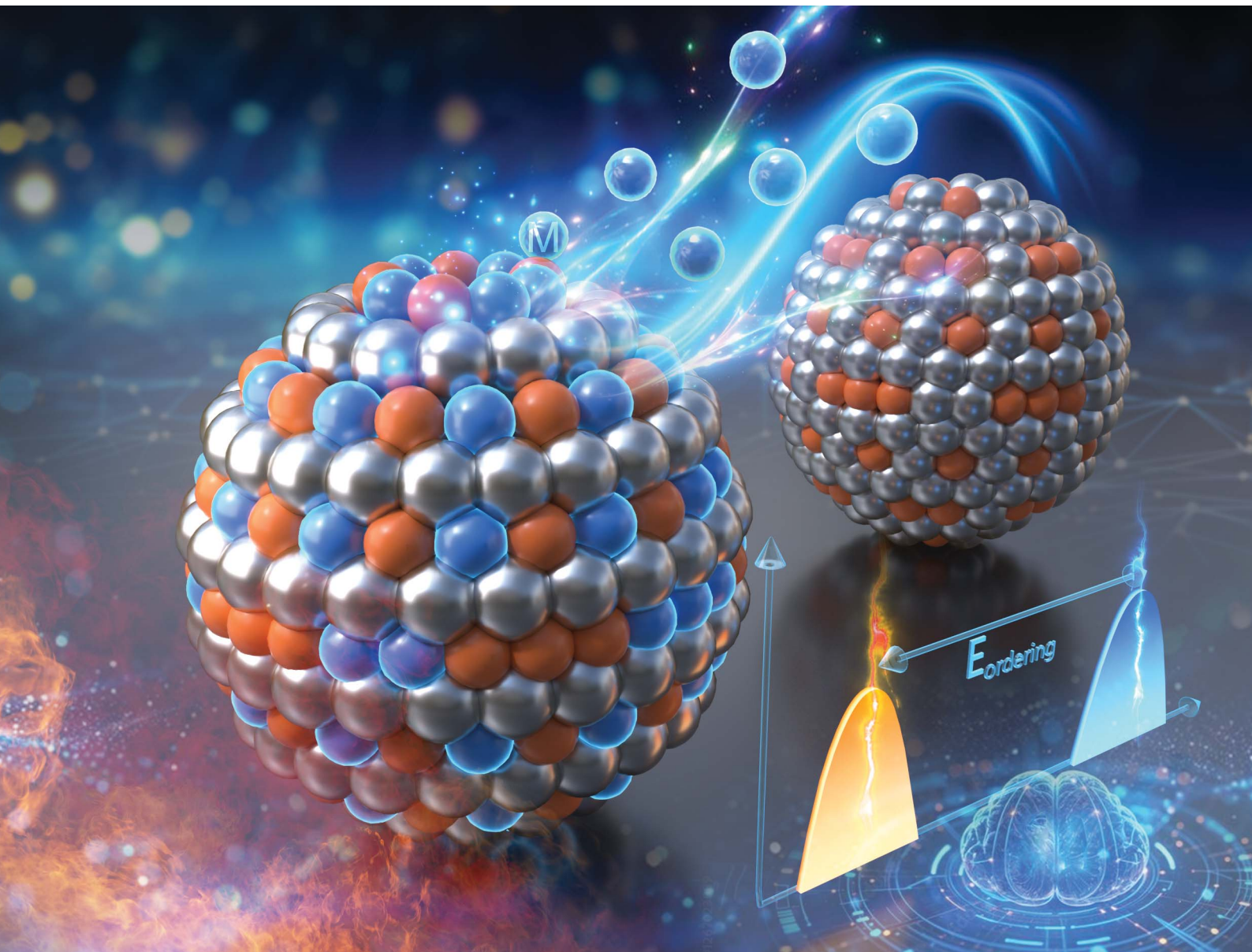


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Unraveling disorder-to-order transitions and chemical ordering in PtCoM ternary alloys using machine learning potential†

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PtCo intermetallic alloy nanoparticles are highly active and stable catalysts for the oxygen reduction reaction (ORR), making them key materials for proton-exchange membrane fuel cells. However, the high-temperature annealing required for ordering into the intermetallic phase often leads to particle growth. In this work, we developed a machine learning interatomic potential to model the disorder-to-order transition in PtCo-based ternary alloys with high accuracy and computational efficiency. Monte Carlo simulations reveal that introducing a third element significantly affects both the ordering process and the critical temperature for the disorder-to-order transition. The thermodynamic driving forces for ordering in various PtCoM alloys were systematically investigated to identify potential high-performance PtCoM catalysts. Kinetic analysis further indicates that the accelerated ordering transition in PtCo alloys is primarily driven by lower migration energy barriers and enhanced directional diffusion. These findings provide valuable atomic-scale insights into the chemical ordering mechanisms and suggest a pathway for designing highly ordered PtCo-based nanoparticles for energy conversion and storage applications.

Introduction

Proton-exchange membrane fuel cells (PEMFCs) are promising clean energy technologies with significant potential in transportation and power generation applications.^{1–5} However, the sluggish kinetics of the oxygen reduction reaction (ORR) at the cathode significantly limit their efficiency, necessitating the use of platinum (Pt)-based catalysts for optimal performance.^{6–8} The high cost and scarcity of Pt remain key barriers to the large-scale adoption of PEMFCs.⁴ Alloy catalysts, which combine Pt with other metals such as Fe, Co, and Ni, present a feasible solution by enhancing catalytic activity through electronic and ligand effects while reducing Pt content.^{9–13} This approach is crucial for achieving high performance at a lower cost, which is essential for the broader commercialization of PEMFCs.

