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Coalescence of AuPd nanoalloys in implicit environments

We develop numerical tools to model the formation processes of metallic and bimetallic nanoparticles. This work investigates the synthesis of AuPd nanoalloys surrounded by various environments. We find that the kinetics of the chemical ordering of the nanoalloy are significantly affected by how strongly the two chemical species interact with the environment itself. The present work provides a foundation for understanding the chemical ordering which occurs during the formation of nanoalloys and hence for the development of nanodevices with a controlled surface chemical distribution.

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Coalescence of AuPd nanoalloys in implicit environments†

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The optimal design of nanoparticles and nanoalloys arises from the control of their morphology which depends on the synthesis process they undergo. Coalescence is widely accepted as one of the most common synthetic mechanisms, and it occurs both in the liquid and gas phases. Coalescence is when two existing seeds collide and aggregate into a larger object. The resulting aggregate is expected to be far from the equilibrium isomer, *i.e.* the global minimum of the potential energy surface. While the coalescence of nanoparticles is well studied in a vacuum, sparse computational studies are available for the coalescence in an environment. Using molecular dynamics simulations, we study the coalescence of Au and Pd nanoseeds surrounded by an interacting environment. Comparing the initial stages of the coalescence in a vacuum and the presence of an interacting environment, we show that the formation kinetics strongly depends on the environment and on the size of the nanoalloy. Furthermore, we show that it is possible to tune the resulting nanoalloys' surface chemical composition by changing their surrounding environment.

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1 Introduction

Since Granqvist's work and the Japanese project Erato,¹ interest in nanoparticles (NPs) has remained high as they have several technological applications. To cite a few, they are the building blocks of sensors, energy harvesting devices, catalysts, and randomly assembled neuromorphic systems.^{2–9} NPs are discrete objects with three dimensions below 100 nm, showing peculiar properties depending on their morphology. Among NPs, metallic nanoalloys (NAs) – referring to NPs made of two or more metals – possess unique chemophysical features.^{10–13} We define NP morphology as the combination of NA size, shape, chemical composition, and chemical ordering of the inner and surface layers.^{14,15} At the core of a rational design process, we must understand how NAs form, evolve, and agglomerate. Numerical methods can shed light on the atomistic processes and the kinetics of the formation/agglomeration processes, providing complementary information to experimental techniques.^{16–18} Generally speaking, the formation of NPs and NAs follows three possible paths: (i) by the subsequent deposition of atoms over an existing core – the so-called one-by-one growth;¹⁹ (ii) by coalescence – the sintering of two or more individual seeds; and (iii) annealing of a melted seed with a specific rate.²⁰ During the coalescence and one-by-one growth

processes, surface rearrangements are mainly driven by adatom diffusion and the formation of small islands. Annealing might be driven by pressure effects and the minimisation of the surface energy contribution. Among the three growth modes, coalescence plays a non-negligible role in gas aggregation cluster sources and liquid laser ablation. The latter is acquiring increasing importance in the large-scale production of Au-based NAs.²¹ In the modelling of coalescence, we appreciate considerable efforts in understanding the coalescence of Au-based NAs.^{22–25} Nonetheless, to our knowledge, those studies refer to coalescence in a perfect vacuum, although the surroundings may likely affect both NA stability, *i.e.*, surface energy, and kinetics, *i.e.*, adatom diffusion. Our interest is in understanding the effect of the environment around NAs during their formation process. We focus on Au–Pd NAs because of their role in heterogeneous thermal catalysis and electrocatalysis, as they exhibit superior activity and enhanced selectivity.^{26–29} The rationale for combining Au with Pd is the plasmonic and versatile character of the Au-seed with the strong catalytic flavour of Pd. Nano Au–Pd is employed in industrial processes such as hydrogen peroxide synthesis from H₂ and O₂, alcohol oxidation,³⁰ vinyl acetate monomer synthesis,³¹ and formic acid de-hydrogenation.³²

Several studies report the complex energy landscape of Au–Pd.^{33,34} They show a tendency to form Au-rich surfaces, with a ball (Pd)–cup (Au) chemical ordering predicted at sizes up to 3 nm.³⁵ We investigate and compare the kinetics of the coalescence of gold and palladium seeds over a few tens of ns both in a vacuum and in a plethora of environments. We use the Cortes-Huerto formalism³⁶ to mimic a uniform medium around the NA, and we can tune the type and the strength of

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† Electronic supplementary information (ESI) available: MEIs selected and supplementary characterization methods: PDDF in sapphire, Au₅₅th–Pd₅₅th, and Au₅₆₁th–Pd₅₆₁th. See DOI: <https://doi.org/10.1039/d4cp00916a>



the interaction for each chemical species. We reveal that the surface chemical composition and the sintering process time-scale depend significantly on the environment.

