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Origin of Spinel Nanocheckerboards via First Principles

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Self-organizing nanocheckerboards have been experimentally fabricated in Mn-based spinels, but have not yet been explained with first principles. Using density-functional-theory, we explain the phase diagram of the $\text{ZnMn}_x\text{Ga}_{2-x}\text{O}_4$ system and the origin of nanocheckerboards. We predict total phase separation at zero temperature, then show the combination of kinetics, thermodynamics, and Jahn-Teller physics that generates the system’s observed behavior. We find the (011) surfaces are strongly-preferred energetically, which mandates checkerboard ordering by purely geometrical considerations.

Experimental observation demonstrates intriguing nanoscale compositional ordering in a variety of material alloys. These include noble-metal-alloy nanocheckerboards [1–3], BaTiO$_3$-CoFe$_2$O$_4$ nanopillars [4], and an assortment of manganite-spinel nanocheckerboards [5–9]. Nanoscale phenomena are inherently difficult to treat with quantum mechanics’ first principles, due to the prohibitive scaling of electronic-structure methods. Previous theoretical studies [10–14] used phase-field models [15, 16] to simulate nanocheckerboard formation. However, those models rely upon coefficients chosen without first-principles justification. In contrast, our work reveals the origin of nanocheckerboards from first-principles.

Here we examine the experimentally well-characterized manganite spinels $A^{2+}\text{Mn}^{3+}_x\text{O}_4^\cdot$. These cooperative-Jahn-Teller crystals, upon doping with certain transition metals, organize into nanocheckerboards. Experiments showed that high-temperature mixing, followed by slow cooling, yields a spontaneously-formed checkerboard whose squares alternate tetragonal Mn-rich and cubic Mn-poor phases. These checkerboards emerge from the cross-section of spontaneously-aligned nanorods. Yeo et al. [5] fabricated self-assembling nanocheckerboards from $\text{ZnGa}_2\text{O}_4$ (ZGO) + $\text{ZnMn}_2\text{O}_4$ (ZMO), comprised of ~ $4 \times 4 \times 70$ nm$^3$ nanorods. Later work grew checkerboards with nanorods over 700 nm long on a MgO substrate [6, 7]. Checkerboards were later extended to other Mn-based spinels, first $\text{MgMn}_x\text{Fe}_{2-x}\text{O}_4$ (MMFO) [8] and then the tunable-sized checkerboards of $\text{Co}_{0.6}\text{Mn}_x\text{Fe}_{2.4-x}\text{O}_4$ (CMFO) [9]. The latter are notable for a patterning that alternates ferro- and paramagnetic phases, which yields potential for ultrahigh-density information storage.

We analyze the nanocheckerboard system $\text{ZnMn}_x\text{Ga}_{2-x}\text{O}_4$, whose relative simplicity renders it a minimal prototype. We sketch the main experimental results of Ref. 5 in Figure 1. ZGO is a cubic spinel while ZMO is a tetragonal spinel ($c/a = 1.14$ [17]), which immediately leads to some anomalous differences between experiment and naive expectations. First, room-temperature experiments reveal solid solutions at non-negligible concentrations, despite the disparate crystal structures of the end-members, in violation of the Hume-Rothery rules. Second, x-ray diffraction experiments show a cubic structure up to 25% ZMO (and 75% ZGO), whereas a non-negligible tetragonality would be expected at this concentration, regardless of the origin of the observed solubility. Third, in the region where ZMGO phase-separates, experiment shows checkerboard formations instead of traditional spinodal decomposition.

To address these issues, we compute the energetics of $\text{ZnMn}_x\text{Ga}_{2-x}\text{O}_4$ using density functional theory as implemented in the Vienna Ab-initio Simulation Package (VASP) [18–21]. We use a plane-wave cutoff energy of 415 eV, the PW91 generalized gradient approximation (GGA) functional [22, 23], and projector augmented wave (PAW) based pseudopotentials [24]. All calculations are initialized with ferromagnetic ordering for simplicity, a well-justified approach for the Néel temperature of merely $\approx 20$ K [17, 25, 26]. We compute a phase diagram via a cluster expansion [27] of the DFT ener-
getics with Monte-Carlo simulations, as implemented in the Alloy Theoretic Automated Toolkit (ATAT) package [28–31].