Intermetallic alloys refer to crystalline structures in which the constituent elements are arranged in an ordered and uniform manner. It is widely reported in the literature that ordered intermetallic compounds exhibit superior activity and stability compared to their disordered counterparts.^{14–17} The enhanced activity and stability of intermetallic compounds are

attributed to stronger electronic interactions between Pt and the doped metals, greater compressive strain, and lower formation enthalpy.^{15,18} Although ordered intermetallic compounds are thermodynamically more stable, the synthesis of multicomponent metallic nanocrystals through wet-chemical methods often results in disordered solid-solution alloys or phase separation due to the varying reduction potentials of the constituent metals.¹⁹ Therefore, annealing at elevated temperatures is commonly employed to facilitate the disorder-to-order transition. However, this process frequently causes catalyst sintering, leading to particle growth and a consequent reduction in the atomic utilization of noble metals.^{20,21}

Several strategies have been explored to mitigate particle sintering during the disorder-to-order transition, including enhancing the interaction between nanoparticles and the support through sulfur atoms,¹⁵ forming protective shells around the particles using molecular additives,^{22,23} and elaborately controlling the heating and cooling rates.²⁴ Additionally, introducing a third element, such as Ga, Cu, or Ni, into bimetallic alloys has proven effective in promoting the ordered transition of the alloy.^{25–28} Previous studies have primarily focused on representative ordered and disordered structures, identifying trends through key indicators such as formation enthalpy and migration energy barriers derived from first-principles calculations.^{24,26,29} The use of empirical potentials and global optimization algorithms has further expanded the scope of investigations, offering alternative approaches to studying the transition process.^{30–33} Despite these advancements, microscopic insights into the processes governing

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disorder-to-order transitions and the chemical ordering of ternary alloys remain limited. These challenges are further complicated by the wide range of alloy configurations and the substantial computational resources needed to achieve statistically significant results with first-principles accuracy.

Recent advances in machine learning potentials have introduced a novel approach for modeling interatomic interactions that combines the precision of first-principles calculations with the computational efficiency of empirical potentials.^{34,35} In this work, we developed a machine learning interatomic potential using transfer learning, based on a pre-trained large atomic model.³⁶ This machine learning potential enables efficient Monte Carlo simulations of the PtCoM disorder-to-order transition process with first-principles accuracy, providing atomic-scale insights into the ordering process. Our work not only advances the understanding of the mechanisms driving chemical ordering but also provides a scalable framework for investigating other alloy systems, facilitating the rational design and optimization of advanced materials.

Results and discussion

Development of machine learning potential

The machine learning potential were trained by fine-tuning the Deep Potential model with a gated attention mechanism (DPA-1), as illustrated in Fig. 1a.³⁶ The dataset for fine tuning includes alloy structures (PtCo and PtCoM) with various degrees of chemical ordering (Fig. S1–S3 and Table S1†). The final dataset contains a total of 36 331 frames from the optimization calculations, which were divided into a training set and a validation set in a 9 : 1 ratio. The convergence process of the model's accuracy is shown in Fig. 1b. A total of 5 million training steps were performed. After these steps, the errors for both the training and validation sets became nearly identical, indicating that no overfitting occurred. Additionally, the model's accuracy plateaued, suggesting that 5 million training steps were sufficient. We evaluated different training strategies using the same training data and the same number of steps but different parameter fine-tuning schemes. The results demonstrate that directly applying the pre-trained DPA-1 model without any modification leads to substantial prediction errors (Fig. 1c). In contrast, fine-tuning all parameters of DPA-1 yields the highest accuracy, outperforming models trained from scratch (Fig. S4†). This performance gain can be attributed to the prior knowledge encoded in the pre-trained model, which provides a better initialization and facilitates more efficient parameter optimization compared to training from scratch. These findings underscore the efficiency and effectiveness of the pre-training and fine-tuning paradigm in developing accurate machine learning potentials.

Through pre-training and fine-tuning, a machine learning potential was developed, achieving an energy error of 0.0022 eV per atom and a force error of 0.04 eV Å^{−1} (Fig. 1d and e). We are confident that the model is sufficiently accurate to replace first-principles calculations, as its energy error surpasses those reported in existing literature for machine learning models applied to alloy systems.^{37–40} Additionally, its force error is

comparable to the force convergence criterion of 0.03 eV Å^{−1} commonly used in DFT structural optimizations. We also performed additional benchmarking against several representative interatomic modeling approaches for PtCoM ternary alloys. Specifically, we compared the energy prediction accuracy of the fine-tuned DPA-1 model with that of a fine-tuned MACE model, as well as more traditional approaches including cluster expansion and descriptor-based regression using Gaussian process regression. As summarized in Table S2,† the fine-tuned DPA-1 model achieves a lower energy MAE (0.0022 eV per atom) compared to the MACE model (0.0029 eV per atom), and outperforms both the cluster expansion (0.0035–0.0082 eV per atom) and the descriptor-based GPR model (0.0036–0.0058 eV per atom), which were trained on representative PtCoCu and PtCoNi subsets. These results highlight the accuracy and transferability of the fine-tuned DPA-1 model across a broader compositional space and support its use in our study.

Thermodynamics of disorder-to-order transition of PtCoM

Using the machine learning potential, we conducted virtual experiments *via* Monte Carlo simulations to investigate the disorder-to-order transition process of PtCoM at different temperatures. To evaluate the temperature dependence, we adopted a simplified thermodynamic model, considering the high computational cost of rigorous phase space sampling methods. Specifically, the free energy change ΔG was approximated as $\Delta G = \Delta H - T\Delta S_{\text{config}}$, where the enthalpy change ΔH was assumed to be temperature-independent, and the entropy contribution was limited to the configurational entropy ΔS_{config} . This approximation neglects vibrational and electronic entropy contributions, which are computationally demanding to compute *via* first-principles methods. Although more accurate evaluations could be achieved using molecular dynamics or metadynamics, these techniques remain impractical for high-throughput alloy screening due to their substantial resource requirements. Despite these simplifications, our approach captures the dominant thermodynamic driving forces and aligns with widely adopted methodologies in prior studies,^{24,41} offering a reasonable approximation for analyzing disorder-to-order transition trends in PtCoM systems. The enthalpy was calculated using the trained machine learning potentials, and the configurational entropy was determined based on chemical ordering using the method proposed by Yang.⁴² The initial atomic occupations in PtCoM were randomly assigned; after that, Monte Carlo simulations were performed to iteratively swap the positions of two atoms, allowing the system to evolve toward the configuration with the lowest free energy. The Warren–Cowley short-range order (SRO) was used to quantify the ordering degree, which is defined as:

$$\alpha^{A-B} = 1 - \frac{P^{A|B}}{C_B} \quad (1)$$

where C_B is the concentration of element B, and $P^{A|B}$ is the probability of finding element B among the neighbors of element A. A negative α^{A-B} value indicates an ordered atomic arrangement, while a positive α^{A-B} value reflects the segregation