2 Methodology

We study the first steps of the coalescence of a 55-atom gold icosahedron (Ih), Au₅₅^{Ih}, against a 55-atom Ih of palladium, Pd₅₅^{Ih}, leading to AuPd₁₁₀. We, then, consider the coalescence of Au₅₆₁^{Ih} against a Pd₅₆₁^{Ih}. In Fig. 1, we report a pictorial representation of the coalescence process.

To understand the effect of the medium, we vary the strength and the type of interaction between metal atoms and the environment (metal–environment interaction, MEI).

We keep the chemical composition equal to 50% and select the same initial geometrical shape, namely an icosahedron, for both seeds. The rationale behind our choice is that often noble and quasi-noble clusters in the 1–3 nm size-range shape as an Ih, which is the closest structure to a sphere.¹⁶

We perform classical molecular dynamics simulations using the open-source package LoDis.³⁷ The systems are thermalised using an Andersen thermostat with a frequency of 10¹¹ Hz. We choose 400 K for the coalescence of Au₅^{Ih}–Pd₅₅^{Ih} and 600 K for Au₅₆₁^{Ih}–Pd₅₆₁^{Ih}. At 600 K, the gold region in any AuPd₁₁₀ almost melts, leading to more mixed and structurally disordered NA independent of the metal–environment interaction. As we want to address the effect of the environment before melting occurs, we do not comment on this case further.

Because we are interested in the first stages of the formation process, the trajectories are collected over $t^* \in (10:20)$ ns, sampling the geometrical configuration every 0.01 ns. Preliminary simulations over 100 ns show that major structural changes occur in the first 20 ns. We average our results over four independent simulations. The Hamiltonian of our system is the sum of two contributions: the metal–metal interaction E_i^{M-M} and the metal–environment E_i^{M-E} , per atom i

$$E_{\text{tot}} = \sum_i E_i = \sum_i E_i^{M-M} + E_i^{M-E} \quad (1)$$

The metal–metal interaction is modelled according to the second moment approximation (SMA) in the tight-binding (TB)

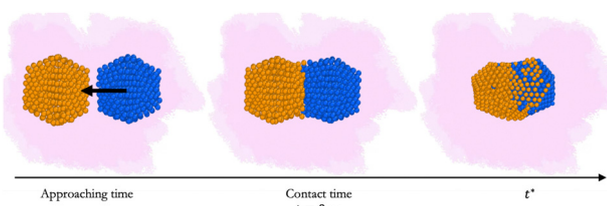


Fig. 1 Pictorial representation of the coalescence process *versus* time, where the pink shadows represent the implicit interacting environment. Orange and blue represent Au and Pd atoms of two NPs, respectively. Along the time axis, we define the approaching time, when the two seeds move towards each other; the contact time, the instant when the two seeds come in contact, set at 0 ns; as the evolution continues, we label with t^* a generic instant after the contact time.

Table 1 TB-SMA parameters used for AuPd nanoalloys from ref. 42

Chemical	p	q	A [eV]	ζ [eV]
Au–Au	10.139	4.033	0.2095	1.8153
Pd–Pd	11.0	3.794	0.1715	1.7019
Au–Pd	10.543	3.8862	0.1897	1.7536

model,³⁸ which is widely used to mimic the growth of NPs and NAs.^{39–41} In such a framework, the total energy of each atom i contains an attractive many-body term and a 2-body repulsive Born–Mayer term as

$$E_i^{M-M} = \sum_{j \neq i}^{n_v} A_{\alpha,\beta} e^{-p_{\alpha,\beta} \left(\frac{r_{ij}}{r_{\alpha,\beta}^0} - 1 \right)} - \sqrt{\sum_{j \neq i}^{n_v} \zeta_{\alpha,\beta}^2 e^{-2q_{\alpha,\beta} \left(\frac{r_{ij}}{r_{\alpha,\beta}^0} - 1 \right)}},$$

where r_{ij} is the atomic pair distances; $r_{\alpha,\beta}^0$ is the reference lattice parameter of individual chemical species α, β , or their arithmetic mean for hetero-pairs; and n_v is the number of neighbours within a radius between the bulk second and third neighbour distances. The parameters $A_{\alpha,\beta}$, $\zeta_{\alpha,\beta}$, $p_{\alpha,\beta}$, and $q_{\alpha,\beta}$, see Table 1, are fitted to reproduce the experimental values of the cohesive energy, elastic constant, lattice parameter and bulk modulus.

