Composition-dependent ordering transformations in Pt–Fe nanoalloys

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Despite the well-known tendency for many alloys to undergo ordering transformations, the microscopic mechanism of ordering and its dependence on alloy composition remains largely unknown. Using the example of Pt85Fe15 and Pt65Fe35 alloy nanoparticles (NPs), herein we demonstrate the composition-dependent ordering processes on the single-particle level, where the nanoscale size effect allows for close interplay between surface and bulk in controlling the phase evolution. Using in situ electron microscopy observations, we show that the ordering transformation in Pt85Fe15 NPs during vacuum annealing occurs via the surface nucleation and growth of L12-ordered PtFe domains that propagate into the bulk, followed by the self-sacrifice transformation of the surface region of the L12 PtFe into a Pt skin. By contrast, the ordering in Pt65Fe35 NPs proceeds via an interface mechanism by which the rapid formation of an L10 PtFe skin occurs on the NPs and the transformation boundary moves inward along with outward Pt diffusion. Although both the “nucleation and growth” and the “interface” mechanisms result in a core–shell configuration with a thin Pt-rich skin, Pt85Fe15 NPs have an L12 PtFe core, whereas Pt65Fe35 NPs are composed of an L10 PtFe core. Using atomistic modeling, we identify the composition-dependent vacancy-assisted counterdiffusion of Pt and Fe atoms between the surface and core regions in controlling the ordering transformation pathway. This vacancy-assisted diffusion is further demonstrated by oxygen annealing, for which the selective oxidation of Fe results in a large number of Fe vacancies and thereby greatly accelerates the transformation kinetics.

Pt-Fe nanoparticles | alloy composition | in situ electron microscopy | chemical ordering

Many alloys tend to undergo ordering transformations from a random solid solution all the way to fully ordered ground-state intermetallics. The driving force of such an ordering process is the chemical attraction between like and unlike atoms, that is, the occupation of neighboring lattice sites by a different type of atom is energetically more favorable than that by the same type of atom (1). Despite this well-recognized pairwise interaction in driving the ordering transformation (2), dynamically understanding the microscopic processes governing the onset, promotion, and termination of an ordering transformation has rarely been attained, mainly due to the lack of suitable tools capable of spatially and temporally tracking the ordering processes including atom mobility, nucleation and growth of ordered domains and the resultant interface dynamics, and possible formation and structural evolution of metastable, intermediate phases. The situation becomes even more challenging for nanoscale alloys, where the significantly increased surface-area-to-volume ratio not only opens up a variety of additional freedoms to initiate an ordering transformation but also allows for kinetic interplay between the surface and bulk due to their close proximity. We provide direct evidence of the microscopic processes controlling the ordering transformation through the surface–bulk interplay in Pt-Fe nanoalloys and new features rendered by variations in alloy composition and chemical stimuli. These results provide a mechanistic detail of ordering transformation phenomena which are widely relevant to nanoalloys as chemical ordering occurs in most multicomponent materials under suitable environmental bias.

Significance

Dynamically understanding the microscopic processes governing ordering transformations has rarely been attained. The situation becomes even more challenging for nanoscale alloys, where the significantly increased surface-area-to-volume ratio not only opens up a variety of additional freedoms to initiate an ordering transformation but also allows for kinetic interplay between the surface and bulk due to their close proximity. We provide direct evidence of the microscopic processes controlling the ordering transformation through the surface–bulk interplay in Pt–Fe nanoalloys and new features rendered by variations in alloy composition and chemical stimuli. These results provide a mechanistic detail of ordering transformation phenomena which are widely relevant to nanoalloys as chemical ordering occurs in most multicomponent materials under suitable environmental bias.

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The authors declare no competing interest.

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tendency (9, 10), nanostructuring of the alloys not only drives the thermodynamic state far away from the bulk phase but also results in shorter transport pathways, faster diffusion, and more rapid transformation kinetics with respect to their bulk counterparts (11). Second, Pt-based alloy NPs are widely used as cathode electrocatalysts for oxygen reduction reaction (12–15) and anode electrocatalysts for oxygen evolution reaction (16), and structurally ordered intermetallic NPs display superior electrocatalytic activity and durability along with decreased Pt loading compared with Pt/C in electrochemical reactions (17–19). However, as-synthesized Pt-based alloy NPs typically form as random solid solutions that suffer from rapid composition and structure degradation during electrocatalytic processes due to the oxidation and dissolution of the alloy components (17). Therefore, posttreatment processing such as high-temperature annealing has long been employed to transform the random solid solution into an ordered intermetallic phase that has improved electroactivity and durability arising from the stronger Pt–M heteroatomic bonding and a more negative enthalpy of formation compared to its disordered state (18). The ordering can change the electron density of states and alter the position of d-band with respect to the Fermi level, where the position of d-band depends on long-range order of the ordered phase (19–22). Therefore, understanding the ordering transformations in these alloy NPs forms the basis for improving the catalytic efficiency and durability. Our current understanding of the ordering processes in such alloy NP catalysts is largely derived from “quench-and-look” studies, where the catalysts are quenched after a certain amount of annealing time, followed by ex situ characterization of the postannealed NPs (23, 24). However, short-lived intermediate states are difficult to capture with this approach and the ordering mechanism may be more complex than that expected from straightforward postmortem studies. Another issue for postexamination is atmospheric contamination that may bring controversies in understanding the reaction mechanism (25–27). To overcome these shortcomings, in situ tools, including PXRD and TEM, have been used to study the ordering processes during thermal annealing (3–6). By monitoring the relative intensity evolution of the superlattice peaks, PXRD is capable of providing important information on the degree of ordering for an ensemble of Pt alloy NPs. In situ TEM can complement the PXRD by providing dynamic information on the atomic structure and morphological evolution of individual NPs upon the heating. However, to the best of our knowledge, none of the studies have attempted to address the ordering transformation mechanism and correlate it with the alloy composition and the annealing atmosphere.

