Atomic short-range order and incipient long-range order in high-entropy alloys

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Atomic short-range order and incipient long-range order in high-entropy alloys

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Within density-functional theory, we apply an electronic-structure-based thermodynamic theory to calculate short-ranged order (SRO) in homogeneously disordered substitutional N-component alloys, and its electronic origin. Using the geometric properties of an \((N−1)\)-simplex that describes the Gibbs (compositional) space, we derive the analytic transform of the SRO eigenvectors that provides a unique description of high-temperature SRO in N-component alloys and the incipient low-temperature long-range order. We apply the electronic-based thermodynamic theory and the new general analysis to ternaries (A1 Cu-Ni-Zn and A2 Nb-Al-Ti) for validation, and then to quinary Al-Co-Cr-Fe-Ni high-entropy alloys for predictive assessment.

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I. INTRODUCTION

Multicomponent metallic alloys constitute an important, widely used class of technological materials. Properties of N-component alloys are sensitive to the state of chemical order in a stable lattice structure. Recently, so-called high-entropy alloys (HEAs) have drawn much attention due to their remarkable properties. HEAs consist of five or more \((N≥5)\) elements with (nearly) equal composition of atomic species. If not too large, their alloy formation enthalpy \(ΔE_0\), dictating ordering \((ΔE_0<0)\) or phase segregation \((ΔE_0>0)\), are easily overcome by the large entropy associated with near-equitatomic compositions, stabilizing solid solutions and suppressing the formation of (small unit cell) intermetallic phases \([1,2]\).

While solid solutions lack chemical long-range order (LRO), they often possess atomic short-range order (SRO) that reveals the high-temperature, incipient chemical ordering tendency, either clustering or ordering. Often the SRO is indicative of the low-temperature LRO, albeit not guaranteed for first-order transitions. For a stoichiometric N-component alloy, in general, there must be \(N−1\) ordering transitions from the homogeneous phase. Hence, SRO can often be used to predict the expected LRO \([3,4]\).

Notably, SRO can be measured in diffuse-scattering (x-ray, electron, and neutron) experiments by extracting Warren-Cowley SRO parameters \([5–7]\), \(α_{μν}(k)\) at scattering wave vector \(k\), which are normalized pair (correlation) probabilities discussed later. For multicomponent alloys, diffuse scattering intensities are given in terms of Laue units \([3,4,8,9]\), i.e.,

\[
f_{μν}(fμ − fν)\left[f_μ − f_ν\right]^2 = \sum_{μ,ν} c_μ(δ_{μν} − c_ν)[f_μ − f_ν]^2 α_{μν}(k),
\]

with atomic scattering form factors \(f_μ\), component labels (μ, Greek lower-case letters), and compositions \(c_μ\). Only off-diagonal SRO parameters may be measured if there is contrast, i.e., \(f_μ − f_ν ≠ 0\). For the well-studied A-B binaries, a single \(α_{ab}(k)\) correlation can be observed, and diagonal values are obtained via sum rules dictated by the optical theorem. Interpretation is straightforward: if not an A atom, then a B atom in a favorable wave-vector ordering sequence. For N-component alloys, \(N(N−1)/2\) pairs must be measured, achieved only if there is sufficient scattering \(f_μ − f_ν\) contrast; but, without all the off-diagonal pairs, the SRO cannot be interpreted.

Here, we do not address the challenge of how one measures all off-diagonal pairs of N-component alloy. We provide a general theoretical method to predict all pair correlations and, in particular, how to interpret uniquely the SRO manifest in \(α_{μν}(k)\). The approach uses the Korringa-Kohn-Rostoker (KKR) Green’s-function electronic-structure method in combination with an inhomogeneous coherent-potential approximation (CPA) developed for binaries \([10,11]\) and extended to ternaries \([3,8,12]\). From the KKR-CPA, we may connect SRO to its electronic origin (competing effects of band filling, hybridization, Fermi-surface nesting, van Hove states, …), and confirm the behavior by direct calculation of formation energies \(ΔE_0\) for partially ordered states.

After some background, we relate in Sec. III the SRO to second-order (infinitesimal) concentration variations to the alloy free-energy (chemical stability matrix). Eigenvectors of the stability matrix reveal the incipient order reflected in the SRO. Describing the N-component system in Gibbs space as an \((N−1)\)-simplex using \(c_μ\) as natural barycentric coordinates, we establish the transform in Gibbs space that yields a correct analysis of SRO in N-component solid solutions. In particular, we obtained the transform’s matrix elements analytically for N-dimensions that guarantees the SRO eigenvectors are properly chemically and geometrically orthogonal. We apply the new SRO analysis to two well-studied ternaries (A1 Cu-Ni-Zn \([3,8]\) and A2 Nb-Al-Ti \([12]\)) to validate, and then to quinary Al-Co-Cr-Fe-Ni alloys. With this general analysis approach, our electronic-structure-based SRO theory can be analyzed using the SRO eigenvectors in a concentration-wave (Fourier) analysis \([3,8,13]\), now rigorously generalized to N-components alloys.

II. BACKGROUND

The study of SRO in disordered solid solutions has a long history, mostly on binaries and a few ternaries. In the...
multicomponent case, the complexity with experiments and interpretation of the pair correlations remains a challenge. In disordered solid-solution alloys, the SRO is a thermally induced infinitesimal concentration fluctuation and directly related to the chemical pair correlations [8,9,13–16]. In terms of site \((i\) and \(j\) occupation variables, pair correlations \(q_{ij}^{\mu
u}(k)\) are mathematically defined as

\[
q_{ij}^{\mu
u}(k) = \langle \delta^{\mu
u}_{k} (\epsilon_{\mu} - c_{\mu}) (\epsilon_{\nu} - c_{\nu}) \rangle = \langle \delta^{\mu
u}_{k} \hat{\epsilon}_{\mu}^{i} \hat{\epsilon}_{\nu}^{j} \rangle - \langle \delta^{\mu
u}_{k} \hat{\epsilon}_{\mu}^{i} \hat{\epsilon}_{\mu}^{j} \rangle, \tag{2}
\]

where \(\langle \ldots \rangle\) represents a thermodynamic average, and \(\hat{\epsilon}_{\mu}^{i}\) is a site-occupation variable; that is, \(\hat{\epsilon}_{\mu}^{i} = 1\) (0) if the site is (is not) occupied by an \(\mu\)-type atom at the site \(i\), hence, \(\langle \epsilon_{\mu}^{i} \rangle = c_{\mu}\).

For an \(N\)-component alloy in a “host” picture, there are \(N - 1\) independent fluctuations, as the host (dependent) variable is dictated by a site single-occupancy constraint, i.e., \(\sum_{\mu} \epsilon_{\mu}^{i} = 1\), forcing \(\sum_{\mu} c_{\mu} = 1\), so there are only \(N - 1\) independent occupations. Notably, a vacancy can be treated as just another independent species (not host). With \(\hat{\epsilon}_{\mu}^{i}\) idempotent \([\langle \hat{\epsilon}_{\mu}^{i} \rangle^{2} = \langle \hat{\epsilon}_{\mu}^{i} \rangle]\), site-diagonal pair correlations \(\langle \epsilon_{\mu}^{i} \epsilon_{\mu}^{j} \rangle\) obey the sum rule

\[
q_{ij}^{\mu
u}(k) = c_{\nu}^{i} (\delta_{\mu\nu} - c_{\nu}^{i}), \tag{3}
\]

where \(\delta_{\mu\nu}\) is Kronecker \(\delta\) function over species. For an \(A - B\) binary, \(c_{A} + c_{B} = 1\), and, if \(c_{A}\) is considered independent, then \(c_{B}\) is the dependent (host) variable, and \(q_{A\Lambda}^{\mu
u} = c_{A}(1 - c_{A})\).

