Accurate estimation of a phase diagram from a single STM image

Kazuhito Takeuchi, Koretaka Yuge, Shinya Tabata, Hiroki Saito, Shu Kurokawa & Akira Sakai

We propose a new approach to constructing a phase diagram using the effective Hamiltonian derived only from a single real-space image produced by scanning tunneling microscopy (STM). Currently, there have been two main methods to construct phase diagrams in material science: ab initio calculations and CALPHAD with thermodynamic information obtained by experiments and/or theoretical calculations. Although the two methods have successfully revealed a number of unsettled phase diagrams, their results sometimes contradicted when it is difficult to construct an appropriate Hamiltonian that captures the characteristics of materials, e.g., for a system consisting of multiple-scale objects whose interactions are essential to the system’s characteristics. Meanwhile, the advantage of our approach over existing methods is that it can directly and uniquely determine the effective Hamiltonian without any thermodynamic information. The validity of our approach is demonstrated through an Mg–Zn–Y long-period stacking-ordered structure, which is a challenging system for existing methods, leading to contradictory results. Our result successfully reproduces the ordering tendency seen in STM images that previous theoretical study failed to reproduce and clarifies its previously unknown phase diagram. Thus, our approach can be used to clear up contradictions shown by existing methods.

Exploring new high-functional materials has remained a greatly challenging theme for several decades. Phase diagrams play very important roles in their manufacture. To obtain phase diagrams, both experiments and simulations have to be used. One can roughly classify the methodologies based on simulations into two main types: one is CALPHAD using thermodynamic information obtained by experiments and/or other simulation methods, and the other is ab initio calculations without experiments. In particular, for estimating alloy phase diagrams based on microscopic information, ab initio calculations based on density functional theory (DFT) are now most widely used to determine the interactions of a given system. DFT has been applied to bulk, surface, and interface systems and has achieved a number of successful results.

Meanwhile, there remain challenging systems where the results of ab initio calculations contradict with those of experiments. Such contradictions can particularly occur in systems where long-range and/or many-body interactions between multiple-scale objects (e.g., atoms and coarse-grained particles) are important because it is typically difficult to determine an effective multiscale Hamiltonian that appropriately describes the behavior of structural phase transitions. For DFT, it can be practically unfeasible to obtain an accurate Hamiltonian, especially when long-range interactions on a coarse-grained scale come into play, mainly due to the restricted size of the cells used; for experiments, essential difficulties arise in dividing thermodynamic information into individual interactions. One representative system, that includes such contradictions is Mg-based long-period stacking-ordered structure (LPSO), which has recently attracted a great deal of attention due to its considerable mechanical performance. At the stacking-fault in LPSO, so-called nano-sized “clusters” consisting of multiple elements are arranged in Mg solvent, and this interface plays a central role in stabilizing the whole LPSO. Through ab initio calculations and Monte Carlo simulation, Kimizuka et al. determined the multiscale Hamiltonian with pair-wise intercluster interaction energies and revealed that the in-plane nanocluster-ordering tendency is mainly due to the repulsive interaction between nanoclusters. Although they partially explained the radial distribution function of clusters, the microscopic ordering tendency seen in STM images, i.e., chain-like cluster ordering shown in Fig. 1(b), was not quantitatively addressed. Minamoto et al. also constructed ternary bulk Mg–Zn–Y phase diagram using CALPHAD. However, since it is difficult to measure the thermodynamic information of a multiscale system, experiments to determine its Hamiltonian are hard to perform. Moreover, CALPHAD cannot show geometrical information concerning the ordering of nanoclusters.

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interactions behave repulsively under very small simulation cells, i.e., very dense cluster compositions. This tendency is similar to the result obtained by the previous DFT work \(^1\), which estimated interactions using a single-cluster approach even when large simulation cells, i.e., very sparse cluster compositions. This tendency is similar to the result obtained by the previous DFT work \(^1\), which estimated interactions using a single-cluster approach even when large simulation cells, i.e., very sparse cluster compositions.

The pair-formation energy of a \((m, L)\) cluster–cluster pair with an \(L \times L\) two-dimensional triangular lattice under a periodic boundary condition is defined as

\[
\Delta E(m, L) = E(m, L) - E_{\text{Mg}} - 2E_{\text{cluster}}(L).
\]

Here, \(E_{\text{Mg}}\) denotes the total energy of a Mg cluster, and \(E_{\text{cluster}}\) denotes total energy of a configuration consisting of a single cluster and Mg atoms on the rest lattice points in a considered cell. The definition of Eq. (1) corresponds with pair interactions used in the previous DFT work \(^1\) and reflects actual \(mNN\) cluster–cluster pair interactions. In Fig. 3, \(\Delta E(m, L)\) is shown for \(m = 6–8\) and \(L = 5–40\).