Herein, we report the real-time, atomic-scale TEM observations of the ordering transformations in Pt–Fe NPs during high-temperature annealing. Two distinct alloy compositions, i.e., Pt85Fe15 and Pt65Fe35, which deviate from the ideal stoichiometry of intermetallic compounds, are chosen to study the ordering mechanism and its dependence on the alloy composition of the NPs. We have performed a comparative study by annealing under vacuum and O2 gas environments, where the former yields the intrinsic ordering behavior for pristine NPs whereas the latter has practical relevance because of the intentional or unavoidable presence of oxidative gas species (O2, H2O, etc.) in the annealing atmosphere for a typical heat-treatment process. By combining with atomistic modeling, this mechanistic study provides microscopic mechanism underlying the onset and progression of the ordering transformation from the surface to the bulk of the NPs and the phase transformation pathways associated with the alloy composition and annealing atmosphere. As compared to pure metal NPs, the complexity is increased manifold for multicomponent NPs, the microscopic insights obtained from the in situ TEM observations open vistas of new opportunities for a synergistic realization of composition and phase via tuning the chemical ordering process through its intrinsic and extrinsic interdependence on the alloy composition and annealing atmosphere.

Results

Pt–Fe alloy NPs were synthesized through a wet chemistry approach (28) (see Materials and Methods and SI Appendix, Figs. S1–S4). Fig. 1A illustrates a representative high-angle annular dark-field (HAADF) image of the as-synthesized Pt85Fe15 NPs, showing the nearly square or rectangular shapes with an average size of ~10 nm. Fig. 1B illustrates an atomically-resolved HAADF-STEM (scanning TEM) image of a typical Pt85Fe15 NP along the [100] zone axis, where the uniform contrast and the absence of superlattice diffraction spots in the diffraction pattern (Fig. 1C, Inset) further evidence the random solid solution nature with a face-centered cubic (fcc) lattice structure. The diffraction spots in the diffractionogram correspond to the (200) reflections and the d-spacing is determined to be ~1.94 Å. Fig. 1D shows the STEM-EDS (energy-dispersive X-ray spectroscopy) line scan across a Pt85Fe15 NP showing the relatively uniform alloy composition without obvious surface segregation of the alloying elements.

Fig. 1 E–G correspond to the HAADF and HRTEM characterization of the as-synthesized Pt85Fe15 NPs, showing features (shape, size, and random solid-solution) similar to Pt85Fe15 NPs. The {200} interplanar spacing measured from the HRTEM image of Pt85Fe15 NPs is ~1.94 Å, slightly smaller than the corresponding value of the {200} planes of Pt85Fe15 NPs. This composition-dependent lattice spacing is further confirmed by synchrotron X-ray diffraction measurements (SI Appendix, Fig. S2 and Table S2), showing that the Pt65Fe35 sample has a slightly smaller d-spacing than that of Pt85Fe15 sample, consistent with Vegard’s law (29, 30). Fig. 1F, Inset is an enlarged view of the surface region of the Pt65Fe35 NP, showing that both the surface region and the bulk display the same fcc lattice of the random solid solution. However, the atomic columns in the outermost two to three atomic layers show much weaker HAADF image intensity than that in the deeper region, which indicates the enrichment of Fe in the surface region, as confirmed by EDS linescan (Fig. 1H) and mapping (SI Appendix, Figs. S3 and S4).

In situ vacuum annealing is performed inside a STEM using a microelectrochemical system (MEMS)-based heating holder. Fig. 2 presents in situ HAADF-STEM images of the ordering-induced atomic structure evolution in a Pt85Fe15 NP, viewed along the [100] zone axis. To avoid any possible effect from the long electron exposure on the ordering process, the beam is on only when acquiring STEM images. As shown in Fig. 2A, the square-shaped NP initially has a solid solution structure with randomly distributed Fe and Pt atoms. After a time elapse of 1 min, nucleation of L12Pt3Fe domains occurs around the surface, as marked by the dashed black lines in Fig. 2B, where the L12 ordering is evidenced by the characteristic arrangement of bright (Pt) and bright/dim (Pt/Fe) atom columns alternate on the {100} planes (more detail in SI Appendix, Fig. S5).
nucleation, growth, and coalescence of L12 Pt3Fe domains propagate into the bulk (Fig. 2 C and D) and eventually transform the NP into a core–shell structure consisting of a thin Pt shell and the L12 Pt3Fe core (Fig. 2 E and F). The existence of the Pt shell is confirmed by EDS maps of the well-annealed NP, showing a larger NP in the Pt map (Fig. 2G) than that in the Fe map (Fig. 2H) and the stronger Pt signal of the outer region in the combined Pt/Fe map (Fig. 2J). The formation of a similar Pt3Fe/Pt core–shell structure is confirmed by examining a large number of well-annealed Pt85Fe15 NPs (SI Appendix, Fig. S6).