We now can define Warren-Cowley (WC) parameters \([5–7]\) as normalized pair probabilities in Laue units, i.e.,

\[
a_{ij}^{\mu
u}(k) = \frac{q_{ij}^{\mu
u}(k)}{c_{\mu} (c_{\mu} - c_{\nu})}, \tag{4}
\]

and in real space this can be understood in terms of pair probabilities:

\[
p_{ij}^{\mu
u} = c_{\nu}^{i} c_{\nu}^{j} [1 - a_{ij}^{\mu
u}(k)], \tag{5}
\]

where \(p_{ij}\) is a point probability:

\[
\sum_{\nu} c_{\nu}^{i} a_{ij}^{\mu
u}(k) = 0 \tag{6}
\]

that allows us to get unmeasured (diagonal) correlations. As shown rigorously elsewhere \([9]\), the theory for SRO for disordered alloys used here calculates the nonsingular \((N - 1) \times (N - 1)\) portion of the inverse of

\[
[q^{-1}(k)]_{\mu\nu} = \left[ \frac{\delta_{\mu\nu}}{c_{\mu}} + \frac{1}{c_{\nu}} \right] - \beta S^{(2)}_{\mu\nu}(k), \tag{7}
\]

where \(\{\mu, \nu\} \in 1, N - 1\) and \(\beta = (k_{B} T)^{-1}\), defined by temperature \((T)\) and Boltzmann constant \((k_{B})\). The expression is exact \([9,17,18]\). Most notably, \(S^{(2)}(k)\)—the chemical fluctuation stability matrix—is a thermodynamic functional that reflects the free-energy cost of a pair fluctuation, and is represented by a symmetric matrix; importantly, it is not a pair interaction, as often assumed \([9]\). At the spinodal temperature \(T_{sp}\) for a specific maximum wave-vector instability, the inverse pair correlation first vanishes (i.e., the pair correlation diverges). Hence, from the stability matrix we can determine which wave-vector instability first becomes unstable, what pair(s) drives this instability, and at what temperature this instability occurs. All this information is useful for predicting and characterizing SRO in any complex solid solution, like any HEA.

From linear-response theory, \(S^{(2)}_{\mu\nu}(k)\) is the second variation of the alloy grand potential with respect to composition fluctuations, evaluated in the high-\(T\) disordered phase \([4,9]\).

That is, expanding the interacting electronic part of the free energy in a functional Taylor’s series relative to the disordered alloy, we find \([9,10,18–20]\) that

\[
\Delta F(\{c_{\mu}^{i}\}) = F(\{c_{\mu}^{i}\}) - F(\{\bar{c}_{\mu}^{i}\}) = \frac{1}{2} \sum_{\mu, \nu, ij} \Delta c_{\mu}^{i} \Delta c_{\nu}^{j} \sum_{k} \frac{\delta_{\mu\nu}^{ij}}{\delta_{\mu\nu}^{ij} \delta_{\nu\mu}^{ij}} |_{c_{\nu}} |_{\Delta c_{\mu}^{i}} |_{\Delta c_{\nu}^{j}} \approx \frac{1}{2} \sum_{\mu, \nu} \int_{BZ} d\mathbf{k} \Delta c_{\mu}^{\mu}(k) S^{(2)}_{\mu\nu}(k) \Delta c_{\nu}(k), \tag{8}
\]

and where the first variation is zero by symmetry of the disordered state. The Brillouin zone (BZ) integral is over that of the solid solution’s Bravais lattice, where the fluctuations \(c_{\mu}^{i} - \bar{c}_{\mu}^{i}\) [i.e., Fourier wave \(\Delta c_{\mu}^{i}(k)\)] happen. The total alloy free energy, \(F = F_{0} + F_{\text{int}}\), can be written as the sum of noninteracting, \(F_{0}\), and interacting, \(F_{\text{int}}\), contributions, where \(F_{0}\) is related to point entropy \((S_{\mu})\) [i.e., \(T S_{\mu} = -k_{B} T \sum c_{\mu}^{i} \ln c_{\mu}^{i}\)] and \(F_{\text{int}}\) comes from electronic structure (ion-ion, band-energy, double-counting, and exchange correlation). \(S^{(2)}_{\mu\nu}(k)\) is determined from \(F_{\text{int}}\) in the solid-solution phase, and encompasses all electronic effects (hybridization, band filling, Fermi-surface nesting, van Hove states, ...).

If evaluated in the homogeneous state, \(a_{\mu\nu}(k)\) is an approximation to the state with actual SRO, and it can be calculated with KKR-CPA linear-response codes \([3,4,12]\). Given calculations of \((6)\), the Warren-Cowley SRO parameters \([5–7]\) are determined from \((4)\). What remains to be determined is ordering fluctuations embodied in \(S^{(2)}_{\mu\nu}(k)\).

## III. CHEMICAL SRO EIGENVECTORS

In an \(N\)-component alloy with \(\sum_{\mu=1}^{N} c_{\mu}^{i} = 1\), the independent compositional fluctuations are described by \(N - 1\) component vectors. The associated Gibbs space is \(N\) dimensional and represented by \((N - 1)\) simplices: for \(N = 2\), a line (one-simplex), for \(N = 3\), a triangle (two-simplex), for \(N = 4\), a tetrahedron (three-simplex), and so on. From sum rules, \([c_{\mu}\] are natural barycentric coordinates as developed for finite-element methods \([21]\), and define coordinates in whole Gibbs space. Figure 1 shows a schematic for ternary and quaternary simplices and arrows labeled by \(\delta c_{\mu}\)’s are the directions of fluctuations in Gibbs space (parallel to directions of increasing concentrations); generally, these axes are oblique to one another—not geometrically orthogonal \([3,9,22,23]\).

The fluctuation energy (in matrix-vector notation) is

\[
\Delta F = \hat{\delta}^{T} S \hat{\delta} \tag{9}
\]

By solving the characteristic equation for \((8)\), the \(S_{\mu\nu}(k)\) eigenvectors \((\epsilon_{\nu})\) exhibit a host dependence, and possibly
unphysical (negative) concentrations. Formally, it is first necessary to transform from Gibbs (δ) to a Cartesian (\(\tilde{\delta}\)) coordinate system (\(\tilde{\delta} = T \delta\)), where \(\tilde{S} = T^\dagger ST\) in (9) has eigenvectors \((\tilde{e}_v)\) that are now host independent but represented in Cartesian space. Then, we must transform \(e_\gamma\) back to Gibbs space \((e_G = T e_v)\) to get the physically proper eigenmodes and concentrations (ordering probabilities). Because compositional representation of Gibbs space based on simplices is oblique, the transformation matrix \(T\) is not unitary, \(T^{-1} \neq T^\dagger\) and \(T T^\dagger \neq 1\), so eigenvectors \(e_\gamma\) are not same as \(e_G\), which are the correct ones in Gibbs space. Notably, with \(T\) known analytically for the \(N\)-dimensional case, we may analyze all SRO directly via \(e_G\).