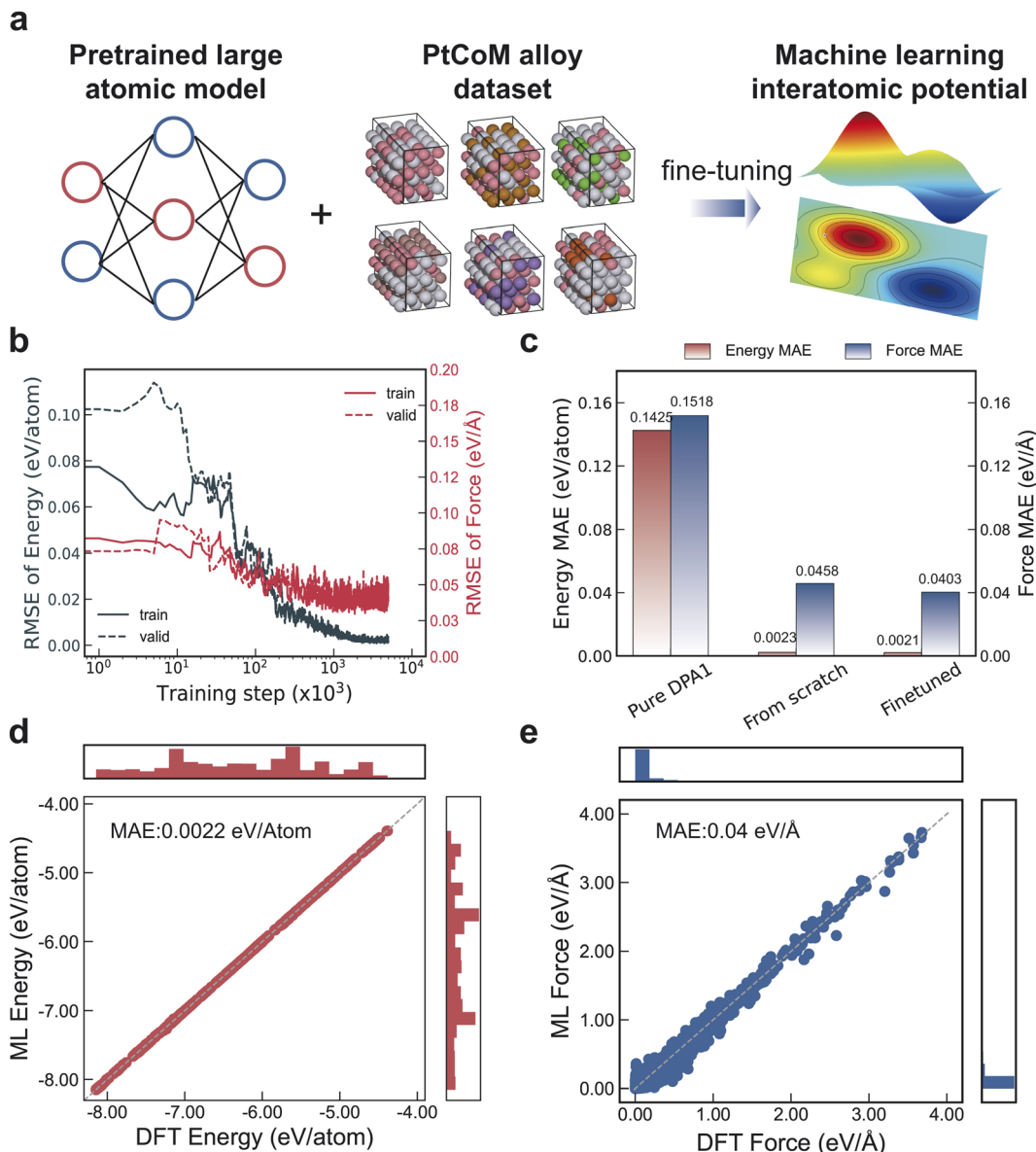


Fig. 1 Development and accuracy of the machine learning interatomic potential. (a) Schematic of the workflow for developing the machine learning potential for PtCoM. (b) Convergence curve of the fine-tuning process. (c) Comparison of accuracy for different machine learning potential models: 'Pure DPA1' refers to the DPA1 model whose parameters are not adjusted; 'From scratch' refers to training a new model with randomly initialized parameters, without inheriting any from the pre-trained DPA1 model; 'Finetuned' refers to training a new model that inherits its initial parameters from the pre-trained DPA1 model. (d) Energy error and (e) force error of the fine-tuned machine learning potential on the validation test.

of A and B. For our bulk model, the minimum and maximum values of $\alpha_{\text{Pt-CoM}}$ are $-1/3$ and $2/3$, corresponding to the complete ordering of Pt with Co/M and the complete aggregation of Pt with Co/M, respectively. Notably, the maximum value of $\alpha_{\text{Pt-CoM}}$, $2/3$, is due to the limited size of the supercells, which prevents it from reaching 1.