Surprisingly, our in situ STEM imaging reveals that the formation of the Pt shell occurs via decomposing the initially formed L12 Pt3Fe domains around the surface during the ordering in the bulk of the NP. This is revealed by the gradual transformation of bright/dim superlattice feature in the surface region (Fig. 2 B and C) into the uniform lattice contrast upon the progressive ordering in the bulk (Fig. 2 E and F). This surface nucleation and growth process of L12 Pt3Fe domains along with the subsequent decomposition of the formed Pt3Fe phase at the surface into the Pt skin provides important clues to the diffusion process for controlling the ordering transformation. For the composition of Pt85Fe15 NPs, the Pt3Fe (L12) ordering requires either the rejection of excess Pt atoms and/or incorporation of additional Fe atoms. The surface region of the NPs has larger atom mobility and is thus kinetically preferred for nucleation and growth of ordered domains by rejecting excess Pt atoms to the outermost surface region along with the outward diffusion of Fe atoms from the bulk to the surface region. As marked by the dashed cyan lines in Fig. 2 A–C, the initially flat surface facet of the NP transforms into a curved shape with the formation and lateral growth of a small protrusion, which can be caused by the surface aggregation of the rejected Pt atoms into a Pt island. The time-sequence HAADF images in Fig. 2 E and F further demonstrate that the islanding of the excess Pt atoms also occurs on the planar surface in the projection view, where the region marked by the red dashed circle shows brighter image contrast and grows larger due to the ordering transformation induced rejection of excess Pt atoms from the bulk to the surface. This is also confirmed by the corresponding EDS mapping, where the region marked by the red dashed circle in the combined Pt and Fe map (Fig. 2J) shows stronger Pt intensity and corresponds to the surface island of Pt atoms.

In addition to the surface rejection of Pt atoms and outward diffusion of Fe atoms from the bulk for the initial stage of the ordering transformation in the surface and subsurface region, our in situ STEM imaging demonstrates that the later-stage ordering transformation in the bulk of the NP involves the inward diffusion of Fe atoms from the surface region into the bulk. As shown in Fig. 2 C–F, the formation of the Pt skin occurs via the progressive transformation of the surface region of the initially formed L12 Pt3Fe into pure Pt. This is due to the counterdiffusion of Fe and Pt atoms, by which the Fe atoms in the surface layer of the Pt3Fe L12 region diffuse inward and the resulting Fe vacant sites are filled by the outward-diffusing Pt atoms from the bulk. The counterdiffusion of Pt and Fe atoms is driven by the composition disparity between the composition Pt85Fe15 in the untransformed bulk region of the NP and the ideal stoichiometry of Pt3Fe intermetallic compound, where the outward diffusion of excess Pt atoms in the bulk in conjunction with the concurrent inward diffusion of Fe atoms from the surface is required to reach the stoichiometry for the formation of L12 Pt3Fe in the core region. This self-sacrifice transformation of L12 Pt3Fe in the surface region into a Pt skin marked by the white dashed circle in Fig. 2 E and F is also favored thermodynamically due to the lower surface energy of Pt than that of L12 Pt3Fe (31–33).

Fig. 2J schematically illustrates the ordering transformation process in Pt85Fe15 NPs. The ordering starts with the nucleation and growth of L12 Pt3Fe domains in the surface region (Fig. 2 J, i). The growth of the ordered domains leads to their coalescence and propagation from the surface region into the bulk (Fig. 2 J, ii). The Pt3Fe L12 ordering in the bulk results in outward diffusion of excess Pt atoms from the core region in conjunction with the concurrent inward diffusion of Fe atoms from the surface region, thereby transforming the surface region of the Pt3Fe L12 phase into a pure Pt skin (Fig. 2 J, iii). The ordering transformation via the counterdiffusion of Pt and Fe atoms results in a core–shell structure consisting of a thin shell of Pt and a Pt3Fe L12 ordered core (Fig. 2 J, iv).

