where, comparing with Eq. (12), we obtain

$$S_M = -1 + \frac{9}{\Omega} \sum_M c_M c_{M + L} = -1 + 9 \langle c_M c_{M + L} \rangle;$$

$$S_{(000)} = -1 + 9(c^2) = 2.$$ 

Now, the probability of finding a pair of Mg atoms separated by a lattice vector $M$ is given by

$$P_{\text{Mg-Mg}}(M) = \langle c_M c_{M + L} \rangle = \frac{1 + S_M}{9}.$$ 

Similarly we can calculate the probabilities of all the other pairs (Mg-Ta/Ta-Mg and Ta-Ta) for any spacing vector $M$, and the result depends only on the $S_M$ correlation coefficients:

$$P_{\text{Ta-Ta}}(M) = \langle (1 - c_M)(1 - c_{M + L}) \rangle = \frac{1}{3} + \langle c_M c_{M + L} \rangle = \frac{4 + S_M}{9},$$

$$P_{\text{mix}}(M) = \langle c_M(c_{M + L}) \rangle + \langle (1 - c_M)c_{M + L} \rangle = \frac{2}{3} - 2\langle c_M c_{M + L} \rangle = \frac{4 - 2S_M}{9},$$

where the subscript “mix” refers to Mg-Ta pairs, in any order. Therefore, if we know the correlation coefficients we can calculate all the properties that depend on the pair interactions.

**E. Scale and background**

Comparing Eq. (4) with Eqs. (13) and (14), and considering $r = 0$, we obtain

$$S_{(000)} = 2 = \frac{1}{\eta^2} \left[ I_B + \frac{I_0}{27} \right] \Leftrightarrow I_0 = 54\eta^2 - 27I_B.$$ 

We define a parameter $t$, so that if $0 \leq t \leq 1$, then

$$I_B = 2\eta^2(1-t); \quad I_0 = 54\eta^2 t.$$ 

This result describes how much of the total diffuse intensity in one Brillouin zone is a constant background and how much comes to form the cubes of diffuse scattering.

As described by Eq. (15), $(1 + S_M)/9$ is a probability; hence it must range between 0 and 1. In order for it to be positive for all $M$,

$$S_M \geq -1.$$ 

For $M = (1,0,0)$, this constraint—using Eq. (5)—implies

$$t \leq \frac{\pi}{3 \sqrt{3}} \approx 0.6046,$$

which tells us that a consistent fraction—at least about 40%—of the diffuse intensity must be spread as a constant background.
while the rest forms the diffuse cubes. This is also a lower bound only, so possibly the background fraction is even larger.

F. Electrostatic energy as a function of the pair correlation function

In Eqs. (15) and (16) we have calculated the probability of finding pairs of atoms of equal or different species (limited to the B sublattice) for each separation vector $M$. That is all we need to calculate the electrostatic energy. In fact, the structure consists of the fixed part (the A sublattice and the oxygen sublattices) and the variable part (the B sublattice). A result of Ref. 27 is that the electrostatic energy does not depend on the interaction terms between the different sublattices; the fixed part then has no influence except for a constant term and one needs to calculate only the interactions between atoms of the B sublattice. Similarly to the scattering density [Eq. (9)], we can separate the charge density $\rho_e(r)$ into the average part and fluctuating part:

$$\rho_e(r) = \rho_{e,A}(r) + \Delta \rho_e(r),$$

(21)

with $\rho_{e,A}$ the average component and $\Delta \rho_e$ a nonperiodic zero-average fluctuation. Again, on Ta-occupied sites we will have $\Delta \rho_e = \eta_Z$ and on Mg sites, $\Delta \rho_e = -2\eta_Z$ with

$$\eta_Z = (Z_{Ta} - Z_{Mg})/3.$$  

(22)