We monitored the evolution of chemical ordering during the disorder-to-order transition process of Pt_2CoCu at 600 K (Fig. 2a). Initially, Pt and non-Pt atoms, as well as Co and Cu atoms, are randomly distributed. As the simulation progresses, Pt and non-Pt atoms arrange in an increasingly ordered

manner, while Co and Cu atoms tend to segregate. This phenomenon can be attributed to the differing interactions between various atomic pairs. Notably, this behavior provides insights into multi-component intermetallic compounds, where certain element pairs still exhibit local disorder despite the overall trend towards ordering. Machine-learning-assisted Monte Carlo simulations provide atomic-level insights into chemical ordering since local short-range order is particularly challenging to observe directly in experiments—especially for elements like Co and Cu, which have similar physicochemical properties.⁴³

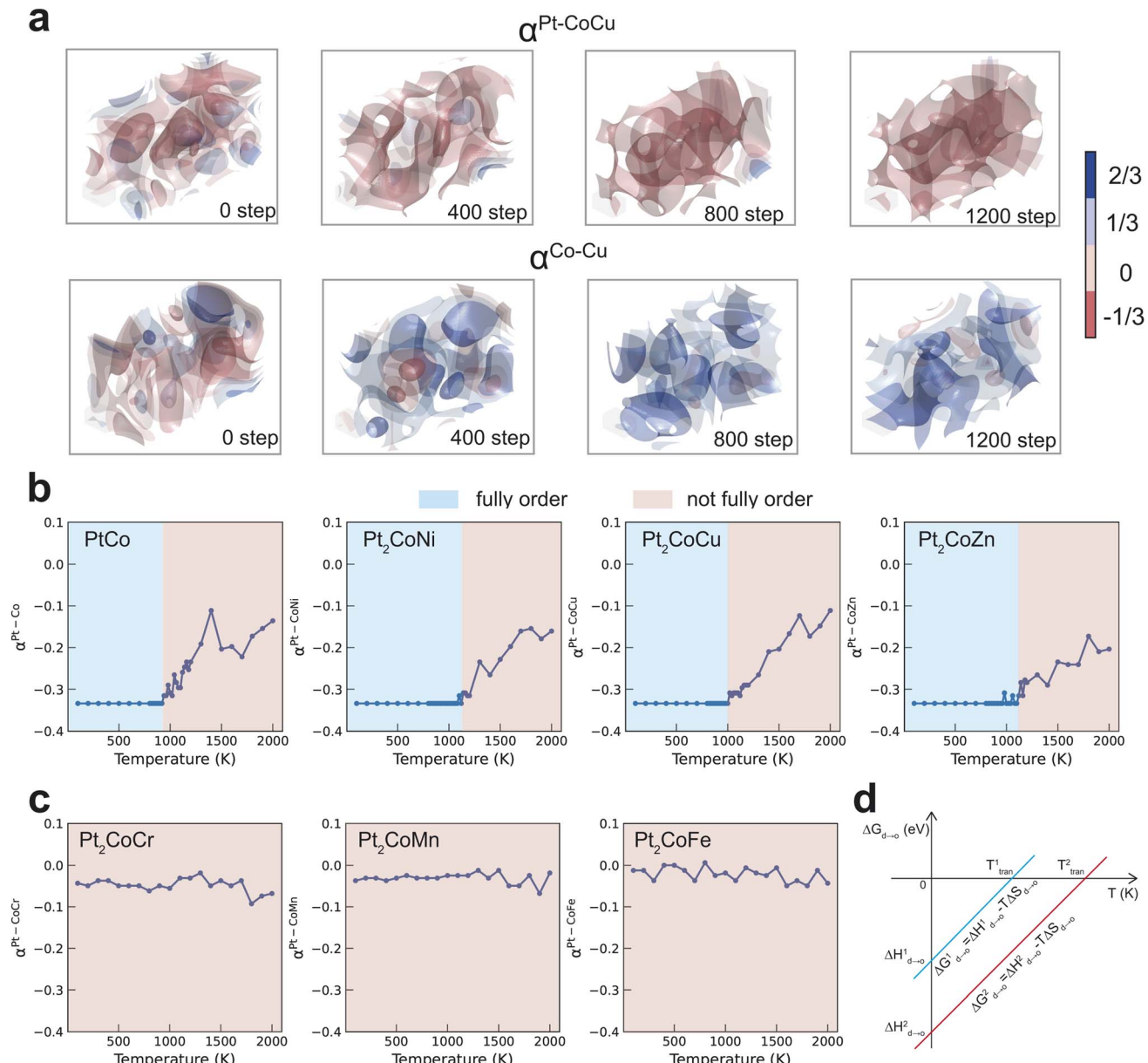


Fig. 2 Thermodynamic insights into the disorder-to-order transition of PtCoM. (a) The 3D distribution of $\alpha^{\text{Pt-CoCu}}$ and $\alpha^{\text{Co-Cu}}$ of PtCoCu at the beginning and after 400 steps, 800 steps, and 1200 steps of the Monte Carlo simulation at 600 K. (b) The final chemical ordering $\alpha^{\text{Pt-CoM}}$ of PtCo, Pt₂CoNi, Pt₂CoCu and Pt₂CoZn as a function of simulated temperature in Monte Carlo simulation. A critical temperature that separates the map into two distinct regimes: fully ordered regime and not fully ordered regime. (c) The final chemical ordering $\alpha^{\text{Pt-CoM}}$ of PtCoCr, PtCoMn and PtCoFe as a function of simulated temperature in Monte Carlo simulations. (d) Schematic diagram of the relationship between Gibbs free energy change ΔG , enthalpy change ΔH , and temperature T during the disorder-to-order transition.