A. Matrix elements of \(T\) analytically for \(N\)-component alloys

The form of \(T\) is well known in terms of barycentric coordinates is well known for any dimension, i.e., \(T^{-1} \neq T^\dagger\) and \(T T^\dagger \neq 1\), so eigenvectors \(e_\gamma\) are not same as \(e_G\), which are the correct ones in Gibbs space. Notably, with \(T\) known analytically for the \(N\)-dimensional case, we may analyze all SRO directly via \(e_G\).

The Jacobian transformation from Cartesian to oblique coordinates is well known for any dimension, i.e., \(T\) is

\[
N - 1 T = \begin{pmatrix}
X_1^N - X_1^1 & X_1^2 - X_1^N & \cdots & X_1^{N-1} - X_1^1 \\
X_2^N - X_2^1 & X_2^2 - X_2^N & \cdots & X_2^{N-1} - X_2^1 \\
\vdots & \vdots & \ddots & \vdots \\
X_N^{N-1} - X_N^1 & X_N^2 - X_N^N & \cdots & X_N^{N-1} - X_N^1
\end{pmatrix}
\]

where \(\{X_j^i\}\) represents coordinates of \(j\)th vertex relative to the host vertex \(\{X_1^N\}\). In a host picture, the \(N\)th vertex in \((N - 1)\) simplex is redundant from the sum rule for barycentric coordinates so the \(T\) matrix will always be rank \(N - 1\). Coordinates of all \((N - 1)\) vertices for Gibbs simplices are derived (see the Appendix) as

\[
X_j^i = \frac{1}{X_j^N} \sqrt{\frac{2(N-1)}{N}} \left[ X_{j-1}^i \cdot X_{j-1}^{N+1} + \frac{1}{N-1} \right]
\]

and each \(i, j\) runs from 1 to \(N - 1\).

From this analytic matrix element for \(T\) for any Gibbs space, we may evaluate easily the specific ternary and quinary cases addressed in the Results section. For a ternary (two-simplex), the \(2 \times 2 \times T\) matrix is

\[
2 T = \begin{pmatrix}
-1 & -1/\sqrt{3} \\
1 & -1/\sqrt{3}
\end{pmatrix}
\]

while, for quinary (four-simplex), the \(4 \times 4\) matrix is

\[
4 T = \begin{pmatrix}
-1 & -1/\sqrt{3} & -1/\sqrt{6} & -1/\sqrt{10} \\
1 & -1/\sqrt{3} & -1/\sqrt{6} & -1/\sqrt{10} \\
2/\sqrt{2} & -1/\sqrt{6} & -1/\sqrt{10} \\
0 & \sqrt{3}/2 & -1/\sqrt{10}
\end{pmatrix}
\]

Notice that \(3 T\) (ternary) is a submatrix of \(4 T\) (quinary), and so too is \(3 T\) the \(3 \times 3\) submatrix (quaternary).

B. Concentration waves in \(N\)-component alloys

Given \(e_\sigma\) (ordering normal modes) and \(T\) (transform matrix), we need to interpret the order reflected in the SRO, accomplished best by Fourier analysis. The occupational vector, \(\hat{n}(r)\), gives the probabilities of an atom to occupy specific sites in a crystal structure. In substitutional solid solutions, \(n_i(r)\) is identical to concentrations, \(e_i(r)\). But, in ordered phases, this depends on the type of order and real-space site coordinates [13]. The vector \(\hat{n}(r)\) for a \(N\)-component alloy, where all sites are represented by the same Bravais lattice, can be expanded in a Fourier series (concentration wave) and written in terms of normal modes as

\[
\begin{pmatrix}
n_1^1(r) \\
n_1^2(r) \\
\vdots \\
n_1^{N-1}(r)
\end{pmatrix} = \begin{pmatrix}
c_1^1(r) \\
c_1^2(r) \\
\vdots \\
c_1^{N-1}(r)
\end{pmatrix} + \sum_{\sigma, r} \hat{n}_{\sigma} e_{\sigma}(k_\sigma) e^{i k_\sigma \cdot r}
\]

For a given crystal lattice vector \(r\), \(c_{\sigma}^{N-1}(r)\) is composition of the \((N - 1)\)th component. The sums run over the star \(s\) (inequivalent wave vectors that define ordering), \(\sigma\) (eigenvector branch of the free-energy quadric), and \(j_\sigma\) (equivalent wave vectors in the \(s\)th star). The other quantities are LRO parameter \(n_{\sigma}^0\) for the \(\sigma\)th branch and \(s\) star; \(v_{\sigma}^{N-1}\) is \((N - 1)\)-component vector of the normal concentration mode \((e_\sigma)\) of \(S_{\sigma\nu}(k)\) for the \(\sigma\)th branch; and the symmetry coefficient \(\gamma_{\sigma}(k_\sigma)\) determined by normalized condition and geometry.

IV. COMPUTATIONAL DETAILS

We use a Korringa-Kohn-Rostoker (KKR) coherent potential approximation (CPA) code [24–26]. For metallic solid solutions, the screened CPA is used to incorporate Friedel screening from charge correlations in the local chemical environment [26]. As our KKR-CPA-based SRO theory is
only coded for the atomic sphere approximation (ASA), we use the KKR-CPA-ASA for all results, and include so-called muffin-tin (MT) corrections to the ASA total energies. We also evaluate electronic properties and total energies using Voronoi polyhedra (VP) integration [27] for spherically averaged radial functions in the site-centered, spherical-harmonic \((Y_L)\) basis. We include \(s,\ p,\ d,\) and \(f\) symmetries in the KKR basis, i.e., truncated at \(L_{\text{max}} = 3\), where \(L \equiv (l, m)\).

Potentials, charge densities, and total energies are obtained using a complex-energy Gauss–Legendre semicircular contour with 24 points, and Brillouin-zone integrations use a special \(k\)-point method [28] with a \(20 \times 20 \times 20\) mesh. We use the vonBarth–Hedin [29] local-density approximation as parametrized by Moruzzi, Janak, and Williams [30]. Self-consistency for potentials and charge densities is achieved with convergence technique based on modified Broyles’ second method [31]. Scalar-relativistic effects are included, but spin orbit is ignored. Because the potential zero \(v_0\), i.e., \(muffin-tin\) zero, can dramatically affect stability prediction for spherical potentials, we use a variational definition [32] that yields kinetic energies that approach those of full-potential methods [32,33].

The first-principles theory of SRO in multicomponent alloys has been presented before [10,11,20]. The complete expression for \(S_{\mu\nu}^{(2)}(q; T)\), one that includes all electronic-structure, charge screening and transfer, has been derived only for binaries [10,11,20]. The importance of metallic screening for SRO calculation in the solid solution phase has been discussed before for binaries [34]. However, at fixed composition, assuming site charges change little with SRO, Pettifor’s force theorem can be applied and \(S_{\mu\nu}^{(2)}(k; T)\) then has contributions only from the band-energy variations [3,8].