If we neglect covalent bonding and assume the formal charges $Z_{Ta} = +5, Z_{Mg} = +2$, then we have $\eta_Z = 1$. It can be shown27 that only the fluctuation density $\Delta \rho_e$, which varies between the different possible phases of BMT, needs be included in the calculations. Therefore, the electrostatic energy is simply expressed—using Eqs. (15) and (16)—as

$$E = \frac{e^2 \eta_Z^2}{8\pi \epsilon_0} \sum_M 4P_{Mg-Mg}(M) + P_{Ta-Ta}(M) - 2P_{\text{mix}}(M)$$

$$= \frac{e^2 \eta_Z^2}{8\pi \epsilon_0} \sum_M S_M |M|,$$

(23)

the prime meaning that $M = (0,0,0)$ is excluded from the summation. An additional factor of 1/2 was included in order not to count twice each bond.

G. Monte Carlo solution

At this point we know that a distribution of B-site cations whose correlation coefficients are given by Eqs. (5) and (6) would produce a diffuse intensity as described by Eqs. (2) and (3), which well matches the observed diffuse scattering cubes. We cannot, however, prove analytically that a distribution of two different cations on a cubic lattice with respective abundances of 1/3 and 2/3 exists that can satisfy Eqs. (5) and (6) [or alternatively, produce a diffuse intensity as in Eqs. (2) and (3)]. In order to prove that such a distribution exists, we decided to try to fabricate one or more by a Monte Carlo simulation. In this way we could also get some structural insight by analyzing the configurations so obtained.

A large cube ($27 \times 27 \times 27$ unit cells with the total number of atomic sites $N = 19683$) was placed on the lattice nodes with a “Mg” atom, represented by a scattering length $-2$, or by a “Ta” atom, represented by a scattering length 1. These values are chosen so that the average density [see Sec. III C, Eq. (9)] $\rho_A$ is zero and only the fluctuation density $\Delta \rho$ is not zero; moreover, we set $\eta = 1$ for simplicity. The FFT power spectrum was evaluated from the intensity $I_{gg}(q)$ in one Brillouin zone. The atoms were moved around until a good agreement with Eqs. (2) and (3) was achieved. The simulation was very successful. In Fig. 3(a) a section of the calculated diffuse intensity is shown for one of the optimal configurations found. The result of the calculations compares well with the experimental data [Eqs. (2) and (3)]. Other sections (not plotted) are consistent with the one shown.

1. Pair correlations and long-range order parameter

Several equivalent configurations were constructed as above and used to evaluate the pair correlations. First, the ratio $I_0/I_B$ for the solutions was found, and it resulted that $I_0/I_B \approx 3.0$. This gives a value of the intensity partition parameter $t \approx 0.1$—actually much lower than its upper bound of 0.6. From that, exploiting Eq. (A4) in the Appendix A3, the value of the long-range order parameter was extracted as $\lambda \approx 0.407$. We can then evaluate $I_B = 1.87\eta^2$, $I_0/27 = 0.2\eta^2$. 

FIG. 3. (Color online) (a) The diffuse intensity calculated in one Brillouin zone from the Monte Carlo solution. The result shown in the $h,k,14/27$ plane. (b) Red bars: Correlations $S_M$—for $M = (k,0,0)$—from the Monte Carlo calculation. As allowed by the cubic symmetry, values have been averaged over $M = (\pm k,0,0), (0, \pm k,0),(0,0, \pm k)$, and plotted vs $k$. Blue line: Correlations $S_M$ calculated (on the same $M$ points) using Eqs. (5) and (6). This is plotted as a double-envelope curve (because of the factor $-1^k$) to guide the eye. Green line: Same as the former, multiplied by a CDF exp($-k/L$) as from Eq. (8) with $L = 1/(2\pi k_0) = 10$. 