Monte Carlo simulations were also performed at various temperatures for different Pt₂CoM systems, and the degree of ordering corresponding to the configuration with the lowest free energy is shown in Fig. 2b and c. These simulations indicate that for PtCo, Pt₂CoNi, Pt₂CoCu and Pt₂CoZn, there is a critical transition temperature. Below this temperature, complete ordering is attainable for Pt and Co/M ($\alpha^{\text{Pt-CoM}} = -1/3$), but above it, such ordering is not feasible ($\alpha^{\text{Pt-CoM}} > -1/3$). This critical temperature, which aligns with findings from other studies, is influenced by two primary factors.^{30,31,44} Firstly,

at elevated temperatures, the increased randomness in the Monte Carlo simulation trajectories hampers convergence to the global minimum energy state. Secondly, based on the free energy relation $\Delta G = \Delta H - T \Delta S_{\text{config}}$, the enthalpy change ΔH during the ordering process is assumed to be temperature-independent. The configurational entropy change ΔS_{config} is negative during the disorder-to-order transition, as disordered states have higher entropy than ordered ones. Consequently, the entropic term $-T \Delta S_{\text{config}}$ is positive and increases with temperature, which raises the free energy ΔG and destabilizes

the ordered phase at higher temperatures. For Pt_2CoCr , Pt_2CoMn and Pt_2CoFe , complete ordering is not achieved at any temperature. As shown in Fig. 2d, theoretically, a higher critical transition temperature T'_{tran} is generally associated with a larger thermodynamic driving force for ordering – namely, a more negative enthalpy change ΔH , which facilitates the disorder-to-order transition. The enhanced ordering observed upon the addition of Ni, Cu, and Zn can thus be attributed to their ability to increase this thermodynamic driving force relative to PtCo. In contrast, the incorporation of Cr, Mn, and Fe reduces the driving force, thereby suppressing the ordering tendency.

Element screening

In addition to the six elements discussed above, we conducted a systematic investigation on the effects of various other elements on the ordering process of PtCo, covering most transition metals from the fourth, fifth, and sixth periods. Lanthanide elements and technetium (Tc) were not included in this study. Tc is extremely scarce in the Earth's crust, and accurately modeling it using DFT presents significant challenges.⁴⁵ In addition, Sc, Y, Zr, and Hf were also excluded, as DFT optimizations for the corresponding Pt_2CoM structures failed to converge. Additionally, a general trend between the critical transition temperature and the thermodynamic driving force for ordering suggests that E_{ordering} can serve as a qualitative indicator for screening alloying elements. A higher E_{ordering} than that of PtCo indicates that incorporation of element M thermodynamically promotes ordering between Pt and Co/M. Even in cases where a well-defined transition temperature is absent—such as for Cr, Mn, and Fe—the ordering energy remains a meaningful measure of chemical ordering tendencies. This E_{ordering} based screening approach enables us to efficiently identify promising alloying elements without relying on time-consuming Monte Carlo simulations.

A total of 180 structures with Pt and Co/M arranged in a random pattern ($\alpha^{\text{Pt-CoM}} = 0$) were generated. Simultaneously, 180 structures with Pt and Co/M arranged in an ordered pattern ($\alpha^{\text{Pt-CoM}} = -1/3$), featuring varying degrees of ordering between Co and M ($\alpha^{\text{Co-M}}$ ranging from $-3/27$ to $25/27$), were also created. Their energies were calculated and compared to that of PtCo. We tested the effect of sampling size on the results, as shown in Table S3,[†] and found that increasing the number of sampled structures beyond 180 has a negligible impact on the calculated energies. As shown in Fig. 3a, compared to the ordered Pt–Co structure, which has a single energy value, the energies of the ordered Pt–CoM structures exhibit a broader distribution due to the varying degree of Co–M ordering. This spread provides an additional thermodynamic driving force that promotes the disorder-to-order transition. As shown in Fig. 3a, for the binary PtCo alloy, the ordering energy E_{ordering} is defined as the energy difference between the ordered structure with $\alpha^{\text{Pt-Co}} = -1/3$ and the average energy of randomly generated configurations with $\alpha^{\text{Pt-Co}} = 0$. For the ternary Pt_2CoM alloy, the concept is extended due to the presence of a third element. The total ordering energy can be expressed as $E_{\text{total}} = E_{\text{ordering}} + E_{\text{extra}}$, where the first term accounts for the energy

difference between the average energy of randomly generated Pt–Co/M disordered configurations ($\alpha^{\text{Pt-CoM}} = 0$) and the average energy of randomly generated Pt–Co/M ordered structures ($\alpha^{\text{Pt-CoM}} = -1/3$). The second term represents the additional stabilization associated with Co–M interactions, which is calculated as the energy difference between the average energy of Pt/CoM ordered structures, with Co/M ordering or disordering, and the average energy of the ordered structures ($\alpha^{\text{Pt-CoM}} = -1/3$).

We first systematically investigated the case where Pt is ordered with Co/M, examining how the chemical ordering of Co and M affects the energy of PtCoM (Fig. S5 and S6[†]). As shown in Fig. S5,[†] for certain elements such as Ti, V, and Nb, the energy increases linearly with increasing $\alpha^{\text{Co-M}}$, indicating a thermodynamic preference for these elements to adopt an ordered arrangement with Co. In contrast, for elements such as Ni, Cu, and Zn, the energy decreases linearly with increasing $\alpha^{\text{Co-M}}$, suggesting a tendency toward phase separation from Co. For a few elements, such as Ru and Re, no significant linear correlation is observed between energy and $\alpha^{\text{Co-M}}$. Fig. S6[†] summarizes the additional thermodynamic driving force arising from either ordering (Fig. S6b[†]) or segregation (Fig. S6c[†]) between Co and M. In Fig. S6b,[†] elements highlighted in red exhibit lower energy when ordered with Co compared to the average energy across different Co–M orderings, indicating a preference for ordering. Conversely, in Fig. S6c,[†] red-highlighted elements show lower energy in a segregated configuration, suggesting a tendency to phase-separate from Co. Overall, early transition metals with fewer d-electrons tend to order with Co, while late transition metals with more d-electrons tend to segregate. This behavior can be attributed to the relative bond strengths among Co–Co, M–M, and Co–M. Specifically, if the sum of the Co–Co and M–M bond energies ($E_{\text{Co-Co}} + E_{\text{M-M}}$) is smaller than twice the Co–M bond energies ($2E_{\text{Co-M}}$), an ordered arrangement is energetically favored. Otherwise, Co and M tend to segregate (Fig. S7[†]).