Both ZGO and ZMO crystallize in the spinel structure, with nominal valences of \( \text{Zn}^{2+}(\text{Ga}, \text{Mn})^{3+}\text{O}_4^{-} \). \( \text{Zn} \) occupies the tetrahedral ("A") sites and \( \text{Ga} / \text{Mn} \) occupy the octahedral ("B") sites, with negligible inversion [32, 33]. Therefore, our study focuses on the effect of \( \text{Ga} / \text{Mn} \) occupation of the B sites. Each B-centered octahedron shares edges with six others, which couples their anionic distortions. ZGO forms a cubic spinel (space group \( Fd\bar{3}m \)) [33], while ZMO is a tetragonal spinel (space group \( I4_1/amd \)) with significant distortion \( c/a = 1.14 \) [17]. The \( T' \) of Figure 1 (high-temperature fully-mixed ZMGO) has \( c/a \approx 1.06 \) [5]. We have provided lattice constants from the literature, and their DFT-calculated analogs, in Supplementary Material (Table S1) [34]; DFT calculations agree with experiment. ZMO’s sizable tetragonal distortion is due to the Jahn-Teller (JT) effect in the \( \text{Mn}^{3+} \) ions, where the \( d^4 \) configuration in a high-spin octahedral environment causes the \( e_g \) orbitals to break symmetry by a tetragonal distortion. This orbital ordering leads to a martensitic cubic-to-tetragonal transition in a variety of crystals, including spinels [35–39]. For consistency, we take [001] to be the JT-distorted direction.

Experiments, summarized in Figure 1, show solid solutions for \( x_{\text{Mn}} \leq 0.25 \) and \( x_{\text{Mn}} \geq 0.85 \). Yet the cubic and strongly-tetragonal crystal structures of the end members are expected to be immiscible: Placing a non-JT octahedron in a tetragonal JT-active environment, and vice versa, costs energy. Our DFT calculations quantitatively verify the qualitative Hume-Rothery rules: We generated 192 supercells (\( \leq 42 \) atoms) then fully relaxed the structures. (See Supplementary Material [34] for further calculation details.) Figure 2(a) shows their formation energies, demonstrating that mixing the disparate crystal structures incurs an energy cost [40]. A ground-state search using the cluster expansion (via the aforementioned ATAT package) revealed no lower-energy supercells. Therefore, the zero-temperature ground state of ZMGO is actually phase separation into bulk ZGO and ZMO, a conclusion absent from previous non-first-principles analyses [11–14]. There is no chemical or physical reason to believe that ZGO and ZMO should mix at anything but elevated temperatures.

To further verify this, we computed a phase diagram via a cluster expansion (CE), using the aforementioned ATAT package. (See Supplementary Material [34] for extensive calculation details, as well as the effect of CE shortcomings.) As shown in Figure 2(b), the zero-temperature stable phases are immiscible bulk ZGO and ZMO. In fact, the observed miscibility at \( x_{\text{Mn}} \leq 0.25 \) and \( x_{\text{Mn}} \geq 0.85 \) is unreasonable at all but extreme temperatures. We therefore attribute the anomalous miscibility to kinetic limitations, i.e. diffusion constraints. This is corroborated by the experimental observation that slower cooling (i.e. better diffusion) leads to larger checkers (i.e. less miscibility) [8, 9]. In fact, kinetics are the obvious origin of the miscible \( T' \) state observed upon rapid quenching (shown in Figure 1). DFT’s prediction for the \( T' \) state for \( x_{\text{Mn}} = 0.5 \) (taken as the mean of all calculated structures) agrees well with the experimental measurements. For example, DFT predicts \( c/a = 1.07 \), comparable with experiment’s \( c/a \approx 1.06 \). (See Supplementary Material [34] for all data.) Similarly, kinetic limitations must cause the apparent solubility of \( x_{\text{Mn}} \leq 0.25 \) and \( x_{\text{Mn}} \geq 0.85 \). For example, if diffusion essentially freezes by e.g. 900 K, the system cannot separate into 100/0% Mn mixtures and will remain a high-entropy frozen solid solution. Unfortunately, quantitatively predicting kinetics (including checker size) requires detailed knowledge of the diffusion mechanisms, coupled with complex JT lattice dynamics, which is beyond the scope of this Letter.

Having explained that the apparent solubility is due to kinetics, we must address the second anomaly, the experimental observation of a cubic structure for \( x_{\text{Mn}} \leq 0.25 \). In contrast, DFT predicts a tetragonal structure (\( c/a \approx 1.03 \) for all calculated structures at \( x_{\text{Mn}} = 0.25 \)) due to the cooperative Jahn-Teller effect.