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any of the equivalent is Mg or Ta, respectively, are shown. In Fig. 4, we plot the fraction where the atom in the origin is Mg or Ta and the neighbor conditions were used. The fractions for all four chemical pairs \( \langle k00 \rangle \) have been calculated as follows. Each site in a cluster with randomly assigned Mg or Ta at each site, fulfilling the choice of the atom at the origin, and amount to 1/2, because \( P_{\text{mix}} \) includes both bond directions, and 3/2, the inverse of the probability of the central atom being Mg. For the same cluster: Red: Mg neighbor fractions for a central Ta atom, \( \langle 3/4 \rangle P_{\text{Ta-Ta}(k00)} \) (here the factor 3/4 is obtained as a product of 1/2, because \( P_{\text{mix}} \) includes both bond directions, and 3/2, the inverse of the probability of the central atom being Ta). Green: Ta neighbor fractions for a central Mg atom, \( \langle 3/2 \rangle P_{\text{Mg-Mg}(k00)} \). Brown: Ta neighbor fractions for a central Ta atom, \( \langle 3/2 \rangle P_{\text{Ta-Ta}(k00)} \). For a cluster with randomly assigned Mg or Ta at each site, but with the exact composition \( \text{Mg}_1/3 \text{Ta}_{2/3} \), we plot in black the Mg neighbor concentrations of any central atom; in violet, the Ta neighbor concentrations of any central atom. These values can be calculated as above, simply substituting the short-range order parameter for the \( k \) side.

**Figure 3(b)** shows the correlations \( S_M \) from the Monte Carlo compared with those evaluated above, for \( M \) along the family of \((1,0,0)\) directions. The good agreement with values calculated from Eqs. (5) and (6). with and/or without an additional PSF with \( L = 10 \), shows more quantitatively that the Monte Carlo converged simulation converged towards the observed pattern.

**2. Crystal chemical information and short-range order**

In order to display some crystal-chemical information in the QLRCO configurations obtained by Monte Carlo, we directly evaluated the probabilities that a Mg (respectively, Ta) atom has a Mg (respectively, Ta) neighbor at a distance \( k \) along any of the equivalent \((1,0,0)\) directions. The results are shown in Fig. 4 and have been calculated as follows. Each site in a cluster is treated as the central atom. The fraction of atoms of a given species over the neighbors having distance \( k \) along the family of \((1,0,0)\) directions was calculated. Periodic boundary conditions were used. The fractions for all four chemical pairs where the atom in the origin is Mg or Ta and the neighbor is Mg or Ta, respectively, are shown. In Fig. 4, we plot the Ta-Ta, Ta-Mg, Mg-Ta, Mg-Mg \((k00)\) chemical pair fractions for the QLRCO cluster. For comparison we add the values for a cluster with randomly assigned Mg or Ta at each site, fulfilling a composition \( \text{Mg}_{1/3} \text{Ta}_{2/3} \). These of course do not depend on the choice of the atom at the origin, and amount to 1/3 when the neighbor is Mg and 2/3 when it is Ta. From the values of the Ta-Mg (or Mg-Ta) chemical pair fractions we immediately obtain the value of the short-range order parameter for the first coordination shell[26,28] for the QLRCO phase. The short-range order parameter \( \alpha(1) \) is found to be \( \approx -0.21 \). For this composition it has a maximum (in absolute value) at \(-0.5\), for the fully ordered state, while it is zero for the random state. Combined with similar results obtained for the long-range order parameter in Sec. III G1 this gives a further justification for the name “quasi-long-range order.”

**IV. FREE ENERGY OF BMT PHASES**

**A. Electrostatic energy calculation**

In this section and the next we shall demonstrate how the electrostatic energy and the configuration entropy, as calculated from the structure and pair correlations for the different possible BMT phases, is sufficient to explain the phase diagram of BMT—in particular, the fact that the QLRCO phase is metastable at room temperature, and the fact that the NaCl phase, although common in similar perovskites, has never been found for BMT.

The electrostatic energy is not the only component of the crystal energy, but has been previously identified as the driving force of the cation ordering in complex perovskites. At least it has been shown[27] that electrostatic energy-based predictions describe quite effectively the ordering behavior of a large class of perovskites. The electrostatic energy was evaluated in a similar fashion as in Ref. 27. As already mentioned in Sec. III F, only the fluctuation term relevant to the B-cation sublattice needs to be calculated, because the other contributions can be considered either as fixed or as zero.