Fig. 3b presents the total ordering energy of Pt_2CoM relative to PtCo. Elements highlighted in red indicate an enhancement in the thermodynamic driving force for ordering, while those in blue suggest a reduction. Among the investigated elements, doping with Ti, V, Cr, Ni, Cu, Zn, Nb, Ag, Cd, Ta, Au, and Hg was found to increase the thermodynamic driving force for ordering between Pt and non-Pt components, suggesting their potential as favorable alloying elements. Notably, ordered phases incorporating Ti, V, Ni, Cu, Zn, Ag, and Au have already been experimentally synthesized, as summarized in Table S4.[†] For example, Liao *et al.* synthesized PtCoTi ternary nanoparticles (~ 3 nm) consisting of a $\text{Pt}_3\text{Co}_{0.6}\text{Ti}_{0.4}$ intermetallic core and three atomic layers of Pt shells using a modified impregnation–reduction approach. This catalyst exhibits superior activity and durability compared with Pt_3Co and Pt in rotating disk electrode (RDE) and single fuel cell tests.⁴⁶ Zhong *et al.* demonstrated the synthesis of highly ordered PtCoV nanoalloys *via* oxidative–reductive thermochemical treatments. At elevated temperatures, enhanced atomic-scale chemical ordering and reduced interatomic distances significantly improve ORR activity, achieving up to 1 A mg^{-1} of Pt in RDE measurements.⁴⁷



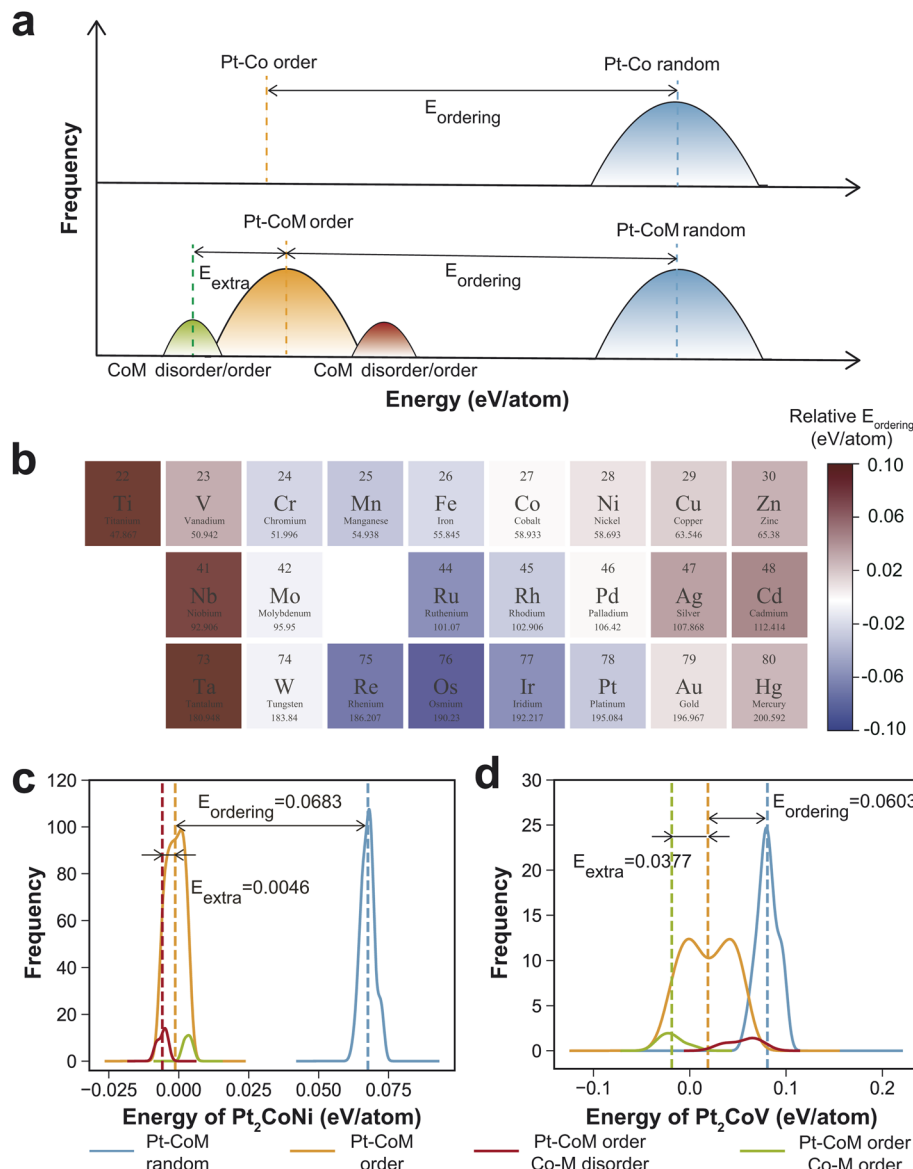


Fig. 3 Element screening in Pt_2CoM systems. (a) Schematic diagram of disorder-to-order driving energy E_{ordering} for Pt_2CoM and PtCo . (b) Comparative energy analysis of ordered ($\alpha_{\text{Pt-CoM}} = -1/3$) and randomly arranged ($\alpha_{\text{Pt-CoM}} = 0$) Pt and Co/M atom configurations, with energy values benchmarked to PtCo . (c) Disorder-to-order driving energy E_{ordering} for Pt_2CoNi . (d) Disorder-to-order driving energy E_{ordering} for Pt_2CoV .

Shen *et al.* reported the synthesis of highly ordered PtCoNi nanoparticles (~ 5.3 nm) through a controlled two-stage confinement strategy that enables both small particle size and high atomic ordering by resolving the conflict between atomic migration and spatial confinement. The resulting intermetallic PtCoZn catalyst exhibits exceptional ORR performance and durability in PEMFCs, attributed to enhanced Pt-Co coupling, high Pt surface exposure, and agglomeration prevention within mesoporous carbon walls.⁴⁸

Considering that the average energy difference between the ordered and random phases of Pt_2CoM relative to PtCo ranges from -0.11 to 0.03 eV per atom (Table S5†), the ordering or segregation of Co and M provides additional thermodynamic driving force, leading to a maximum energy decrease of 0.12 eV per atom for the ordered phase (Table S6†). This highlights the

critical role of Co-M chemical ordering in facilitating the disorder-to-order transition of Pt and non-Pt elements. As illustrated in Fig. 3b and c, Co-Ni segregation contributes a thermodynamic driving force of 0.0046 eV per atom in Pt_2CoNi , while Co-V ordering contributes a notably higher value of 0.0377 eV per atom in Pt_2CoV .