Figure 3 shows that positive values of \(\Delta E(6, 5)\), \(\Delta E(7, 6)\) and \(\Delta E(8, 7)\) imply that 6–8NN cluster–cluster interactions behave repulsively under very small simulation cells, i.e., very dense cluster compositions. This tendency is similar to the result obtained by the previous DFT work \(^1\), which estimated interactions using a single-cluster approach even when large simulation cells, i.e., very sparse cluster compositions. This tendency is similar to the result obtained by the previous DFT work \(^1\), which estimated interactions using a single-cluster approach even when large simulation cells, i.e., very sparse cluster compositions.

In order to confirm validity of \(V_m\) in Fig. 2(a), we investigated the pair-formation energy of a single \(mNN\) \((m = 6, 7, 8)\) cluster–cluster pair with an \(L \times L\) two-dimensional triangular lattice under a periodic boundary condition. The pair-formation energy of \(\Delta E(m, L)\) is defined as

\[
\Delta E(m, L) = E(m, L) - E_{\text{Mg}} - 2E_{\text{cluster}}(L).
\]

Here, \(E_{\text{Mg}}\) denotes the total energy of a Mg cluster, and \(E_{\text{cluster}}\) denotes total energy of a configuration consisting of a single cluster and Mg atoms on the rest lattice points in a considered cell. The definition of Eq. (1) corresponds with pair interactions used in the previous DFT work \(^1\) and reflects actual \(mNN\) cluster–cluster pair interactions. In Fig. 3, \(\Delta E(m, L)\) is shown for \(m = 6–8\) and \(L = 5–40\).

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disagreement is originate from insufficient consideration of dependence on cluster compositions in the previous DFT work. Therefore, the 6NN cluster–cluster interaction behaves attractively for dilute clusters. Although the previous result successfully showed qualitative landscapes of the radial distribution function of nanoclusters, it could not reproduce the quantitative microscopic ordering tendency seen in STM images (solid

Figure 2. (a) Energy coefficients estimated by geometrical information, i.e., the SROs of Zn–Y clusters evaluated from a single STM image on the right-hand side of Fig. 1. (b) Schematic illustration of mNN pairs \( (m = 4–8) \) at the interface.

Figure 3. Pair-formation energy of a single \( m \)NN cluster–cluster pair, as defined by Eq. (1), for each \( L \times L \) two-dimensional triangular lattice. The inset emphasizes the behavior of pair-formation energy in \( L = 5–8 \) simulation cells.
lines in Fig. 1). This is because the previous DFT work determined cluster–cluster interactions by dense-cluster simulation cells, whereas we found that interactions behave differently between dense and dilute cluster concentrations, as shown in Fig. 3. Now, let us confirm that our multiscale interactions in Fig. 2(a) can capture the characteristics of SRO of clusters at the interface. Figure 4 shows thermodynamically averaged SRO at equilibrium, as obtained by Monte Carlo simulation (see Methods for a more detailed procedure), and this SRO corresponds to the radial distribution function of clusters. The strong SRO of 6NN pair means that the number of Mg–Mg and cluster–cluster pairs are relatively larger than that of Mg–cluster pairs, i.e., clusters are arranged so as to be at each other’s 6NN position. Our SRO landscape agrees well with the radial distribution functions obtained by previous DFT and experimental work, especially for the peaks in 6, 9, and 15NN pairs. For further discussion, we took a snapshot at the interface in the simulation and compared it with an STM image in Fig. 1. From the STM image, we can clearly confirm chain-like cluster ordering, unlike the previous DFT work, which showed a uniform arrangement of clusters. As above, our snapshot quantitatively captures the features of STM images and supports the validity of our multiscale interactions.

The Mg$_{0.9489}$Zn$_{0.0511}$ interface phase diagram obtained by different three STM images is presented in Fig. 5(a), which shows the ordering tendency in the cluster-rich phase (Fig. 5(b)) through a first-order order-disorder phase transition. Since, the interplane interactions are relatively weak in 18R-type LPSO$^{19}$, it is valid to regard the interface of LPSO as an isolated two-dimensional system. As a result, the closed circle in Fig. 5(a), showing the point where the STM image in Fig. 1(b) is obtained, indicates that Fig. 1(b) is not a single phase but a order-disorder phase coexistence state.