We have used a relative permittivity \( \varepsilon = 1 \) and a cubic cell side \( a = 4 \) \( \AA \) and the formal charges Ta\(^{+5}\), Mg\(^{+2}\), Ba\(^{+2}\), O\(^{-2}\) which gives \( \eta = 1 \). It is easy to scale the results to more realistic values of the parameters later on.

The electrostatic energy was calculated using different ways. In all cases, the zero of the energy was fixed to the energy of a reference perovskite, BaTiO\(_3\), which has a +4-charged cation (equal to the average charge of Mg\(_{1/3}\)Ta\(_{2/3}\)) on the B site. Electrostatic energies were calculated—apart from the reference BaTiO\(_3\) perovskite, labeled “uniform”—for all the different B-site orderings discussed in this paper. The BMT phase (we label it “QLRCO” for conciseness), the structure with NaCl-type partial B-site order (label “NaCl”), the trigonal phase with full B-site order (label “\( P_3m 1 \)”), and the phase having fully random B-site occupancies (label “random”) have been calculated.

As a first method of calculation, we created large clusters of atoms and computed the interactions directly, summing up all atom pair contributions. Calculations of this kind were performed for clusters of increasing size up to \( 27 \times 27 \times 27 \) cubic cells, at which limit the results were well stabilized and appeared to have reached their asymptotic value (with residual oscillations smaller than \( 0.1 \) eV/formula unit). Fractional O- and Ba-site occupancies at the boundary have been used to keep the neutrality of the clusters. This fact, together with the cubic cluster symmetry, explains the very rapid convergence of the electrostatic energy sum.

As a second approach, the Madelung method[29] was used. This gave consistent results within \(<0.1 \) eV/formula unit.
Third, we use the knowledge of the correlation coefficients $S_M$ (see Sec. III D). In this case, the electrostatic energy is simply expressed as in Eq. (23). Such sums were evaluated in all cases from the analytical $S_M$. Again, we evaluated them on cubes of increasing size (up to $500 \times 500 \times 500$ unit cells) in order to verify the convergence—which was always excellent. Calculation times in this case were also extremely fast—less than one minute on a single PC for the largest cubes with $\approx 10^8$ nodes. With this latter method for the QLRCO case, the effect of a PSF spreading the diffuse scattering cubes was also evaluated (see Fig. 5). It turns out that even with small correlation lengths ($L \approx 10$ unit cells) the energy was within a few percent from the value as $L \to \infty$.

For the QLRCO, NaCl, and random cases (which are not deterministic) we have repeated the calculation for many configurations, evaluating the average values and their standard deviations. In all cases the number of trials was chosen high enough for the standard deviations to be below $0.1 \text{ eV/formula unit}$. Permittivity was taken as $\varepsilon = 1$, and the “true” energy levels can be obtained by dividing the results by the permittivity of the material. The results for the uniform structure could be compared with those evaluated using the Madelung constants (see Ref. 29) and turn out to be quite accurate. Also the electrostatic energy of the “$P\bar{3}m1$” structure—calculated with the Madelung method by assuming that the site geometry remains cubic—is found in Ref. 30 and compares well with our results. Furthermore we are not interested in absolute values but in comparative ones. For each structure we evaluate $\Delta E_{\text{structure}} = E_{\text{structure}} - E_{\text{BaTiO}_3}$. The results show that with respect to the uniform structure, in order of increasing stability,

1. the random structure has the same energy within 0.1%,
2. the QLRCO structure has 1.1% lower energy or about $\Delta E_{\text{QLRCO}} = -1.902 \text{ eV/formula unit}$;
3. the NaCl structure has 1.8% lower energy or about $\Delta E_{\text{NaCl}} = -3.229 \text{ eV/formula unit}$;
4. the $P\bar{3}m1$ structure has 3.0% lower energy or about $\Delta E_{\text{P\bar{3}m1}} = -5.295 \text{ eV/formula unit}$.