Kinetics of disorder-to-order transition of PtCoCu

The previous section investigated chemical ordering in thermodynamic equilibrium using Monte Carlo simulations based on atomic exchanges. However, atomic exchanges are primarily a computational tool for identifying stable configurations and do not directly reflect physical atomic migration mechanisms. In reality, atomic movement occurs *via* vacancy-mediated

diffusion, and the presence of vacancies plays a key role in facilitating the ordering process.⁴⁹ To enhance the accuracy of migration barrier predictions, we fine-tuned the machine learning potential by incorporating a small dataset of transition-state configurations obtained from climbing-image nudged elastic band (CI-NEB) calculations of vacancy migration in PtCo. As shown in Fig. 4a, the fine-tuned model achieves a prediction error of just 0.07 eV for the vacancy migration energy barrier in PtCo.

In FCC crystals, each atom has 12 nearest neighbors, giving rise to 12 possible vacancy migration pathways (Fig. 4b). By

combining the fine-tuned machine learning potentials with the CI-NEB method, we calculated 5000 migration energy barriers for Pt, Co, and Cu atoms migrating to neighboring vacancies in Pt₂CoCu and PtCo alloys, thereby investigating the kinetics of the ordering process. As shown in Fig. 4c, Cu atoms exhibit, on average, lower migration energy barriers compared to Pt and Co atoms in Pt₂CoCu, suggesting that the incorporation of Cu can accelerate the disorder-to-order transition. Furthermore, the average migration barriers of Pt and Co atoms in Pt₂CoCu (Pt: 1.07 eV; Co: 1.02 eV) are notably reduced compared to those in PtCo (Pt: 1.33 eV; Co: 1.17 eV), indicating that the presence of