Hence, for present purpose to showcase the prediction and interpretation of SRO for \(N\)-component metallic solid solutions, we calculate all \(S_{\mu\nu}^{(2)}(q; T)\) results using the \(band\-energy\-only\) expression; that is, double counting terms and exchange correlation are neglected by invoking the force theorem. (We will add these variations in the future.) The scalar-relativistic KKK-CPA-ASA potentials, charge densities, and scattering matrices for a given solid solution are used to then evaluate the linear-response expression for \(S_{\mu\nu}^{(2)}(q; T)\). The expression for \(S_{\mu\nu}^{(2)}(q; T)\) are evaluated on a log mesh along the Matsubara poles, and interpolated to the correct poles (temperature) for use in response functions [10,11].

Thermodynamically, the \(Warren-Cowley\) parameters \((4)\) [5–7] must obey the optical theorem and conserve the particle number associated with the through beam, i.e., at the \(i\)th site (see the review [9])

\[
\alpha_{\mu\nu}^{ii} = 1 + \frac{\delta_{\mu\nu}}{c_{vi}} = \frac{1}{V_{\text{BZ}}} \int d\mathbf{k} \alpha_{\mu\nu}(\mathbf{k}).
\]

Any mean-field approximation to \(S^{(2)}(q; T)\) does not guarantee this sum rule [9], as is true of the CPA. As shown from (6), modifying \(S_{\mu\nu}^{(2)}(q; T) \equiv S_{\mu\nu}^{(2)MF}(q; T) - \Lambda_{\mu\nu}(T)\) satisfied the sum rule [8–10] with

\[
\Lambda_{\mu\nu}(T) = \frac{1}{V_{\text{BZ}}} \sum_{\beta} \int d\mathbf{k} S_{\mu\nu}^{(2)MF}(\mathbf{k}, T) \alpha_{\beta\nu}(\mathbf{k}).
\]

This coupled set of equations may be solved by Newton-Raphson techniques, using multidimensional mapping for inversion of tensors. \(S^{(2)}(q; T)\) is typically very weakly temperature dependent arising from the Fermi factors (see below), while \(\alpha_{\mu\nu}(\mathbf{k})\) strongly depends on temperature, diverging at \(T_{\text{sp}}\), see (6). The correction is historically called the Onsager cavity field correction [35], which renormalizes the thermodynamic excitation energies to conserve the diffuse intensity over the Brillouin zone. Although not commonly used as a more proper mean-field theory, this single-site fix to mean-field theory corrects the topological error in mean-field phase diagrams, such as Bragg-Williams (Ising) models [36].

Besides energy- and species-dependent matrix elements \(M_{\mu\nu}^{(2)}(\epsilon)\) and Fermi factors \(f(\epsilon)\), \(S^{(2)}(q; T)\) is found from the KKR scattering path operator, \(\tau_{LL}(k; \epsilon)\), which determines the Green’s function and embodies all electronic-structure effects [3,8,10,11]. In brief, \(S^{(2)}(q; T)\) is a generalized susceptibility, and, roughly [18], in terms of the Bloch spectral functions \(A(k; \epsilon) = -\text{Im} \epsilon(\mathbf{k}; \epsilon) / \pi\) (dispersion), we may suggestively write it as

\[
S_{\mu\nu}^{(2)}(q; T) \sim \int d\mathbf{E} M_{\mu\nu}(\mathbf{E}) \int d\mathbf{E}' f(\mathbf{E}; T) f(\mathbf{E}'; T) \int d\mathbf{k} A(k; \epsilon) A(k + q; \epsilon') \int d\mathbf{k} L_{\text{BZ}} \int d\mathbf{k} A(k; \epsilon) A(k + q; \epsilon').
\]

All valence states contribute to (17). If only hole and electron states near the Fermi energy, \(E_F\), dominate, the bracketed \([\ldots]\) term collapses the energy integrals to

\[
S^{(2)}(q; T) \rightarrow \int d\mathbf{k} A(k; E_F) A(k + q; E_F),
\]

a convolution integral of the Fermi-surface states and the origin for so-called Fermi-surface “nesting” [3,8,18]. Due to alloying, even in a metallic system, hybridized states well below \(E_F\) in (17) can drive ordering, as for NiPt [37], or van Hove features at \(E_F\) contribute, as for CuPt [38]. In short, the nature and electronic origin of the SRO may be determined directly.

For completeness, using a laptop computer with an Intel i7 (2.3 GHz quad core) processor, we provide some timings involved in evaluating the self-consistent potentials and linear-response \(S^{(2)}(q; E)\) for 24 energy points on the semicircular or Matsubara contour. For pure bulk (one-atom) elemental ground states, KKR-CPA SCF calculations require 10–20 iterations with \(\sim 10\) sec per iteration. However, for bulk solid solutions (effectively one-atom cells for \(A1\) or \(A2\)), KKR-CPA SCF calculations require 30–50 iteration with \(\sim 25–50\) sec per iteration, due to cost of the CPA convergence that often takes longer than charge convergence. Typically, as the number of components (disorder) increases, the CPA is easier to converge and uses fewer iterations. For linear-response calculations in serial mode for a binary, the \(S^{(2)}(q; E)\) requires \(\sim 3600\) sec. Postprocessing analysis requires another \(\sim 20\) sec. For the \(N\)-component case, the time grows linearly with \(N - 1\) due to extra matrix elements in \(S^{(2)}\). For the magnetic case in local-spin density approximation, the timings are double that for the \(N\)-component case, the time grows linearly with \(N - 1\) due to extra matrix elements in \(S^{(2)}\).
constants can be done to get the minimum-energy potentials, or use the experimental lattice constants, and those potentials are then used for SRO predictions.

So, it is now possible both to predict SRO from the electronic structure and to interpret the SRO correctly by extracting the proper eigenvectors of $S^{(2)}(k, T)$ for arbitrary $N$-component alloys.

V. RESULTS

The prediction and characterization of chemical ordering instabilities in multicomponent alloys is of great practical and fundamental interest. The temperature-dependent chemical order is determined by the balance between ordering energy (favoring long-range order) and entropy (favoring disorder). The ordering energy largely reflects the underlying electronic effects within the disordered alloy. We have divided this section into two parts: the first validation of updated $S^{(2)}(k)$ code and generalized transformation approach to interpret SRO eigenvectors using well-studied binary and ternary systems, and the second focuses on the analysis of the high-entropy alloy candidate Al-Co-Cr-Fe-Ni quinary system. We report $k$-space wave vectors in units of $2\pi/a$ and real-space site coordinates in units of $a$. We calculate the KKR-CPA formation energy for a solid solution in a phase $x$ (e.g., $A1, A2,$ or $A3$) as $\Delta E_f^x (V) = E^x (V) - \sum_o c_o E^x_o (V^o),$ where the $E^x_o (V^o)$ is the energy of the alloying element in its ground-state phase and equilibrium volume. As discussed by Alam et al. [39], formation energy provides a good estimate of miscibility gap ($T_c = \Delta E_f^x / \delta_{ps}$) and order-disorder temperature ($T_{sp} = \Delta E_f^x / k_b$), where there is a cancellation in entropy to second-order from above and just below the transition at fixed concentration [39].