Noteworthy is the level spacing ratio $\zeta = \frac{\Delta E_{\text{QLRCO}} - \Delta E_{\text{NaCl}}}{\Delta E_{\text{NaCl}} - \Delta E_{\text{P\bar{3}m1}}} = 0.642 < 1$, which is smaller than $\zeta' = \frac{\Delta E_{\text{Random}} - \Delta E_{\text{NaCl}}}{\Delta E_{\text{NaCl}} - \Delta E_{\text{P\bar{3}m1}}} = 1.56 > 1$.

The importance of the level spacing ratios is that combined with the values of the entropy they regulate the important features of the phase diagrams. In particular, as is shown in Sec. V, they determine whether the free energy curves $U_{\text{ph}} = S_{\text{ph}} T$ of the various phases (subscript ph) intersect or not in a certain $T$ range.

There are several factors which will influence the absolute values of these energy levels, and the relative values of $\zeta$ and $\zeta'$ as well. These are as follows:
(1) The energies have to be divided by the material’s relative (static) permittivity \( \varepsilon \), which can be large in perovskites. For \( \text{BaMg}_{1/3}\text{Ta}_{2/3}\text{O}_3 \) it is found that \( \varepsilon \approx 24 \).\(^{31}\) This factor would not change the ratio.

(2) The electrostatic energy is evaluated using the full formal charges of the ions. Covalency corrections may play a role. A value of \( \eta_1 = 0.913 \) can be extrapolated from the valence sum corrections evaluated for BMT in Ref. 3. This entails an absolute change of the energy levels, not affecting the ratio \( \xi \).

(3) The electrostatic energy is inversely proportional to the cubic lattice parameter \( a \). Appropriate scaling is needed. However, for the known \( \text{BaMg}_{1/3}\text{Ta}_{2/3}\text{O}_3 \) phases, \( P\bar{3}m1 \) and QLRCO, \( a \) (evaluated from the cubic main reflections) is the same within one part in \( 10^3 \).\(^{3} \) This effect cannot change the ratios.

(4) Geometric distortions are present in the ordered NaCl and \( P\bar{3}m1 \) structures, but they affect the distances by only a few parts in \( 10^5 \),\(^{3} \) and as their effect on the energies is of comparable order, we can neglect them altogether.

(5) A PSF is observed, leading to smoothed cube edges. This term is possibly “accidental,” depending on the detailed crystallization kinetics, so it will not be considered quantitatively in the following. Its effect is also small enough (see Fig. 5) that it cannot qualitatively affect our results.

B. Entropy

The crystal structure can be stabilized by the entropy. For disordered structures configurational entropy plays an important role. A mixed-occupancy atomic site with occupancy \( x \) for one species and \( 1-x \) for another gives an entropy contribution of

\[
S = -k_B \left[ x \ln(x) + (1-x) \ln(1-x) \right].
\]

In the \( P\bar{3}m1 \) structure no mixed sites exist; therefore there have no entropic contribution to the Gibbs free energy. There will be an entropy term related to the domain distribution, but that is smaller by many orders of magnitude.

The maximal configurational entropy per formula unit is expected in the random structure and it amounts to

\[
S_0 = -k_B \left[ \frac{x}{2} \ln \left( \frac{x}{2} \right) + \frac{1-x}{2} \ln \left( \frac{1-x}{2} \right) \right]
= 54.85 \mu\text{eV/K}.
\]

For the QLRCO structure, as discussed, the entropy value is slightly smaller, but the difference can be evaluated numerically (in a simple but tedious way) from the correlation coefficients and the result is that the correction is negligible, so the entropy is \( \approx S_0 \). For the NaCl structure, only 50% of the sites are mixed so the entropy is \( S_0/2 \).