Fig. 3 presents in situ HAADF-STEM image snapshots of the ordering transformation in a Pt85Fe15 NP at 700 °C in vacuum. The as-prepared NPs are of the fcc random solid solution of Fe-Pt, as shown in Fig. 1 and SI Appendix, Fig. S7. Fig. 3A shows an
HAADF image of the NP, captured right after the temperature reaches 700 °C, showing that the entire surface of the NP is already covered with a continuous skin of the ordered intermetallic compound. The intermetallic skin comprises two atomic layers bright (Pt) atom columns in the outermost layer and dim (Fe) atom columns in the second layer, which match the arrangement of Pt and Fe atoms in the $f001$ planes of $L1_0$ PtFe (more detail in SI Appendix, Figs. S7 and S8). The presence of the outermost atomic layer of Pt is also confirmed by the intensity profiles taken across the HAADF image (SI Appendix, Fig. S7) by exploiting the atomic sensitivity of the Z contrast in the HAADF-STEM imaging. As marked by the white dashed lines in Fig. 3 B and C, the transformation boundary between the $L1_0$ PtFe skin and the random solid solution advances inward, which results in $L1_2$ ordering ($Pt_3Fe$) from the subsurface to the bulk. Meanwhile, the initially formed $L1_0$ PtFe skin gradually transforms to $L1_2$ Pt$_3$Fe (Fig. 3D). As a result, the entire NP is dominated by the lattice feature of $L1_2$ Pt$_3$Fe (Fig. 3D). Upon the continued annealing, however, the center region of the NP as marked by the red dashed circle in Fig. 3E further transforms into a different ordered structure in which the arrangement of Pt and Fe atoms as well as the interplanar spacings match well with $L1_0$ PtFe (detail in SI Appendix, Fig. S8). As marked by the red dashed circle in Fig. 3F, the PtFe domain grows outward by consuming the surrounding Pt$_3$Fe. Meanwhile, the superlattice feature of the $L1_2$ Pt$_3$Fe is gradually lost in some of the surface and subsurface regions (as marked by the purple dashed lines in Fig. 3 E and F), indicating the self-sacrifice transformation of the outer region of the $L1_2$ Pt$_3$Fe into a Pt skin in conjunction with the Pt$_3$Fe → PtFe transformation in the bulk and suggesting the tendency to form a PtFe/Pt core–shell structure upon the transformation pathway (more details in SI Appendix, Figs. S8 and S9). This is demonstrated in Fig. 3G, showing the formation of the core–shell structure for a well-annealed Pt$_{65}$Fe$_{35}$ NP that consists of a 1-nm-thick Pt shell and the $L1_0$ PtFe core. As shown in Fig. 3 H–J, the core shell structure is also compositionally confirmed by the corresponding STEM-EDS mapping of the NP. The core/shell boundary position is roughly marked with the white dashed circle in the combined Pt/Fe map (Fig. 3J) and the shell region shows a
weak intensity of Fe signal in the map, indicating the presence of a small amount of Fe in the Pt shell. Similar PtFe/Pt core–shell structure is confirmed by examining a large number of well-annealed Pt65Fe35 NPs, as shown in SI Appendix, Figs. S8 and S9.

The in situ STEM imaging in Fig. 3 shows that the ordering transformation in the Pt65Fe35 NPs starts from the rapid formation of an L10 PtFe skin that further transforms to L12 Pt3Fe. The subsequent inward growth of the L12 Pt3Fe shell toward the Pt65Fe35 solid solution core requires the supply of Pt atoms from the bulk in conjunction with the rejection of excess Fe atoms from the Pt3Fe/Pt65Fe35 transformation boundary toward the core region. This counterdiffusion of Fe and Pt atoms in the course of the inward growth of the L12 Pt3Fe shell results in the transfer of Pt atoms out of the core but Fe atoms into the core region. Therefore, the composition in the core region evolves to the stoichiometric ratio of Pt and Fe, thereby resulting in the nucleation of the L10 PtFe domain in the core region (Fig. 3 K, iii). The outward growth of the PtFe core requires the outward rejection of Pt atoms from the PtFe/Pt3Fe interface to the surface region and inward supply of Fe atoms from the surface region, thereby transforming the NP into a PtFe/Pt core–shell configuration by simultaneously decomposing the initially formed Pt3Fe into PtFe in the core and into Pt in the surface region of the Pt3Fe shell, which eventually leads to the formation of the PtFe/Pt core–shell structure consisting of the Pt shell and the L10 PtFe core, as confirmed by ex situ HAADF imaging and EDS mapping in Fig. 3 G–J and in situ HRTEM imaging (SI Appendix, Fig. S9).

Fig. 3K schematically illustrates the entire process of the ordering transformation in the Pt65Fe35 NPs. The ordering starts with the rapid formation of an L10 PtFe skin on the parent NP (Fig. 3 K, i). This is followed by the transformation of L10 PtFe skin to L12 Pt3Fe and its inward growth via the outward diffusion of Pt atoms from the Pt65Fe35 core to the Pt3Fe/Pt65Fe35 interface and inward diffusion of Fe atoms from the interface to the core (Fig. 3 K, ii). The counterdiffusion of Pt and Fe makes the composition in the core region evolve into the near-equal atomic ratio of Pt and Fe, thereby resulting in the nucleation of the L10 PtFe domain in the core region (Fig. 3 K, iii). The outward growth of the L10 PtFe core requires the outward rejection of Pt atoms from the PtFe/Pt3Fe interface to the surface region and inward supply of Fe atoms from the surface region, thereby transforming the NP into a PtFe/Pt core–shell configuration by simultaneously decomposing the initially formed Pt3Fe into PtFe in the core and into Pt in the surface region (Fig. 3 K, iv).

The in situ STEM imaging of the NPs shown above is performed under the vacuum annealing condition that results in the removal of the O-rich organic capping layer and oxygen on the NPs (more detail in SI Appendix, section SI-4), thereby rendering the intrinsic behavior of ordering transformations for...
FeOx shell and a Pt-rich random solid solution core. The gray, purple and red spheres represent the Pt-rich solid solution, Fe and O atoms,

$pO_2^{NP}$ during O2 annealing. (Fig. 4.)