A. Validation and example analysis

We first validate our updated $S^{(2)}(k; T)$ calculations and new generalized SRO analysis by investigating a clustering binary and two ordering ternaries to show previous calculations and experiments are reproduced quantitatively.

$A1 Cu_{90}Ni_{50}$. Diffuse scattering and phase diagram experiments show that Ni-Cu has a clustering (phase segregation) tendency above the observed miscibility gap at 615 K [40]. An earlier SRO calculation [8] reported [000] instability at a spinodal temperature ($T_{sp}$) of 564 K. We repeated this and also found a $k_{sp} = [000]$ mode at $T_{sp}$ of 559 K. The KKR-CPA calculation of the $Cu_{90}Ni_{50}$ solid solution $\Delta E_f$ finds $2.90$ mRy, a positive value indicating phase segregation with estimated miscibility gap of 660 K. The solid solution $\Delta E_f$ and SRO-assessed $T_{sp}$ indicate segregation on a similar energy scale.

$A2 Nb-Al-Ti$. We apply the new code and analysis to $A2 Nb-Al-Ti$ ternaries, and compare the SRO results and cluster variation method (CVM) calculations by Johnson et al. [12], neutron scattering results by Jacob et al. [41,42], and TEM-ALCHEMI (atom location by channeling enhanced microanalysis) measurements by Fraser et al. [43–46]. We also found $B2$-type ordering for Nb-Al-Ti alloys with $[111]$ instability dominated by Ti-Al correlation, arising simply from hybridization and band filling, as found earlier. The eigenvectors at the instability estimate the occupational probability of each species at different sublattices in the ordered structure using (14).

Due to a SRO instability at $k_{sp} = [111]$, $A2 NbAlTi_2$ is unstable to ordering below the order-disorder temperature with a concentration wave given by

$$n^{Al}_{\eta} (r) = \frac{0.25}{0.81} \times e^{2\pi i (111) \cdot r}$$

and

$$n^{Al}_{\eta} (r) = \frac{0.25}{0.81} \times e^{2\pi i (111) \cdot r}$$

giving the probability distribution for $A2$ sites, i.e., cube corner at (000) and center at $(1/2, 1/2, 1/2)$. From (19) the first site probability that vanishes for a given $\eta(T)$ occurs for $n^{Al}(r = 000)$, giving a maximum LRO of $\eta_{max} = 0.25 (0.5 \times 0.81)^{-1} = 61.7\%$; it follows that the probability occupation of (Nb, Al, Ti) is $21.3$, $0$, $78.7\%$ at the cube corner and $(28.7, 50, 21.3)\%$ at the cube center. Clearly, the Nb/Al concentrations are reduced (enhanced) at corner (central) sites, and, as a result, the Ti increases to a maximum of $79\%$ at corners—a partially ordered $B2$ structure.

Analytically the ratio of eigenvectors of independent components (with respect to host) at unstable ($k_{sp}$) is

$$c^{Al}_{sp} (\{111\}) - c^{Al}_{sp} (000) = \frac{e^{2\pi i (111) \cdot k_{sp}}}{e^{2\pi i (111) \cdot k_{sp}}}$$

and, for a ternary, they may be graphically represented by the slope of a line in the Gibbs triangle, Fig. 2, known as ordering tie lines (OTLs) [46]. Our SRO results compare well with calculated and measured results. The calculated $T_{sp}$ of 1610 K is in good agreement with the measured 1713 K order disorder [41].

$A1 Cu_{2}NiZn$. Hashimoto et al. [47] and Van der Wegen and co-workers [48,49] showed first-order structural transitions from $A1 Cu_{2}NiZn$ occur at 774 K to a $[100]$-type partially ordered $L1_2$ structure. Althoff et al. [8] predicted $[100]$-type ordering corresponding to the partially ordered $L1_2$ phase with $T_{sp} = 980$ K, driven by strong Ni-Zn correlations arising directly from Fermi-surface nesting features, as also found here. With our newly developed analysis approach, we find a

![FIG. 2. (Color online) For A2 Nb-Al-Ti, concentration-wave polarizations (OTLs) are plotted for lines of short black (old theory), short blue (new theory), and green (ALCHEMI [43–46] extended to maximal permitted values). Neutron Rietveld-refinement results [41,42] are plotted by dotted lines. CVM B2-sublattice concentrations at $T/T_c = 0.9, 0.8, 0.7$ given by circles, squares, and triangles, respectively.](image-url)
similar partially ordered state at $T_p = 840$ K. The lower $T_p$ arises from the use of an optimal basis set in KKR-CPA-ASA.

### B. High-entropy alloy: Al-Co-Cr-Fe-Ni

For HEA formation, Zhang et al. [50] proposed three empirical criteria:

1. Mix $N \geq 5$ atoms in near-equiaatomic ratio for higher $\Delta S_p = -k_B \sum c_i \ln c_i \geq 1.60 k_B$.
2. Have atomic size ratio $\delta < 4.6$ (like Hume-Rothery size effect rule for solid solutions); here, $\delta = \sqrt{\frac{\sum c_i (1 - r_i/R)}{\sum c_i r_i}}$ for elemental radii $r_i$ and average radii $R$.
3. Small $-2.05 \delta - 1.94 < \Delta E_1 < -0.98 \delta + 4.14$ mRy to void compound formation.

Interestingly, so far, many multicomponent systems following the above criteria form simple solid-solution phases, e.g., A1, A2, or A3. For AlCoCrFeNi, using an empirical Miedema model by Ren et al. [51], we find $\Delta E_1 = -1.79$ mRy and $\Delta S_p = 1.61$, so this system obeys the criteria.

Chou et al. [52], in their observation for Al-Co-Cr-Fe-Ni, found that increasing Al % plays the role of phase stabilizer. For the reader’s convenience, we define a parameter $\Delta$ (in mole fraction) that controls the Al concentration in Al$_{\Delta/5}$CoCrFeNi$_{1-\Delta/5}$. Similar to Chou et al. [52] and Zhang et al. [50], we focused our attention to three regions: (1) An Al-poor region for $\Delta < 0.5$ with the A1 phase; (2) an Al-intermediate region for $0.5 \leq \Delta < 1.25$, which exhibits a mixed A1 + A2 phase; and (3) an Al-rich region for $1.25 < \Delta < 2.0$ with the A2 phase [53,54].

We performed KKR-CPA calculations in each region to study relative phase stability (by both VP and ASA methods to assess errors), and then used the KKR-CPA-ASA potentials and charge densities in the SRO calculations (the restriction in the SRO code). We chose one composition within each region: (1) a near-atocatomic quinary (N=5) for Al-poor; (2) an Al-intermediate region with $0.5 \leq \Delta < 1.25$, which exhibits a mixed A1 + A2 phase; and (3) an Al-rich region with $1.25 < \Delta < 2.0$ with the A2 phase [53,54].