One more important contribution is the phonon entropy. A different B-site degree of order can in principle affect the phonon entropy. However, this effect is much smaller than the configurational term. This is supported by the scarce literature; see, e.g., Ref. 32. The specific heat \( C_v \) for BMT has been measured;\(^{33}\) however the phase was not fully characterized at the time (being, however, either QLRCO or \( P\bar{3}m1 \)). We can only compare the entropy evaluated from the experimental specific heat with data for Pb containing relaxor perovskite (\( \text{PbMg}_{1/3}\text{Ta}_{2/3}\text{O}_3 \), PMT).\(^{33,34}\) The phonon entropy of PMT extrapolated to \( T \gg \Theta_D \) exceeds that of BMT by 59 \( \mu\text{eV/K} \).

This large difference can be explained either by the polar nanoregions which are only present in PMT\(^{33} \) or by the difference in the masses of Pb and Ba.

V. DISCUSSION AND CONCLUSIONS

Entropy is important at high temperature for stabilizing different structures. For instance, considering the various free energies as calculated in the previous section, we can construct a phase diagram of BMT (Fig. 6). We have taken, as previously stated, realistic values of \( \varepsilon = 24 \) (assumed constant with \( T \)), \( \eta_1 = 0.913 \), and a relative linear expansion coefficient of \( 9 \times 10^{-6} \). We have not included any contribution from imperfect ordering (namely, the effect of a CDF limiting the spatial extent of correlations) because it is not easy to evaluate quantitatively and it is not an intrinsic effect but rather a kinetic effect while its contribution is small (Fig. 5) and the leading approximation would only compress the energy range.

The entropy term is canceled by the energy difference between QLRCO and \( P\bar{3}m1 \) phases at about 1600 °C, that is about the melting point. Above this temperature the QLRCO state would be stable while the \( P\bar{3}m1 \) state would be stable below. We can also see from Fig. 6 that there is no region that is stable for the NaCl-ordered phase, and in fact, this phase has never been observed in BMT, at variance with PMT.\(^{4,5}\)

Due to the slow kinetics associated with cation ordering, chemical ordering is usually achieved only after annealing for a long time at very high \( T \), where the equilibrium state may be dictated by the entropy. Cooling—due to the slow kinetics—cannot change the high-\( T \) equilibrium state so that it would remain preserved (metastable) at room temperature.

The random structure is always less stable than the QLRCO structure, because of the tiny entropy difference and the large energy difference that favors the latter structure. Therefore the random structure is a disordered metastable state. The ground state—or the thermodynamic equilibrium state at 0 K—is the \( P\bar{3}m1 \) structure. The true high-temperature thermodynamic equilibrium state is QLRCO, because the electrostatic energy difference with the \( P\bar{3}m1 \) structure is overcome by the entropy term at high \( T \), even if the dielectric constant is of moderate size. The NaCl structure is intermediate between these two. Given that its Gibbs free energy decreases with increasing temperature half as fast as the free energy of the QLRCO state, the ratio \( \xi \) becomes crucial to determine whether the NaCl

![FIG. 6.](Color online) Schematic phase diagram of BMT.
order can be the thermodynamic equilibrium state in some temperature range. In fact for $\xi < 1$ the Gibbs free energy of QLRCO is always lower than that of the NaCl structure, while for $\xi > 1$ there exists a temperature interval where NaCl structure is the equilibrium state. The QLRCO structure is favored above that temperature and the $P3\overline{3}m1$ structure below.

The arguments above are valid in principle for all $AB_1B_2$ perovskites with $A$ and $B$. $B''$ having strongly ionic character and charges $+2$, $+2$, $+5$, respectively, and with no other strong interaction. One most striking point is that the NaCl-type ordering could never appear in such cases. Therefore, a brief word is needed to explain why in PMT this is not the case, as mentioned above. We discuss here the results from Refs. 4, 5, 21, and 34 and literature therein. The difference between the two materials is the replacement of Ba$^{2+}$ with Pb$^{2+}$ in the $A$ site. The latter ion has much less ionic character and larger anisotropy in PMT, such that at sufficiently low temperatures the Pb atom never resides in the $A$ site but is always consistently displaced from it. Moreover, in a broad temperature range the Pb displacement vectors tend to co-align along the $(1,1,1)$, forming cigar-shaped polar nanoregions, while at higher temperature $>700$ K the Pb displacements are uncorrelated. This has large consequences both on the internal energy and on the entropy (as mentioned above). Therefore the phase diagram presented here has no reason to be valid to PMT, as too many important factors are missing. In addition, an important piece of information on the correlation between the partly ordered $A$ and $B$ sites is still largely unknown. In order to extend the discussion to PMT, such factors should be considered as well.