We propose that the cubic structure of the Mn-poor phase is caused by noncooperative JT distortions at finite temperatures. It is well-known that, as in many spinels,
ZMO undergoes a phase transition to a cubic spinel above approximately 1323 K, due to noncooperative JT distortions (each octahedron distorting in a random direction) [32, 35, 36]. The transition temperature scales with doping: Within mean-field theory it is approximately linear with doping [41]; experimentally, a variety of spinels transition at $T_c \approx 9260(c/a - 1) \text{ K}$ [36], where $c/a$ is the tetragonal distortion induced by the cooperative distortion [42]. As illustrated in Figure 3, both these trends imply that at room temperature, structures of $x \lesssim 0.25$ are cubic due to this transition, while $x \gtrsim 0.25$ are tetragonal. Similar “critical concentrations” of JT-ions have been documented in the literature [41, 43]. This is also consistent with the conclusion of Noh et al. [44] that although $x_{\text{Mn}} = 0.25$ has an XRD pattern of a cubic spinel, it has a rather large JT splitting of $\sim 0.7 \text{ eV}$, i.e. a high-spin electronic configuration, which leads to JT distortion. (See Supplementary Material [34] for detailed analysis of the reliability of GGA’s high-spin prediction.) Therefore, low-temperature measurements should reveal DFT’s tetragonally-distorted structure for $x \leq 0.25$.

![Figure 3](image-url)

**FIG. 3.** Phase diagram for the noncooperative Jahn-Teller transition in spinels. DFT calculates the distortion (left axis) of our ZMGO structures (as in Figure 2(a)) at 0 K. The critical temperature for the tetragonal-to-cubic transition (right axis) is obtained based on the fitting $T_c \approx 9260(c/a - 1) \text{ K}$, empirically accurate for a variety of spinels [36]. Therefore at room temperature, $x_{\text{Mn}} \lesssim 0.25$ is cubic.

Therefore, the cubic structure is due to the high-entropy noncooperative distortion. We should note that these finite-temperature Jahn-Teller lattice dynamics are expected to enhance miscibility, as noted by other experiments [45]. For example, whereas experiment places the boundaries at 25/85% Mn, our phase diagram (Figure 2(b)) shows that ZMO is more tolerant of Ga-doping than ZGO of Mn-doping. This discrepancy is likely due to the lack of noncooperative distortions in the cluster-expansion model, although a quantitative treatment is beyond the scope of this Letter. However, this alone would not suffice to cause appreciable solubility near room temperature.

Now we turn to $0.25 < x_{\text{Mn}} < 0.85$, where slowly-cooled samples phase-separate on a diffusion-limited scale. We seek to explain how this leads to nanoflakes rather than traditional spinodal decomposition. We calculate the energy of joining a slab of ZGO to a slab of ZMO along a particular surface; lower energies indicate stable interface directions [46]. Figure 4(a) shows the energies of formation for various slab thicknesses. (Larger thicknesses are inaccessible due to large, convergence-challenged supercells.) In agreement with experiment, DFT prefers the (011) surfaces. The energy relative to the next-preferred surface is about 10 meV/B (160 meV per cubic unit cell) at only 1.5 nm, and presumably larger for the 4 nm checkerboard.

![Figure 4](image-url)

**FIG. 4.** (a) Formation energies of ZGO/ZMO slabs layered in the given direction, normalized per B-site, plotted against the average thickness $t$ of the ZGO and ZMO slabs. The $t \to \infty$ limit corresponds to the contribution of B-sites far from the interface; finitely valued $t$ averages the contribution of all sites, including those near the interface. See also Figure 5(b-c). (b) Same, multiplied by the number of B-sites ($N$) and divided by the interface area ($A$), giving a per-area normalization. (c) Fitting cubic (C) and tetragonal (T) domains with a coherent interface along (011) surfaces. Arrows show the tetragonally elongated direction in corresponding domains. Geometrically, coherent (011) C-T interfaces require a slight angular rotation (shown) between adjacent C and T regions. The consequent zigzag interface can match either an extended equivalent domain (illustrated on left of subfigure) or a commensurate domain (on right), forming a checkerboard structure. This generates tetragonal domains rotated by 90° and cubic domains rotated by $\theta$ (see text).