For stability of A2 relative to A1, as shown in Fig. 3, both KKR-CPA VP and ASA results show that increasing Al% stabilizes the A2 phase in agreement with the experimental observations made by Chou et al. [52] and calculations of Zhang et al. [55] using CALPHAD (calculation of phase diagram) techniques [56,57]. The VP results provide a better description of charge and integrations within each VP site, improving the total energy. (For the ASA, errors in relative phase stability, e.g., A2 versus A1, is a known issue.) The small shift in $E^{A2} - E^{A1}$ from VP versus ASA improves agreement with that measured experimentally; see Fig. 3 (bottom). The common tangent to enthalpy curves shows Al % composition ($0.5 \leq \Delta < 1.25$) where two-phase A1 + A2 equilibria occurs, which lowers the overall free energy of the homogeneous system into a weighted mix of two phases.

For selected compositions, we performed the linear-response calculations to identify unique SRO modes. We calculated $S^{(2)}_{\alpha\beta}(k, T)$ and determined $\alpha_{\mu\nu}(k)$ for each composition. The $S^{(2)}_{\alpha\beta}(k, T)$ stability matrix is formulated in a “host” picture for mathematical and computational expediency. For ease of interpretation, we convert from host to the “off-diagonal” representation [22] so that the SRO corresponds to all individual pairs directly.

Notably, for binaries, the unstable wave vector in $\alpha(k)$ is the same as the favorable modes in $S^{(2)}(k)$. However, for complex ($N > 2$) alloys, this need not be the case. Due to the inversion in (6) and the competition between modes and intensity conservation (15), the eigenvectors in $S^{(2)}(k)$ may be different in $\alpha(k)$. In this case a careful analysis in real space can be helpful. In fact, unlike in a binary, a multicomponent alloy can have negative intensity in $\alpha_{\mu\nu}(k)$ relative to the homogeneous disordered case, exactly because of this competing nature between modes and intensity conservation. Examples of the effect appear below. In short, the instability in $\alpha_{\mu\nu}(k)$ may not reflect the pair’s driving instability, which shows the importance of present thermodynamic theory.

#### 1. A1 and A2 equiatomic quinary ($\Delta = 1.0$)

**A1 phase.** We first focus on the Warren-Cowley SRO parameters [5–7] $\alpha_{\mu\nu}(k)$ (or pair correlations) that may be measured experimentally. The diffuse maximal peaks at $k_{\text{max}} = \{100\}$ (X points) for $\alpha(k)$ indicates the periodicity of the ordering instabilities in the disordered alloy (not Bragg reflections). At $T_p$, particular elements of $\alpha(k)$ become unstable (diverge) and indicate second-order instability to LRO. The instability in $\alpha(k)$ is related to the peak in the stability matrix $S^{(2)}(k)$ in select pairs. The strongest pair in $S^{(2)}(k)$ driving ordering is Cr-Al, but Co-Cr is the dominant mode in $\alpha(k)$. Clearly, the dominant mode in $\alpha(k)$ is not the same as $S^{(2)}$, but Cr is involved in both competing modes and the strong ordering for one element must be accommodated by the pairs sharing that element.

Just like phonon modes, $k$-space representation provides relative stability of ordering modes. However, the $k$ space does not necessarily give a direct picture of underlying pair correlations in complex systems. Then, Fourier decomposition of such quantities into real space becomes important. The real-space-pair-correlation energies for the select pairs are shown...
in Table I, which are calculated via Fourier (shell-by-shell) transform,

\[ S_{\mu\nu}^{(2)}(k) = S_{\mu\nu}^{(2)}(0) + \sum_{i\sigma} S_{i\sigma,\mu\nu}^{(2)} e^{i k \cdot R_i}, \]

where \( n \) represents the shell number. Being much larger, Cr-Al pair is a dominant mode. The shell-by-shell calculation gives the strength and spatial extent of \( S^{(2)}(k) \).

The normal modes in Table II are shown at 1.2\( T_B \) (\( T_B = 448 \) K), which are the eigenvectors of \( S^{(2)}(k) \) driving divergence in SRO. The free energy cost to establish one of these modes vanishes at \( T_B \). Above \( T_B \) all eigenvalues remain positive, costing energy to the disordered state for substantiating concentration modulations while below \( T_B \), the critical eigenvalue establishes the anticipated probability distribution.

Similar to the NbAlTi\( _2 \) example, the probability distribution can easily be determined for systems with any number of components. Because SRO instability for A1 occurs at the star of wave vector \( k_{so} = (001) \), i.e., \((100), (010), \) and \((001)\), the system is unstable to ordering into a \( L_1 \)-like superstructure, where \( \gamma = \frac{1}{2} \) by symmetry. The concentration wave is then

\[ \left[ \begin{array}{c} n_{Al}(r) \\ n_{Co}(r) \\ n_{Cr}(r) \\ n_{Fe}(r) \end{array} \right] = \left[ \begin{array}{c} 0.20 \\ 0.20 \\ 0.20 \\ 0.20 \end{array} \right] \times \left[ \begin{array}{c} 0.20 \\ 0.20 \\ 0.20 \\ 0.20 \end{array} \right] + \frac{1}{4} \eta(T) \times \left[ \begin{array}{c} -0.21 \\ +0.28 \\ +0.32 \\ -0.60 \end{array} \right] \]

\[ \times [e^{2\pi i(100)\cdot r} + e^{2\pi i(010)\cdot r} + e^{2\pi i(001)\cdot r}]. \]

which gives the probability distribution for A1 sites, i.e., the cube corner at \((000)\) and faces at \([0, \pm \frac{1}{2}, \pm \frac{1}{2}]\). The maximum possible LRO is given by the site probability that vanishes first; here, that occurs for \( n_{Fe} \) at \((000)\) giving \( \eta_{max} = 0.21 \times 0.61^{-1} = \frac{1}{4} = 0.44 \). Thus, the maximum probability occupation of Al, Co, Cr, and Ni at the cube corner is 13, 29, 31, 0, and 27%, respectively. Similarly, the occupation at other sites can be evaluated from (22).

For A2 with \( k_{so} = [111] \), the system is unstable to a \( B_2 \) superstructure with \( \gamma = \frac{1}{4} \) by symmetry. The concentration wave, corresponding to the eigenvector of \( A^{(2)}(k) \) driving divergence in SRO, i.e., \( E_4 \), is

\[ \left[ \begin{array}{c} n_{Al}(r) \\ n_{Co}(r) \\ n_{Cr}(r) \\ n_{Fe}(r) \end{array} \right] = \left[ \begin{array}{c} 0.20 \\ 0.20 \\ 0.20 \\ 0.20 \end{array} \right] + \frac{1}{2} \eta(T) \times \left[ \begin{array}{c} +0.20 \\ -0.46 \\ +0.04 \\ -0.27 \end{array} \right] \times e^{2\pi i(111)\cdot r} \]

TABLE II. With Ni as host, the normal modes in Gibbs space for A1 AlCoCrFeNi at 1.2\( T_B \) \((\text{Fe} = 448 \text{ K})\). \( E_3 \) is the mode corresponding to vanishing eigenvalue (highlighted) and used for concentration wave analysis in (14).