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APPENDIX

1. More choices of point-spread functions

A particular type of PSF/CDF pair has been shown in Eqs. (7) and (8). In this section some different PSF-CDF pairs will be discussed. PSFs are normalized so that their sum is 1. First, as in Eqs. (7) and (8), we consider the symmetry of the PSF to be cubic. A Lorentzian PSF and the corresponding exponential CDF are presented in Eqs. (7) and (8). A simpler PSF would be a sinc function, corresponding to a box function CDF.

The negative parts of the sinc tails are, however, a problem. Therefore we consider a sinc-squared function. Recalling $\text{sinc}(x) = \sin(x)/x$, $B_n(x) = 1$ if $|x| < w$, $B_n(x) = 0$ otherwise [see Eq. (1)] we have

$$
\text{PSF: } D(q) = (\pi k_0)^{-3} \sum_{a=1}^{3} \text{sinc}^2(q_a/k_0);
$$

$$
\text{CDF: } \tilde{D}(r) = \prod_{a=1}^{3} (1 - \pi |r_a/k_0| B_{\text{sinc}(k_0)}^{-1}(r_a)).
$$

Correspondingly, if the symmetry is spherical, the PSF/CDF are given by the radial Lorentzian or exponential:

$$
\text{PSF: } D(q) = \left(2\pi^2 k_0^2\right)^{-1} \left[1 + (q/k_0)^2\right]^{-2};
$$

$$
\text{CDF: } \tilde{D}(r) = e^{-2\pi r k_0},
$$

and the analog of the squared sinc/box pair:

$$
\text{PSF: } D(q) = 3\left(2\pi^2 k_0^3\right)^{-1} \left[\sin(q/k_0) - (q/k_0) \cos(q/k_0)\right]^2 / (q/k_0)^3;
$$

$$
\text{CDF: } \tilde{D}(r) = (1 + \pi r k_0/2)(1 - \pi r k_0)^2 B_{\text{sinc}(k_0)}^{-1}(r).
$$

2. Correlation coefficients for ordered and semiordered BMT phases

For the case in which the $B$ site is fully unordered, i.e., when all $c_L$ are randomly assigned, $S_{(000)} = 2$, but

$$
S_M = -1 + 9(c^2) = 0 \quad \text{whenever } M \neq (0,0,0),
$$

so the Fourier transform $I(q)$ is a constant, $I(q) = 2\eta^2$.

The trigonal superstructure has as the 3-fold axis one of the equivalent $(1,1,1)$ cubic axes. This means that it exists in four different orientations, where the 3-fold axis $T$ is $T = (1,1,1)$, $T = (-1,-1,-1)$, $T = (-1,1,-1)$, and $T = (-1,-1,1)$, respectively. The $B$-site superstructure is obtained by positioning in a lattice node $M$ on a Mg atom if $M \cdot T = 0$ modulo 3. Therefore, we will have pure Mg planes alternating with pairs of Ta planes, stacked orthogonally to $T$. In any crystallite, domains of the four different orientations will be present in equal volumes, therefore conserving the global cubic symmetry, in agreement with Landau’s theory. The correlations can be calculated for each orientation and then averaged. However, one can also calculate them from the scattering pattern, as an average over domains. The superstructure satellites—considering equipopulated domains—are eight equal Bragg peaks (each with intensity $I_{P_{3m1}}(m)$ at $\pm 1/3, \pm 1/3, \pm 1/3$).