Due to symmetry, only five directions are calculated directly. We search formation energies of multilayer slabs of thickness $t$ oriented in an arbitrary direction $\mathbf{k}$ by performing an expansion in the symmetry-adapted (tetragonal) harmonics, a standard group-theory methodology.
It is apparent from Figure 4(b) that the total formation energy can be approximated by \( E(\mathbf{k}, t) = A(c_a(\mathbf{k}) + t c_v(\mathbf{k})) \), where \( A \) is the cross-sectional area and \( c_a, c_v \) are constants. We perform an expansion of \( c_a, c_v \) in the four lowest-order harmonics via a least-squares fit for the five calculated values of \( \mathbf{k} \). (Fitting data appear in Supplementary Material [34].) We thus confirm that \( \{011\} \) and equivalently \( \{0\overline{1}1\} \), are the lowest-energy surfaces.

Hence physics dictates that when our system coherently mixes cubic and tetragonal phases, it forms \( \{011\} \) and \( \{0\overline{1}1\} \) interfaces. Now pure geometry dictates checkerboard configurations for coherent interfaces. As shown in Figure 4(c), due to bent angles at the interfaces, the only configuration that retains a coherent lattice is the checkerboard formation. Experiments observed that alternating cubic phases are rotated by a few degrees, while alternating tetragonal phases are rotated by 90° [5, 6, 9]. The origin of these measurements appears clearly in Figure 4(c), with the cubic-phase angle of rotation \( \theta = \pi/2 - 2 \tan^{-1} a/c \). This agrees with measured values within \( < 1° \) (see data in Supplementary Material, Table S2 [34]). In this, our checkerboards are directly related to the CoPt cubic-tetragonal nanostructures [1, 10] and other lattice-induced interface rotations [49].

Whence does the \( \{011\} \) preference originate? Previous work showed the importance of both strain and local ionic distortions (i.e., \( \mathbf{k} = 0 \) phonons) to the Jahn-Teller effect [50]. Our slabs’ formation energy consists of strain energy, due to biaxial lattice-matching, and contact energy, due to local ionic distortions and local binding energies. Specifically, for two slabs of thickness \( t \) joined in direction \( \mathbf{k} \), we write:

\[
E_{\text{formation}} = NE_{\text{strain}}(\mathbf{k}, t) + AE_{\text{contact}}(\mathbf{k}, t) \tag{1}
\]

where the units now ensure that the latter two \( E \) terms converge for for \( t \to \infty \). As shown in Figure 5(a), the strain energy can be computed directly by relaxing ionic coordinates for ZGO and ZMO separately, with strained lattice vectors (the “strained bulk” calculation). Then the contact energy is simply the difference between the formation energy and the strain energy. (Detailed equations appear in Supplementary Material [34].)

These energies are shown in Figure 5(b-c). We note that \( \{111\} \) and especially \( \{011\} \) have negative contact energies, meaning the layered-slab heterostructure is more stable than the strained ZGO and ZMO separated bulk. Hence the local ionic distortions couple beneficially to the lattice strain, i.e. atomic rearrangements partially alleviate the energy penalties of lattice strain. This is predominantly concentrated in the breathing and tetragonal-distortion local modes \( q_2 \) and \( q_4 \) in the notation of Ref. 51). Remarkably, the contact energy does not converge in the accessible thicknesses, due to the long range of the Jahn-Teller effect. Rather, even a remarkable distance from the surface, interoctahedron coupling leads to intracellular atomic displacement. Therefore the \( \{011\} \) preference originates in a beneficial coupling between strain and long-range atomic displacements (contrary to previous non-first-principles work neglecting the latter [10–14]).

In conclusion, we have presented the physics of nanocheckerboards based on first-principles calculations. We established that the thermodynamic ground state is complete phase separation. The incomplete separation originates in diffusion limitations, leading to nanoscale phase domains. We explained the observed cubic crystall structure at \( x_{\text{Mn}} = 0.25 \) based on noncooperative Jahn-Teller distortions at room temperature. Therefore, although ZMO’s ground state is bulk-incoherent, the diffusion-limited structure is bulk-coherent (using the terminology of Ref. 52). This kinetically-driven bulk coherence leads to phase separation of cubic and tetragonal phases along \( \{011\} \) surfaces, which we showed from first principles. This, in the presence of kinetic constraints, automatically leads to checkerboards, based on pure geometry. The preference for \( \{011\} \) surfaces originates in beneficial coupling between local distortions and lattice strain. Further quantitative understanding will require robust models for the Jahn-Teller effect in doped mate-
materials at finite temperatures.

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