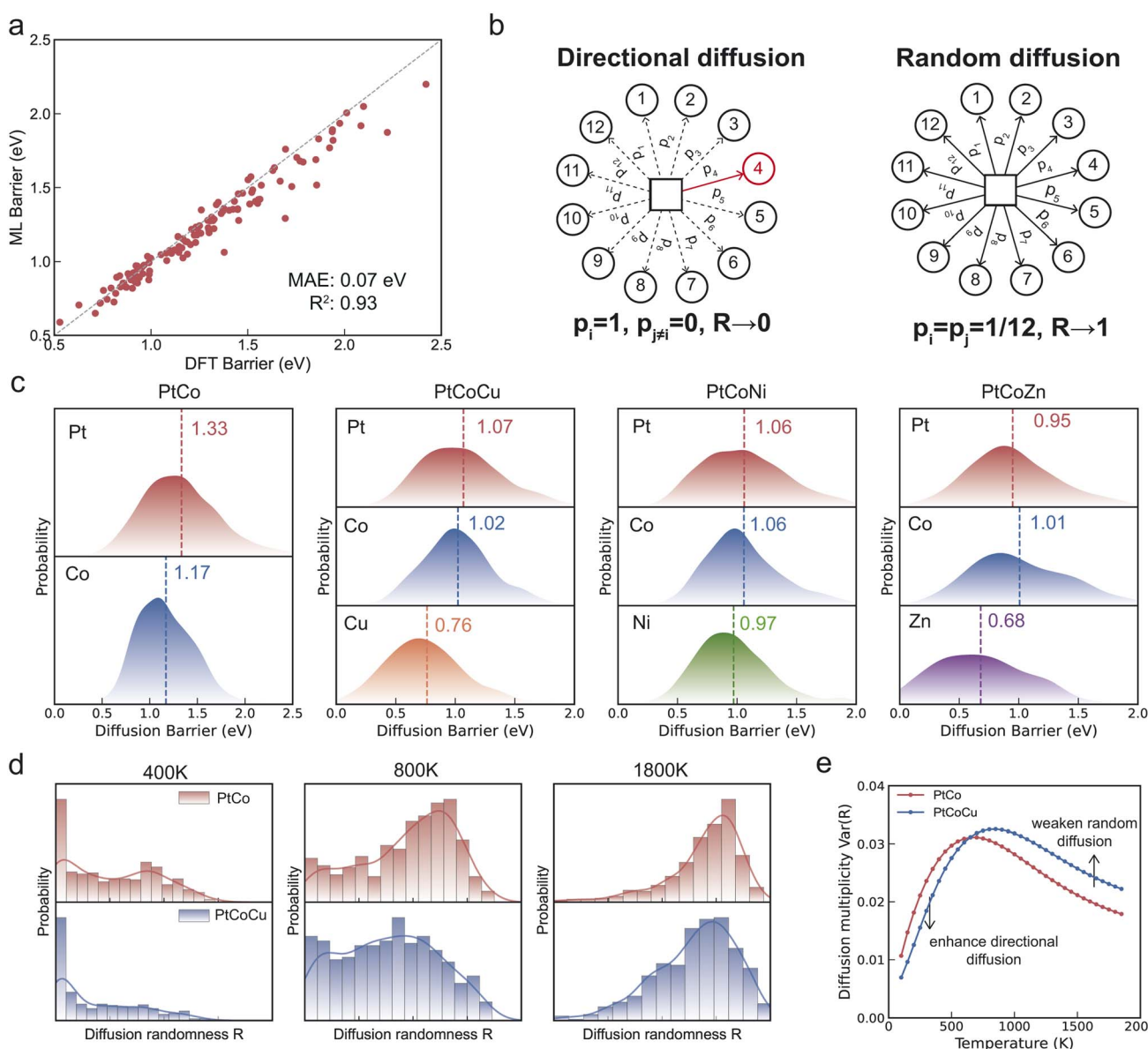


Fig. 4 Kinetic insights into the disorder-to-order transition of PtCoCu. (a) Comparison between the vacancy migration barrier of PtCo calculated using DFT and the machine learning potential. Distribution of atomic migration energy barriers for Pt, Co, and Cu. (b) Schematic representation of two extreme atomic migration modes. In one case, one of the twelve possible paths dominates with directional migration, $R = 0$; in the other case, all twelve paths have equal probability in a random migration scenario, $R = 1$. (c) Statistical distribution of machine learning predicted atomic migration energy barriers for Pt, Co, and M in PtCo, Pt₂CoCu, Pt₂CoNi and Pt₂CoZn; the horizontal dashed line in the figure represents the average migration energy. (d) The distribution of R of PtCo and Pt₂CoCu at different temperatures: 400 K, 800 K and 1800 K. (e) Migration multiplicity $\text{Var}(R)$ as a function of temperature for PtCo and Pt₂CoCu.



Cu not only enhances the mobility of Cu atoms themselves but also facilitates the diffusion of Pt and Co atoms, further promoting the ordering kinetics. To assess whether this kinetic promotion effect extends to other dopants, we performed additional vacancy-mediated diffusion barrier calculations for Pt₂CoNi and Pt₂CoZn. The results show that Ni and Zn also exhibit lower migration barriers compared to Pt and Co, with mean values of 0.97 eV for Ni and 0.68 eV for Zn, respectively. Importantly, the presence of these dopants similarly reduces the barriers for Pt and Co diffusion (Pt₂CoNi : Pt/Co = 1.06/1.06 eV; Pt₂CoZn : Pt/Co = 0.95/1.01 eV), indicating that the barrier-lowering effect is not unique to Cu but may be generalized to other dopants (Table S7†).

The probability of vacancy migration along each path is given by:

$$p_i = \frac{\exp(-E_i/k_B T)}{\sum_{i=1}^{12} \exp(-E_i/k_B T)} \quad (2)$$

where E_i is the migration energy barrier for path i , k_B is the Boltzmann constant, and T is the temperature. To characterize the distribution of these migration probabilities, we define the parameter R :

$$R = 1 - \sigma(\mathbf{p})/\max(\sigma(\mathbf{p})) \quad (3)$$

where $\sigma(\mathbf{p})$ denotes the standard deviation of the probability vector \mathbf{p} . At low temperatures, migration is dominated by the path with the lowest energy barrier, resulting in $p_i = 1$, $p_{j \neq i} = 0$ and $R \rightarrow 0$. In contrast, at high temperatures, the migration probabilities become uniform ($p_i = p_j = 1/12$), leading to $R \rightarrow 1$ (Fig. 4d). To quantitatively compare the migration behaviors in PtCo and PtCoCu alloys, we adopted the diffusion multiplicity metric proposed by Cao,⁵⁰ which is defined as the variance of R . Fig. 4e shows the diffusion multiplicity of PtCo and PtCoCu as a function of temperature. It can be observed that the diffusion multiplicity first increases and then decreases with temperature, exhibiting a critical temperature at which $\text{Var}(R)$ reaches its maximum value. In the low-temperature directional migration region, PtCoCu shows a lower $\text{Var}(R)$ than PtCo, indicating that the introduction of Cu enhances directional migration in this temperature range. In the high-temperature random migration region, PtCoCu exhibits a higher $\text{Var}(R)$ value than PtCo, suggesting that Cu reduces random migration at high temperatures. Moreover, the position of the $\text{Var}(R)$ peak for PtCoCu shifts to the right compared to PtCo, indicating stronger directional migration. Additionally, the intensity of the peak increases, revealing greater diffusion heterogeneity. These observations collectively suggest that the introduction of Cu accelerates the ordering kinetics.

Conclusions

In conclusion, by fine-tuning a pretrained large atomic model, we developed a machine learning potential model that accurately predicts the energy of PtCoM ternary alloys with various chemical orderings. This model serves as a powerful tool for

uncovering thermodynamic and kinetic insights into the disorder-to-order transition in PtCoM alloys. Rather than building separate machine learning models for each PtCoM composition, as commonly done in earlier work, we employ a transferable, fine-tuned machine learning potential capable of handling a broad range of elements across the periodic table. This enhanced generality enables efficient screening not only of the 16 elements considered in previous studies but also of most transition metals in the 3rd, 4th, and 5th periods, thereby substantially expanding the discovery space for new alloy candidates.

Thermodynamically, we find that introducing third elements such as Ni, Cu, and Zn increases both the thermodynamic driving force and the critical transition temperature for the disorder-to-order transformation compared to binary PtCo. Our screening results identify several promising PtCoM alloy candidates—PtCoTi, PtCoV, PtCoNi, PtCoCu, PtCoZn, PtCoNb, PtCoAg, PtCoCd, PtCoTa, PtCoAu, and PtCoHg—that exhibit a stronger thermodynamic driving force for ordering than PtCo. These findings highlight the important role of chemical ordering between the third element M and Co in enhancing the overall ordering tendency of PtCoM alloys.

Kinetically, migration barrier calculations reveal that Cu not only possesses a lower migration energy barrier compared to Pt and Co but also facilitates the migration of neighboring atoms. This enhanced atomic mobility promotes structural rearrangement and accelerates the ordering process during annealing. These insights not only deepen our understanding of the underlying atomic mechanisms but also provide valuable guidance for the design of high-performance, cost-effective multicomponent catalysts for energy conversion and storage applications.

Data availability

Data for this article, including the training dataset, the trained machine learning model, the code for Monte Carlo simulations, and the raw data for the figures are available in the GitHub repository at <https://github.com/ZhangLabTHU/disorder-to-order>.

Author contributions

X. N.: data curation, investigation, methodology, visualization, writing – original draft. S. Z., R. Z., and J. L.: project discussion. L. Z.: conceptualization, supervision, methodology, writing – review & editing, project administration and funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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