In summary, we suggested a new approach based on microscopic geometric information for constructing phase diagrams in cases where this is difficult for conventional simulation methods. Using STM images, we demonstrated our approach through the interface of 18R-type Mg–Y–Zn LPSO. We clarified the contradictory information concerning cluster arrangements presented by experiments and simulations and successfully reproduced an interface phase diagram consistent with that obtained by experiment. Our approach is expected to become a powerful tool for modeling isolated systems subjected to experimental observations such as STM.

Methods
Sample preparation for STM images. The directional solidification process was applied to a Mg$_{85}$Zn$_6$Y$_9$ ingot. The ingot was annealed at 773 K for 168 h and quenched with water. We confirmed the presence of a diffraction spot corresponding to 18H-type LPSO by TEM observation.

Samples for STM observation were prepared by cutting the alloy ingot, typically into a reed shape of 8 mm $\times$ 5 mm $\times$ 0.5 mm. After the introduction of the sample into the preparation chamber of the low-temperature ultrahigh-vacuum STM (Unisoku 1200), we cooled the sample with a liquid-nitrogen flow for 10–15 min. Finally, we cleaved the sample using a pushing rod in the UHV chamber. The cleaved sample was immediately transferred into a pre-cooled observation chamber. All STM observations were carried out at liquid-nitrogen temperature (up to 77 K).

STM itself has enough spatial resolution to resolve each atomic position on the surface. However, due to some noise and thermal drift, the positions of the clusters determined by our method have some uncertainty. To demonstrate the accuracy of our method, we plotted relative positions of the clusters measured from all the clusters in a single STM image in Fig. 6. As clearly seen, the relative positions of the clusters construct a lattice; which indicate that the accuracy of positions is good enough to estimate SRO.

Generalized Ising model. In recent alloy studies with ab initio calculations, the generalized Ising model$^{20}$ (GIM) has often been used for constructing a coarse-grained Hamiltonian, which captures the characteristics of a system within a given accuracy. Since we are interested in how clusters would be arranged at the interface with
Figure 5. (a) $\text{Mg}_{1-x}\text{(Zn}_x\text{Y}_8)_x$ interface phase diagram. Black lines denote the phase-coexistent line between the ordered and disordered phases with error bars. Closed circle denotes a point where the STM image in Fig. 1(b) is obtained. (b) Ordered phase of the interface. Large and small circles denote Zn–Y clusters and Mg, respectively.

Figure 6. Relative positions of the clusters measured from all the clusters in a single STM image in Fig. 1 (Right).
Mg solvent, for simplicity, it is natural to consider the clusters as coarse-grained atoms, i.e., Mg and the center of cluster as +1 and −1 Ising-spin variables, respectively. In GIM, the total energy of a given configuration, \( \sigma \), is expanded by a set of complete and orthogonal basis functions, \( \{ \xi_k \} \), with indices

\[
E(\sigma) = \sum_k V_k \xi_k(\sigma),
\]

where \( V_k \) denotes an energy coefficient and \( k \) is an index specified as a symmetrically nonequivalent figure such as 1NN and 2NN pairs. In particular for a binary system, \( \xi_k \) corresponds to the average product of spin variables \( s_\alpha \) on \( k \) over all symmetrically equivalent \( k \) in an Ising-spin configuration

\[
\xi_k(\sigma) = \frac{1}{N_k} \sum_{k \in \sigma} \prod_{\alpha \in k} s_\alpha,
\]

where \( \alpha \) denotes a site on a given lattice and \( N_k \) denotes the number of \( k \) in the configuration. \( V_k \) is defined as the inner product between total energy and \( \xi_k \):

\[
V_k = \langle \xi_k(\sigma) \rangle E(\sigma) = \rho_k \sum_\sigma \xi_k(\sigma) E(\sigma) = \rho_k \sum_\sigma \xi_k(\sigma) \left( \sum_\sigma E(\sigma) \right).
\]

Here \( \langle \rangle \) denotes inner product for configuration space, \( \rho_k \) denotes a normalized constant for the inner product, and the term in \( \langle \rangle \) denotes over all possible configurations except for pair \( k \) (i.e., \( \sigma_{N-k} \)). Transformation from the second to the third line means that summation is taken in terms of possible configurations on pair \( k \) (i.e., \( \sigma_k \)) instead of a summation over all possible configurations of \( \sigma \). This transformation explicitly shows us that when a given system includes many-body interactions \( \sum \sigma_{N-k} E(\sigma) \) should include the many-body interactions except for that of \( k \). Therefore, even though in this study pair-wise \( k \) is only considered, our method correctly treats essential many-body interactions.