<table>
<thead>
<tr>
<th>Comp.</th>
<th>( e_{Al} )</th>
<th>( e_{Co} )</th>
<th>( e_{Cr} )</th>
<th>( e_{Fe} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( E_1 )</td>
<td>+0.15</td>
<td>-1.09</td>
<td>-0.78</td>
<td>-0.20</td>
</tr>
<tr>
<td>( E_2 )</td>
<td>+0.83</td>
<td>+0.77</td>
<td>-0.83</td>
<td>-0.08</td>
</tr>
<tr>
<td>( E_3 )</td>
<td>-0.21</td>
<td>+0.28</td>
<td>+0.32</td>
<td>-0.60</td>
</tr>
<tr>
<td>( E_4 )</td>
<td>+1.11</td>
<td>-0.39</td>
<td>+0.78</td>
<td>-0.03</td>
</tr>
</tbody>
</table>

FIG. 4. (Color online) For \( \Delta = 1.0 \) (20% Al) in A1 phase, \( \alpha_{\mu\nu}(k) \) (upper) and \( S_{\mu\nu}^{(2)}(k, T) \) (lower) plotted along \( \Gamma-X-W-K-L-\Gamma \) of the fcc Brillouin zone.

FIG. 5. (Color online) For \( \Delta = 1.0 \) (20% Al) in A2 phase, \( \alpha_{\mu\nu}(k) \) (upper) and \( S_{\mu\nu}^{(2)}(k, T) \) (lower) plotted along \( \Gamma-P-N-\Gamma-H \) of the bcc Brillouin zone.
TABLE III. With Ni as host, the normal modes in Gibbs space for \( A_2 \) AlCoCrFeNi at 1.27\( T_g \) (\( T_g = 1705 \) K). \( E_4 \) is the mode corresponding to vanishing eigenvalue (highlighted) and used for concentration wave analysis in (14).

<table>
<thead>
<tr>
<th>Comp.</th>
<th>( e_{\text{Al}} )</th>
<th>( e_{\text{Co}} )</th>
<th>( e_{\text{Cr}} )</th>
<th>( e_{\text{Fe}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( E_1 )</td>
<td>-0.46</td>
<td>-0.84</td>
<td>-0.99</td>
<td>-0.15</td>
</tr>
<tr>
<td>( E_2 )</td>
<td>-0.45</td>
<td>+0.47</td>
<td>+0.22</td>
<td>-0.55</td>
</tr>
<tr>
<td>( E_3 )</td>
<td>-0.41</td>
<td>-0.94</td>
<td>+0.98</td>
<td>-0.01</td>
</tr>
<tr>
<td>( E_4 )</td>
<td>+1.20</td>
<td>-0.46</td>
<td>+0.04</td>
<td>-0.27</td>
</tr>
</tbody>
</table>

maximum probability of Al, Co, Cr, Fe, and Ni at the cube center is 0, 27.7, 19.3, 24.5, and 28.5%, respectively.

2. \( A_1 \) Al-poor quinary (\( \Delta = 0.395 \))

In Fig. 6, the \( \alpha(k) \) instability occurs at \( k_{us} = [000] \) (\( \Gamma \) point), indicating clustering involving two competing Cr-Al and Cr-Ni pair correlations. Clearly, however, \( S^{(2)}(k) \) shows that the strongest exchange energy is in the Cr-Ni pair, which drives clustering and slightly weaker Cr-Al (X-point) ordering energy. This energetics is manifest in \( \alpha(k) \) through Cr-based pair correlations (Cr-Al and Cr-Ni) to accommodate favorable \( S^{(2)}(k) \) clustering energy in Cr-Ni and weaker ordering energy in Cr-Al, so they are coupled. This effect is driven by the strongest Cr-Ni pair in \( S^{(2)}(k) \), but manifest in \( \alpha(k) \) in two closely competing modes, i.e., Cr-Al and Cr-Ni.

3. \( A_2 \) Al-rich quinary (\( \Delta = 1.6 \))

In Fig. 7, the instability in \( \alpha(k) \) occurs at \( k_{us} = [111] \) (\( H \) points) indicating ordering tendency in Al-Ni and Co-Al pairs. However, the peak in \( S^{(2)}(k) \) at the \( H \) point is driven by competing pairs Al-Ni, Co-Al, and Fe-Ni. Clearly, a small change in pair energies can significantly affect the Warren-Cowley parameter [5-7].

In the Al-rich region, the SRO for \( A_2 \) occurs at \( k_{us} = [111] \) indicating an ordering instability to the B2-like superstructure with \( \gamma = \frac{1}{2} \) by symmetry. The concentration wave, corresponding to the eigenvector of \( S^{(2)}(k) \) driving divergence in

maximum probability of Al, Co, Cr, Fe, and Ni at the cube center is 0, 27.7, 19.3, 24.5, and 28.5%, respectively.

4. Electronic origin of SRO

For better understanding, KKR-CPA density of states results were plotted for \( \Delta = 0.395, 1.0, \) and 1.6 for \( A_1 \) and \( A_2 \) Al-Co-Cr-Fe-Ni systems and tried to connect general behavior to its electronic origin.

In Fig. 8, \( A_2 \) (\( \Delta = 1.0 \) and 1.6) has strong hybridized states compared to \( A_1 \) (\( \Delta = 0.395 \) and 1.0) which enhances the filling of bonding-type states below \( E_f \), while simultaneously pushing antibonding (or nonbonding) states above \( E_f \). But for the \( A_1 \) phase, the disorder broadening leads to weak hybridization, reducing the ordering strength. Clearly, increasing Al % stabilizes the \( A_2 \) relative to \( A_1 \), due to increased hybridization.

TABLE IV. With Ni as host, the normal modes in Gibbs space for Al-rich \( A_2 \) at 1.27\( T_g \) (\( T_g = 1190 \) K). \( E_4 \) is the mode corresponding to vanishing eigenvalue (highlighted) and used for concentration wave analysis in (14).

<table>
<thead>
<tr>
<th>Comp.</th>
<th>( e_{\text{Al}} )</th>
<th>( e_{\text{Co}} )</th>
<th>( e_{\text{Cr}} )</th>
<th>( e_{\text{Fe}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( E_1 )</td>
<td>-0.40</td>
<td>-0.82</td>
<td>-1.03</td>
<td>-0.15</td>
</tr>
<tr>
<td>( E_2 )</td>
<td>-0.32</td>
<td>+0.40</td>
<td>+0.23</td>
<td>-0.58</td>
</tr>
<tr>
<td>( E_3 )</td>
<td>-0.41</td>
<td>-0.97</td>
<td>+0.94</td>
<td>-0.02</td>
</tr>
<tr>
<td>( E_4 )</td>
<td>+1.25</td>
<td>-0.48</td>
<td>+0.04</td>
<td>-0.20</td>
</tr>
</tbody>
</table>
in the A2 phase resulting in an increased order-disorder temperature.