As a consequence of the annealing, the O2 ambient is further con

strong resistance to the ordering transformation in the NPs in

driven tilting of the NP at the elevated temperature, which

as a FeOx skin. Consistent with the EDS mapping, the corre-

around the NP surface indicates that Fe is selectively oxidized

ation. As a result, the O2 annealing transforms the Pt85Fe15 NP

structural evolution from the outermost surface layer to deeper

ubiquitous presence of O2 gas (more or less) in the annealing

oxygen on ordering transformations in the NPs because of the

clean NPs. We further examine the possible effect of ambient

oxygen on ordering transformations in the NPs because of the

ubiquitous presence of O2 gas (more or less) in the annealing

atmosphere for a practical heat-treatment process. Our in situ

experiments are performed within the environment TEM that

allows for temperature-, pressure-, and time-resolved observa-

tions of the structural transformation in the NPs by flowing O2

gas in the sample area while simultaneously monitoring atomic

structural evolution from the outermost surface layer to deeper

atomic layers of the NP. Fig. 4 A–C show in situ HRTEM images of a Pt85Fe15 NP (viewed along the [001] zone axis) during the annealing at 700°C in the O2 flow of $5 \times 10^{-3}$ Torr. The NP maintains the random solid solution without showing any noticeable ordering transformation throughout the annealing period (the slight loss of the two-dimensional lattice fringe image contrast in Fig. 4 C is caused by thermally driven tilting of the NP at the elevated temperature, which results in the slight deviation from the [001] zone axis). This strong resistance to the ordering transformation in the NPs in the O2 ambient is further confirmed by prolonged annealing. Fig. 4D illustrates a STEM-HAADF image of a Pt85Fe15 NP annealed at 700°C in the O2 flow of $5 \times 10^{-3}$ Torr for 2 h, where the annealing time is more than twice of that shown in Fig. 2. The HAADF image shows the uniform image contrast of atom columns, indicating the random distribution of Pt and Fe atoms without undergoing chemical ordering (more examples given in SI Appendix, Fig. S10). Fig. 4E presents the STEM-EDS elemental mapping of a well-annealed NP in the O2 ambient, in which the stronger intensity of Fe and O signal around the NP surface indicates that Fe is selectively oxidized as a FeOx skin. Consistent with the EDS mapping, the corresponding electron energy-loss spectroscopy maps (see SI Appendix, Fig. S11) further confirm the selective oxidation of Fe from the O2 annealing and resultant elemental redistribution. As a result, the O2 annealing transforms the Pt85Fe15 NP into a core–shell structure consisting of an ultrathin FeOx shell and a Pt-rich solid solution core. The selective oxidation of the minor element (Fe) results in the depletion of Fe in the alloy; therefore, the solid solution in the core becomes so dilute that it is effectively the pure component of Pt. Fig. 4F schematically illustrates the Pt–FeOx core–shell structure of the NP resulting from the O2 annealing.

Fig. 5 A–E display time-sequence HRTEM images along the [001] zone axis, showing the structural evolution of a Pt65Fe35 NP during the annealing at 700°C in pO2 = $5 \times 10^{-3}$ Torr. As marked by the dashed cyan circles in Fig. 5B, the superlattice domains become visible, signaling the onset of the ordering transformation via the nucleation and growth of L12 Pt3Fe domains in the initially homogeneous solid solution NP. Meanwhile, an ultrathin, amorphous-like FeOx skin develops around the NP surface (marked by the red dashed lines in Fig. 5B) due to the selective oxidation of Fe. Upon the continued O2 annealing, the L12 Pt3Fe domains expand and result in the formation of a L12 Pt3Fe core. The selective oxidation of Fe is further evidenced by the expansion of the L12 Pt3Fe domains and the thickening of the amorphous-like FeOx shell around the NP, as marked by the dashed cyan and red circles, respectively, in Fig. 5E. As can be seen in Fig. 5E, an intermediate layer of the Pt-rich random solid solution exists between the inner L12 Pt3Fe core and the outer FeOx shell. The intermediate Pt-rich layer is formed due to the selective oxidation of Fe to form the FeOx shell. Fig. 5F is a diffractogram of the HRTEM image in Fig. 5E, in which the sharp (100) and (110) superlattice reflections confirm the high degree of ordering in the L12 Pt3Fe core. Fig. 5G presents a STEM-HAADF image of a well-annealed Pt65Fe35 NP (2 h at 700°C in pO2 = $5 \times 10^{-3}$ Torr), showing the dominant formation of the L12 Pt3Fe core along with the Pt-rich solid solution in the some of the subsurface region. The FeOx formation at the outer surface is also visible in some region because of its large thickness, where the absence of crystalline lattice contrast in the HAADF image confirms the amorphous nature of the oxide. Fig. 5 E and G also show that the FeOx clusters formed in the O2 annealing tends to spread away from the Pt65Fe35 NPs (more examples in SI Appendix, Fig. S12), a behavior similar to the oxidation of Pt0.5Co0.5 NPs (34). Fig. 5H displays the corresponding STEM-EDS mapping, in which the strong intensity of O and Fe in the surface region corresponds to the FeOx shell and the stronger intensity of Pt than that of Fe in the bulk region is related to the L12 Pt3Fe intermetallic compound. The intermediate layer of the Pt-rich solid solution is not distinguishable in the EDS mapping due to its relatively low spatial resolution. More examples of the EDS mapping of the O2 annealed Pt65Fe35 NPs are given in SI Appendix, Fig. S13, showing large Fe–O regions composed of Fe–O flocs and newly formed FeOx clusters from the O2 annealing. Fig. 5I schematically illustrates the resulting sandwich structure consisting of the L12 Pt3Fe core, the intermediate layer of the Pt-rich solid solution, and the FeOx shell from the O2 annealing of Pt65Fe35 NPs.