**Short-range-order in our simulation.** We calculated the SROs of clusters using the Metropolis algorithm with a 100 × 100 two-dimensional triangular lattice under fixed composition. Note that, to explicitly consider the size of the nano-sized clusters without loss of validity, interactions of 1–3NN pairs are configured as having a very high so as not to overlap with each other.

**Determination of many-body interactions from a single STM image.** We first assume that the measured structure obtained by STM is in thermodynamic equilibrium. Then, the expectation value of the configurational density of states in an Ising-spin configuration on \( k \) over all symmetrically equivalent \( k \) is

\[
\xi_k(T) = Z^{-1} \sum_\sigma \xi_k(\sigma) \exp \left( \frac{-E(\sigma)}{k_B T} \right),
\]

where summation is taken over all possible configurations \( \sigma \), and \( Z \) denotes a partition function.

Under these conditions, we have recently derived the relationship between the structure, \( Q(T) = \{ (\xi_i(T), \ldots, \xi_j(T)) \} \) and the energy coefficients in Eq. (4), \( V = \{ V_1, \ldots, V_5 \} \), in an explicit matrix form:

\[
Q_{\text{ave}} + \Gamma(T) \cdot V \simeq Q(T)
\]

\[
\Gamma_\alpha(T) = -\frac{1}{k_B T} S_\alpha,
\]

where \( Q_{\text{ave}} \) represents a linearly averaged configuration and \( S_\alpha \) denotes an element of the covariance matrix for the configurational density of states in \( (\xi_i, \xi_j) \) two-dimensional space for a non-interacting system. Both of these quantities can therefore be known a priori without any information about energy or temperature. From Eq. (6), we can thus directly determine the energy coefficients defined in Eq. (4) from a measured structure using the equation

\[
V \simeq \Gamma^{-1} \cdot (Q(T) - Q_{\text{ave}}).
\]

**Constructing the interface phase diagram.** Recently, we proposed a new method based on the Wang-Landau algorithm to construct an alloy phase diagram. Since the Wang-Landau algorithm can accurately estimate phases that conventional Monte Carlo methods overlook, this method is suitable for estimating the unknown interface phase diagram.
In this study, the interface is regarded as a Mg–cluster pseudo-binary alloy. To construct the alloy phase diagram, a semi-grand-canonical (SGC) ensemble is more often used than grand-canonical or canonical ensemble. In an SGC ensemble, compositions can vary under the number of atoms, \(N\), kept fixed, i.e., we handle the difference of chemical potentials for each constituent element, \(\mu = \mu_{\text{cluster}} - \mu_{\text{Mg}}\), in spite of handling compositions. The total energy, \(E_{\text{SGC}}\), is defined as \(E_{\text{SGC}} = E - \mu_N x\), and the free energy, \(\phi\), is defined as \(\phi = -k_B T \ln Y\), with Boltzmann constant \(k_B\) and the partition function in the SGC ensemble \(Y\) defined as:

\[
Y(T, \mu) = \sum_{E_{\text{SGC}}} W(E_{\text{SGC}}) \exp \left( \frac{-E_{\text{SGC}}}{k_B T} \right),
\]

where \(W\) denotes density of states (DOS). Note that Helmholtz free energy \(F\) in the canonical ensemble is related to the free energy \(\phi\) in the SGC ensemble through Legendre transformation:

\[
\phi = F - \mu_N x.
\]

We can obtain \(x\) using partial differentiation by interpolating \(\phi\) for each chemical potential:

\[
x = -\frac{\partial \phi}{\partial (\mu N)}. \quad (10)
\]

The above procedure compensates for a disadvantage of the Wang-Landau algorithm, which cannot not directly calculate a thermodynamically averaged value unlike the Metropolis algorithm.

We estimated the DOS for a \(30 \times 30\) two-dimensional triangular lattice with single-spin-flip update so as to prevent nanoclusters from overlapping with each other. Using the DOS, Eqs (8) and (10), we obtained the free energy \(F\) in the canonical ensemble and estimated its phase diagram.

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Author Contributions
S. Kurokawa and A. Sakai conceived and designed STM experiments and estimated short-range-order of clusters. H. Saito prepared the samples and took STM images. S. Tabata did data processing of STM images and determined the position of the clusters. K. Yuge modeled the effective Hamiltonian via the method with geometrical information. K. Takeuchi calculated thermodynamic properties and constructed the phase diagram using the effective Hamiltonian.

Additional Information
Competing Interests: The authors declare no competing interests.

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