From our KKR-CPA-ASA, we can compare the $\Delta E_f$ and their changes with composition and structure (due to the ASA, trends with $\Delta$ and structure can be shifted from full potential results). At $\Delta = 0.395$, the positive $\Delta E_f = +5.19$ mRy for A1 shows the clustering nature of the alloy, with an estimated miscibility gap of 523 K. At $\Delta = 1.0$, the negative $\Delta E_f$ for A1 ($-4.26$ mRy) and A2 ($-7.66$ mRy) shows the ordering nature with estimated order-disorder temperature 679 and 1217 K, respectively. At $\Delta = 1.6$, negative $\Delta E_f$ for A1 ($-5.86$ mRy) also shows ordering behavior with estimated order-disorder temperature of 932 K. The estimated $T_{sp}$ from SRO calculations are 448, 437, 1705, and 1190 K, respectively. The spinodal temperature for A2 increases because of stronger hybridization. Note that, if SRO linear response had charge fluctuations incorporated (i.e., all double counting terms), then Madelung screening, if relevant, can change the magnitude of $S^{(2)}(k)$ and $T_{sp}$. Here, for the Al-Co-Cr-Fe-Ni systems only hybridization and band filling are significant in driving ordering or clustering.

VI. CONCLUSION

We have developed an algorithm based on mathematics of $(N-1)$ simplex to analyze uniquely the chemical SRO in $N$-component solid-solution alloys, i.e., thermodynamically induced ordering fluctuations. The eigenvalues associated with the SRO can be interpreted easily within a concentration-wave framework. In addition, we utilized a KKR-CPA based thermodynamic linear-response theory to predict the SRO for $N$-component solid solutions described by an inhomogeneous coherent potential approximation. We validated the new SRO code and analysis on experimentally and theoretically well-studied binary and ternary systems. We investigated SRO in quinaries Al-Co-Cr-Fe-Ni systems in three regions with different Al %, that is, Al poor with $\Delta < 0.5$, Al intermediate with $0.5 \leq \Delta \leq 1.25$, and Al rich with $1.25 < \Delta \leq 2.0$, all of which show simple solid-solution phases, i.e., A1, A2, or both. The thermodynamic predictions from our electronic-structure-based theories of formation energy and short-range order from linear response agree with all known measurements. Given that SRO is difficult to measure in HEA alloys, and has yet to be attempted, our validated theory provides predictiv methods to guide experiment and to assess properties for design of complex alloy systems.

For future design purposes, we propose to combine SRO prediction over the entire Gibbs space with structural stability (A1 versus A2 versus A3) and mechanical property estimates using stacking fault energy to narrow the search space for desired chemical and mechanical behavior. Notably, stacking fault calculations must include the Suzuki effect, [58,59] where solute is attracted (repelled) from the defect to lower (maintain) the defect energy, or otherwise the mechanical strength can be highly overestimated [60].

ACKNOWLEDGMENTS

We would like to thank B. Haber (University of Illinois Urbana-Champaign) for useful discussions on barycentric coordinates, and F. Pinski (University of Cincinnati) for helping to resurrect elements of our old SRO code. The work was supported by the U.S. Department of Energy (DOE), Office of Science, Basic Energy Sciences, Materials Science and Engineering Division. The application of this method to the specific quinary was partially supported by the DOE Office of Fossil Energy (Cross-cutting Research program). The research was performed at the Ames Laboratory, which is operated for the U.S. DOE by Iowa State University under Contract No. DE-AC02-07CH11358.

APPENDIX : CARTESIAN REPRESENTATION OF REGULAR SIMPLICES IN ARBITRARY DIMENSION: AN $(N-1)$ SIMPLEX

In geometry, a triangle or higher-order polygon can be generalized into a polytope of arbitrary dimension, also known as simplex [61]. A $(N-1)$ simplex is defined as a geometric object that has $N$ vertices and $N(N-1)/2$
edges. By convention, a \((N-1)\) simplex has \(N\)-barycentric coordinates defined to sum to 1, and, as such, the \(N\)th coordinate is redundant (linearly dependent).

A regular \((N-1)\) simplex is a polygon with equal edge lengths and \(N\) vertices, so compositions \(\{c_i\}\) of constituents in an \(N\)-component alloy are natural barycentric coordinates with \(c_N\) a dependent variable, as to be used in a "host" picture. For a ternary system, for example, the Gibbs (equilateral) triangle in composition space uniquely describes compositions of the alloy problem.

### 1. General formula for the height of \(N\) simplex

First, we consider two-simplex, i.e., equilateral triangle, with \(A_1A_2 = A_2A_3 = A_3A_1 = 1\) (unit length); see Fig. 9. The height of such a triangle is \(h = \cos \frac{\pi}{3}\). Angle \(X = \angle A_2A_1A_3\) is the dihedral angle of the regular two-simplex and defined by scalar product, \(A_1A_2 \cdot A_1A_3 = \cos X = \frac{1}{2}\), which results in \(h = \sqrt{\frac{1}{2}(1 + \cos X)} = \frac{\sqrt{3}}{2}\).

The relation between the height of a regular \((N-1)\) simplex and its dihedral angle is the same as for a two-simplex,

\[
h = \cos \frac{X}{2} = \sqrt{\frac{1}{2}[1 + \cos X]} = \frac{N}{2(N-1)}.
\]

Here, we use the elementary result for the dihedral angle [62], i.e., \(\cos X = (N-1)^{-1}\).

### 2. Cartesian coordinates of vertices of \(N\) simplex

In \((N-1)\) simplex, the Cartesian coordinates of each vertex \(A_N\) with respect to centroid "\(o_N^{-1}\)" can be determined with following conditions:

1. For a regular simplex, the distances of each vertex from center will be a constant.
2. The scalar product of two vertices of \((N-1)\) simplex with respect to its center \((\frac{1}{N-1}, \frac{1}{N-1}, \ldots, \frac{1}{N-1})\) will give the angle subtended, i.e., \(\arccos \left[-\frac{1}{N-1}\right]\) [62,63].

Here, in Fig. 9, \(A_1, A_2, \) and \(A_3\) are equivalent vertices with length "\(a\)" from centroid "\(o_2\)" and angle subtended is \(\arccos \left[-\frac{1}{2}\right]\).

In a host picture for a \(N\)-component alloy, the \(T\) matrix is rank \((N-1)\) and must be evaluated with respect to \(N\)th vertex of regular \((N-1)\) simplex. Here, the coordinate of all vertices are known in terms of center as the origin, so we translate the origin from center to the \(N\)th vertex. Next, the vertex coordinates are scaled by the height of \((N-1)\) simplex calculated analytically, i.e., \(\sqrt{N/2(N-1)}\). This cumulative result can be used for any alloy without change.

It follows that the coordinates of \((N-1)\) vertices are analytically given by the relation (11)

\[
X_i^j - X_i^N = -\frac{1}{X_j} \sqrt{\frac{2(N-1)}{N}} \left[X_{i-1}^j \cdot X_{i-1}^j + \frac{1}{N(N-1)}\right],
\]

where each \(i,j\) runs from 1 to \(N-1\). For example, \(T\) for binary (12), quaternary, and quinary (13) are given in the text. This \(T\) matrix for \(N\)-component alloy ensures that the Gibbs' eigenvectors, after transformation, are both algebraically and geometrically orthogonal (see the main text). This transformation is used to determine eigenvectors, SRO, and concentration waves, and guarantees that \(0 \leq c_i \leq 1\).