$$
S_M = (-1)^{k+1} \frac{\eta^2}{9},
$$

where $k$ is the number of components of $M$ that are equal to zero, or $k = \#\{m_a = 0\}$ modulo 3, $\alpha = (1,2,3)$. From the condition $S_{(000)} = 2$, we obtain $I_{P_{3m1}} = \eta^4/4$, so that

$$
S_M = (-1)^{k+1}\frac{\eta^2}{9}.
$$

The hypothetical NaCl-type superstructure is simpler. There is only one Bragg peak at $(1/2,1/2,1/2)$ with intensity $I_{\text{NaCl}}$ plus a small constant background $I_b$ due to residual randomness on half of the sites. So one obtains

$$
S_M = (-1)^m I_{\text{NaCl}} + I_b, \quad S_{(000)} = \frac{I_{\text{NaCl}} + I_b}{\eta^2}.
$$

As half the sites contribute to the background and the occupancy of the random sites is always divided as $1/2$, $I_b = \eta^2$ and again using the condition $S_{(000)} = 2$, we have

$$
I_{\text{NaCl}} = \eta^2.
$$
3. Long-range order parameter

In this section we show how the former results can be related to the formalism of the long-range order parameter. A simple way of describing systems where there is a fully ordered state is by the long-range order parameter. Here the cubic $B$-site lattice is divided into 3 sublattices, denoted (a), (b), (c) for convenience of which one is occupied by Mg and two by Ta. There are four ways of choosing the threefold axis of the trigonal structure and for each three permutations of the Mg sublattice between (a), (b), (c). Suppose that we chose one axis and that the Mg sublattice is (a). In the ordered state all Mg atoms are actually on the (a) sublattice; if we chose one axis and that the Mg sublattice is (a). In the ordered state, 0 in the random one, and assumes intermediate values for all intermediate states. Using the same method lead to the expressions for the nearest-neighbor pair frequencies as a function of $X$. If the interaction is fundamentally short-ranged, or if the interactions beyond the first neighbors are not very important, one can then use different interaction hypotheses and calculate the internal energy. Finally, from the stochastic occupancy of each site, it is possible to evaluate the configurational entropy. The nearest-neighbor bond fractions turn out to be

$$v_1 = \frac{1}{N}(1 - X^2) \quad \text{Mg-Mg pairs},$$
$$v_2 = \frac{1}{N}(4 - X^2) \quad \text{Ta-Ta pairs},$$
$$v_3 = \frac{1}{N}(2 + X^2) \quad \text{Ta-Mg pairs}.$$

As nearest-neighbor bond vectors are just the (1,0,0) family, it is easy to see that

$$v_1 = \frac{1}{N} \sum L c_L c_{L'},$$

where $X$ is a parameter that is 1 in the ordered hypothetical NaCl-type phase. Obviously $X = 0$ for the random $B$-site phase, while for the QLRCO phase, using the $t \approx 0.1$ value estimated from the Monte Carlo runs, we have $X = 0.407$, which is well above the value of zero for the completely random structure. This, combined with similar results obtained for the short-range order parameter of Ref. 26 in Sec. III G2, is a further justification for the name "quasi-long-range order."

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This is similar in reverse to the case of a perfect crystal, where perfect periodicity in direct space means that the intensity is concentrated in the Bragg peaks, which are a weighted set of Dirac deltas located on the reciprocal lattice. The restriction of the periodic pair correlation function to one unit cell can be then evaluated as a discrete Fourier transform of the Bragg intensities and it is named the “Patterson function.”

For the $k$th B-B coordination shell in the $AB'_x\text{B}''_{1-x}O_3$ perovskites, the short-range order parameter is defined as (Ref. 28) $\alpha(k) = 1 - P_{B'}(k)/x$, where $P_{B'}(k)$ is the fraction of $B'$ atoms in that shell. The positive values indicate a preference for like neighbors, while the negative values, as obtained for BMT, for unlike ones.