Discussion

The as-synthesized Pt–Fe NPs are nearly square or rectangle shapes, dominated by (100) facets (Fig. 1). During the annealing process, the sharp corner regions evolve into the round shape driven by surface diffusion of atoms and have the tendency to further evolve into relatively flat (110) facets (i.e., round and truncated shapes), as shown in Figs. 2 and 3 and SI Appendix, Figs. S5–S9. Such annealing-induced shape changes to the NPs are consistent with other studies (3, 35, 36). More importantly, the in situ STEM imaging (Figs. 2 and 3) demonstrates the composition-dependent ordering transformations in Pt–Fe NPs. That is, the ordering transformation in Pt85Fe15
The surface nucleation and growth of L12 PtFe atoms from the bulk toward the surface and short-range diffusion of Fe and Pt atoms without the need for a long-range supply of Fe from the bulk. The Pt surface termination is driven by the lower surface energy of Pt than Fe ($\gamma_{Pt} \sim 2.79$ J/m$^2$, $\gamma_{Fe} \sim 3.55$ J/m$^2$) (39–41). The surface segregation of Pt atoms from the bulk is accompanied by the inward migration of Fe atoms, which therefore subsequently transforms the L1$_0$ PtFe shell into the Pt-richer L1$_2$ Pt$_3$Fe.

The following inward advancement of the L1$_2$ Pt$_3$Fe involves the outward diffusion of Pt atoms from the core to the Pt$_3$Fe/Pt$_6$Fe$_{35}$ interface and inward diffusion of excess Fe atoms from the moving Pt$_5$Fe/Pt$_6$Fe$_{35}$ interface toward the core region. This counterdiffusion of Fe and Pt drives the alloy composition in the core region to evolve to the stoichiometric ratio of intermetallic PtFe, greatly facilitating the nucleation of the L1$_0$ PtFe in the core region. The nucleated L1$_0$ PtFe domain grows outward by rejecting excess Pt atoms from the PtFe/Pt$_3$Fe interface toward the surface region and the inward supply of Fe atoms from the formed Pt$_3$Fe in the surface region, thereby resulting in the Pt surface shell by sacrificing the existing PtFe and eventually leading to the formation of the PtFe/Pt core-shell structure with the L1$_0$ PtFe core and the Pt shell, as shown in Fig. 3. For Pt$_6$Fe$_{35}$ NPs, this Pt$_3$Fe $\rightarrow$ PtFe transformation is driven by the strong Pt–Fe pairwise interactions and should be thermodynamically favorable because it leads to the maximized Pt–Fe bonds. By comparison, Pt$_6$Fe$_{35}$ NPs have a lower overall content of Fe and do not show the Pt$_3$Fe $\rightarrow$ PtFe transformation in the final configuration. This is because the L1$_2$ Pt$_3$Fe phase is thermodynamically more favorable than PtFe since the former allows for more abundant pairwise Pt–Fe atomic interactions for the relatively dilute Fe content in the Pt$_6$Fe$_{35}$ NPs.

To gain further understanding of the diffusion processes underlying the composition-dependent ordering transformations, we carried out density functional theory (DFT) calculations to predict the diffusion energetics in the two random Pt–Fe alloys. We used periodic supercells containing 108 atoms to model the Pt–Fe random alloys with the specific compositions of Pt$_{65}$Fe$_{35}$ and Pt$_{65}$Fe$_{35}$. In both cases, special quasi-random structures were generated using the ATAT code (42) to distribute Fe and Pt atoms randomly in an fcc lattice. Our DFT calculation predicts the equilibrium lattice parameters of an fcc unit cell to be 3.91 Å for Pt$_{65}$Fe$_{35}$ and 3.88 Å for Pt$_{65}$Fe$_{35}$, which are in good agreement with Vegard’s law (29, 30). Moreover, we predict the vacancy formation energies and migration energies for vacancy-assisted diffusion in the Pt–Fe random alloys. The vacancy formation energies are calculated as the energy difference between the system containing a vacancy and the perfect crystal, taking into account the chemical potential change associated with the vacancy formation. In this work, the chemical potential of a Fe or Pt atom is adopted as the energy per atom in the pure body-centered cubic (bcc) Fe or fcc Pt crystal. It should be noted that the vacancies in the random alloys have different local chemical environments and hence different formation energies. Consequently, we calculate the formation energies for each type of the vacancies at four different chemical environments in terms of the varying local Fe concentration. The predicted vacancy formation energies are depicted in Fig. 6A for Pt$_{65}$Fe$_{35}$ and Fig. 6C for Pt$_{65}$Fe$_{35}$ alloys. We predict the average value of the formation energies to be 0.39 eV for a Fe vacancy and 1.26 eV for a Pt vacancy in Pt$_{65}$Fe$_{35}$ alloy, suggesting that Fe vacancies are favored to form in this alloy than Pt vacancies. In contrast, the average values of vacancy formation energies are found to be 1.84 eV for a Fe vacancy and 1.62 eV for a Pt vacancy in Pt$_{65}$Fe$_{35}$. It is worth mentioning that these predictions are close to the reported vacancy formation energies of 1.85 eV for a Fe vacancy and

**Fig. 5.** In situ TEM imaging of Pt$_{65}$Fe$_{35}$ NPs during annealing at 700°C in pO$_2$ = 5 x 10$^{-3}$ Torr. (A–E) Time-sequence HRTEM images, showing that the selective oxidation of Fe along with the ordering transformation in the bulk of the NP during O$_2$ annealing results in a sandwich structure consisting of an amorphous-like FeO$_x$ shell, a thin intermediate layer of Pt, and the L1$_2$ Pt$_3$Fe core. (F) Diffractogram of the HRTEM image in E, showing the (100) and (110) superlattice reflections associated with the L1$_2$ Pt$_3$Fe core. (G) HAADF-STEM image showing the presence of the L1$_2$ Pt$_3$Fe core in the NP annealed for 2 h at 700°C in pO$_2$ = 5 x 10$^{-3}$ Torr. The regions marked with the cyan dashed circles in B, E, and G correspond to the L1$_2$ Pt$_3$Fe structure. The red dashed circles mark the FeO$_x$/Pt boundary. (H) STEM-EDS mapping of an O$_2$-annealed NP, showing the presence of a FeO$_x$ shell. (I) Schematic illustration of the FeO$_x$/Pt/Pt$_3$Fe sandwich structure of the NP from the O$_2$ annealing. The yellow, purple, and red spheres represent Pt, Fe, and O atoms, respectively.
1.70 eV for a Pt vacancy in Fe50Pt50 predicted using the coherent potential approximation (43, 44).

The observed Pt segregation at the NP surface is believed to be a result of a directional migration of the Fe or Pt atoms through the vacancy-exchange mechanism in the Pt–Fe NPs. Hence, we calculate the migration energies for the vacancy-assisted diffusion of Fe and Pt atoms in the two Pt–Fe alloys using the nudged elastic band (NEB) method (45). In this work, the migration path is defined as one atom moving to an adjacent vacancy. There are four types of diffusion paths, namely, a Pt atom to a Pt vacancy, an Fe atom to a Pt vacancy, a Pt atom to a Fe vacancy, and an Fe atom to a vacancy, in the alloys. Our DFT results of the migration energy of Fe and Pt atoms are illustrated in Fig. 6B for Pt65Fe35 and Fig. 6D for Pt65Fe35 alloy. In Pt65Fe35 alloy, we predict that the migration energy of a Pt atom to an adjacent Pt vacancy could be as high as 4.36 eV, implying a very difficult movement of Pt atoms along this path. In addition, the predicted migration energy barrier of a Pt atom to a Fe vacancy (2.55 eV) is higher than those for a Fe atom to either a Pt vacancy (1.37 eV) or a Fe vacancy (2.28 eV). Consequently, our DFT results indicate that Fe is the faster-migrating species in Pt65Fe35. This corroborates well with our experimental observations showing that the ordering transformation in Pt65Fe35 NPs is initiated by the nucleation and growth of L12 Pt3Fe domains at the surface, which requires the outward diffusion of Fe atoms from the bulk to the Pt-rich surface (Figs. 1D and 2A–D). The later-stage ordering transformation in the core region of the NP requires the inward diffusion of Fe atoms from the surface region into the core (Fig. 2E and F).

In Pt65Fe35, our DFT calculations show that the formation energy of a Pt vacancy is lower than that of a Fe vacancy. However, the migration energy of a Fe atom to an adjacent Fe vacancy is lower (0.95 eV) than the barriers for the migration of a Fe or Pt atom to a Pt vacancy (1.42 to 1.62 eV) (Fig. 6D). In the initial stage of the ordering transformation, the rapid formation of the L10 PtFe skin (Fig. 3A) can be related to the small energy barrier for the inward migration of Fe atoms from the Fe-rich surface region. For the subsequent growth of Pt3Fe and PtFe, the diffusion of Fe and Pt atoms can be equally important. This is because the smaller vacancy formation energy of Pt than that of Fe favors the generation of more Pt vacancies than Fe vacancies (thereby facilitating the diffusion of Pt and Fe atoms to Pt vacancies), whereas the lower energy barrier for the Fe migration of Fe vacancies promotes Fe diffusion.

As discussed above, the ordering transformations are controlled by the vacancy-assisted diffusion of Pt and Fe atoms. Therefore, the availability of atomic vacancies in the alloys plays an important role in influencing the ordering kinetics. This effect is particularly clear by comparing the ordering transformation in Pt65Fe35 NPs for annealing in vacuum and O2 ambient. As shown in Fig. 3A–D, it takes ~60 min to fully transform the Pt65Fe35 NP into L12 Pt3Fe for vacuum annealing at 700 °C. By contrast, the full transformation of the Pt65Fe35 NP into L12 Pt3Fe takes only ~2 min for the O2 annealing at 700 °C. This significantly enhanced ordering kinetics in the O2 ambient can be related to the selective oxidation of Fe to form FeOx, which results in a large number of Fe vacancies in Pt65Fe35. As shown from our DFT calculations, the energy barrier for the migration of a Fe atom to a neighboring Fe vacancy is 0.95 eV, which is lower than all other diffusion paths (Fig. 6D) for Pt65Fe35. Therefore, the formation of a large number of Fe vacancies from the selective oxidation of Fe significantly promotes the diffusion of Fe in the
alloy, thereby dramatically accelerating ordering kinetics. By contrast, no ordering transformation is observed for the O2 annealing of Pt85Fe15 NPs (Fig. 4). On one hand, the selective oxidation of Fe results in more Fe vacancies in the alloy and therefore enhances the mobility of atoms for potential ordering transformation. On the other hand, the selective oxidation of Fe also depletes Fe from the Pt85Fe15 alloy that is already Fe-poor, and therefore impairs the ordering because of the lack of sufficient Fe to form a Pt–Fe intermetallic compound.

In summary, we have used in situ TEM to directly monitor the ordering transformations in Pt65Fe35 and Pt85Fe15 NPs at the atomic scale. Our in situ atomic-scale observations demonstrate that the ordering in Pt85Fe15 NPs occurs via the surface nucleation and growth of L12 Pt3Fe domains by long-range diffusion of Fe atoms from the bulk to the surface region. The ordering propagates inward with continued Pt3Fe growth in the bulk and self-sacrifice transformation of the surface region of the formed L12 Pt3Fe into a Pt skin, resulting in a core–shell structure consisting of the L12 Pt3Fe core and the Pt shell. By contrast, the ordering in Pt65Fe35 NPs starts with the rapid formation of an L10 PtFe skin via short-range diffusion of Fe and Pt atoms in the surface region. The subsequent inward growth of the L12 Pt3Fe shell results in the Fe enrichment in the core and thus drives the nucleation of L10 PtFe in the core. The outward growth of the L10 PtFe core is accompanied by the decomposition of the surface region of the formed L12 Pt3Fe into a Pt skin, leading to a final core–shell structure consisting of the L10 PtFe core and a Pt shell. Our DFT calculations identify the composition-dependent vacancy-assisted counterdiffusion of Pt and Fe atoms between the surface and core regions in controlling the ordering transformation pathway. This vacancy-assisted diffusion is further demonstrated by annealing in O2 ambient, for which the selective oxidation of Fe results in a large number of Fe vacancies in the alloy, thereby greatly accelerating the ordering kinetics that transforms the Pt65Fe35 NPs into a sandwich structure consisting of a FeOx shell, an intermediate layer of Pt and an L12 Pt3Fe core. By contrast, the O2 annealing transforms Pt85Fe15 NPs into a core–shell structure composed of a FeOx shell and Pt-rich solid solution core that is depleted with Fe for the selective oxidation of Fe in the alloy. These results provide unprecedented microscopic insights into the ordering transformation phenomena and their dependence on the alloy composition and chemical stimuli. Structurally ordered nanoalloys constitute an important subclass of metals and alloys and have attracted wide attention from the perspective of both fundamental understanding and potential applications, the results reported here are of wide relevance and considerable practical importance for a wide range of alloys, properties, and reactions, including metallurgy, nanostructure synthesis, corrosion, oxidation, and heterogeneous catalysis.

Materials and Methods

Synthesis of Random Solid Solutions of Pt–Fe NPs. Pt–Fe NPs were prepared using a modified method reported previously (28) and more detail is given in SI Appendix, sections SI 1–3. Composition analysis of the as-synthesized NPs was performed using an Optima 7000 DV ICP-OES (inductively coupled plasma optical emission spectrometry) spectrometer, EDS, and synchrotron X-ray diffraction measurements on the lattice spacings. In addition to the EDS linescan analysis performed on individual NPs (Fig. 1 D and H), EDS mapping of a collection of as-synthesized NPs (SI Appendix, Fig. S3) further confirms the more Fe enrichment in the surface region of Pt65Fe35 NPs than in Pt85Fe15 NPs. The EDS mapping also shows that the as-synthesized Pt85Fe15 and Pt65Fe35 NPs are covered with an O-rich organic capping layer. This capping layer can be largely removed by vacuum annealing (SI Appendix, Fig. S4), consistent with other studies (37, 38). The as-synthesized Pt65Fe35 NPs have a small amount of Fe–O flocules, as confirmed by XRD (SI Appendix, Fig. S2), HAADF and EDS mapping (SI Appendix, Fig. S4), and ICP-OES (SI Appendix, Table S1). These Fe–O flocules remain stable without decomposition under the vacuum annealing (SI Appendix, Fig. S4). By contrast, the as-synthesized Pt85Fe15 NPs are free of Fe–O flocules.

In Situ STEM and TEM Characterization. The as-prepared Pt–Fe NPs were loaded on SiN targets and then transferred into the TEM for STEM and TEM observations.

DFT Calculations. The DFT calculations were performed using the Vienna ab initio simulation package (VASP) (46). The projector augmented wave pseudopotential (47) was used to describe the core electrons of the elements. The energy cutoff was set as 500 eV to expand wave function. The generalized gradient approximation with the Perdew–Burke–Ernzerhof (48) exchange-correlation functional was used to evaluate the exchange-correlation energy. In all calculations, the total energy of system was converged within 10−6 eV. A Monkhorst-Pack k-point grid of 4 × 4 × 4 was used in these calculations. All the structures were fully relaxed until the force acting on each atom was lower than 0.01 eV Å−1. The transition states of vacancy migration were located using the NEB method (46), in which the forces along and perpendicular to the path were relaxed to less than 0.05 eV Å−1. In each NEB calculation, we used three intermediate images to locate the transition state and calculated the migration energy as the energy difference between the transition state and the initial state.

Data Availability. All study data are included in the article and/or SI Appendix.

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