We model the metal–environment interaction (MEI), E_i^{M-E} , via the CortesHuerto, Goniakowski, Noguera formalism^{36,43,44} that approximates the presence of a non-inert, homogeneous environment, in terms of the number of unsaturated bonds of surface atoms i , concerning their positioning in bulk $CN_{\text{bulk}} = 12$. The contribution to the atom i of chemical type α is represented by $E_{i \in \alpha}^{M-E}$,

$$E_{i \in \alpha}^{M-E} = -\eta_{\alpha} (CN_{\text{bulk}} - CN_{i \in \alpha})^{\rho_{\alpha}}, \quad (2)$$

CN_i being the coordination of atom i of type α . Eqn (2) is an improvement with respect to implicit electrostatic potential as it accounts for the site-specific adsorption and hence changes the relative stability of facets relevant during the coalescence process. As in previous works,^{36,43} we fix the solvation number to 7 and eqn (2) stands when the number of unsaturated bonds exceeds the difference between the bulk coordination and the solvation number. We should stress that in the present work, the $CN_{i \in \alpha}$ is an analytic function of the distance of i and its neighbours, as proposed in ref. 20 for metadynamics studies.

The strength of the interaction is tuned by the parameter η_{α} while ρ_{α} controls the type of interaction. For example, $\rho_{\alpha} = 1$ stands for a pairwise interaction; while covalent interaction occurs when $\rho_{\alpha} < 1$ and $\rho_{\alpha} > 1$ is for strongly-interacting environments.^{36,44}

For AuPd NAs, we have four parameters ρ_{Au} , ρ_{Pd} , η_{Au} , η_{Pd} to vary. The choice of η_{α} and ρ_{α} has been made by exploring different types of interactions. We identify 37 MEIs listed in Table S1 of the ESI.† Several of them identify poorly interacting environments and have a negligible effect on the kinetics of the coalescence process. Environments interacting with only one metal have a significant effect on both the nanoalloy energy and the kinetics of the coalescence process. When $\rho_{\alpha} = 1.5$ and η_{α} exceeds 0.04 eV per atom, the interaction could approximate the presence of strongly interacting capping agents, as CTAB on Au or C₂-polyol on Pd. Nonetheless, we



should note that eqn (2) is a mean-field approximation without reproducing any specific environments.⁴⁴ The choice for η_α and ρ_α could alter the relative stability of Miller-index surfaces, as it favours low coordinated Pd (Au) atoms. An estimate of the surface energy of a generic plane p , γ_p , can be approximated as $\gamma_p \sim k_\alpha (\text{CB}_{\text{bulk}} - \text{CN}_p) \left[1 - \frac{\eta_\alpha}{k_\alpha} (\text{CB}_{\text{bulk}} - \text{CN}_p)^{(\rho-1)} \right]$, with k_α the energy per surface atom of a certain material. For ρ_α below or equal 1, the surface energy order is still (111) < (100) < (110). In any event, η_α below 0.04 eV per atom preserves the same order. Apart from the vacuum (set 0), we split MEIs into three groups.

- MEI-A: Only Au atoms interact with the environment, setting $\eta_{\text{Pd}} = 0$ eV. ρ_{Au} is 0.5, 1.0, 1.5 and η_{Au} between 0.02 and 0.08 eV per atom, respectively.

- MEI-P: Only Pd atoms interact with the environment, $\eta_{\text{Au}} = 0$ eV. ρ_{Pd} is 0.5, 1.0, 1.5 and η_{Pd} equal to 0.02 eV per atom, 0.08 eV; $\eta_{\text{Pd}} = 0.06$ eV with $\rho_{\text{Pd}} = 1.5$.

- MEI-AP: Both Au and Pd atoms interact. We fix the Au values to (0.02 eV per atom, 0.5). Pd-E interactions vary with ρ_{Pd} equal to 0.5, 0.75, 1.0, and 1.5 and $\eta_{\text{Pd}} = 0.02, 0.04, 0.06, 0.08,$ and 0.2 eV per atom.

For the coalescence of $\text{Au}_{561}^{\text{lh}}-\text{Pd}_{561}^{\text{lh}}$ we select fewer MEIs among group P and AP: (i) P-group: $\rho_{\text{Pd}} = 1.5$ and $\eta_{\text{Pd}} = 0.08$ eV per atom; (ii) AP: ρ_{Pd} is 0.5, 1.0, 1.5 and $\eta_{\text{Pd}} = 0.02$ eV, and 0.08 eV per atom. Such reduction is motivated by the results obtained on the 110-atom system and to save computational time. In the main text, we discuss four cases: the vacuum, MEI-AP, MEI-P, and MEI-A. All the results from other environments are available in the ESI.†

2.1 Characterisation tools

We analyse and characterise the classical MD trajectories using the Sapphire suite.⁴⁵ We select a few descriptors to quantify the structural and chemical changes during the AuPd coalescence. Our aim is to show the effect of the environment on the kinetics of the coalescence and the surface chemical ordering. We focus on the evolution of the neck region, *i.e.*, the spatial region where coalescence occurs, the MEI effect on NA shrinking, and any significant change in the surface chemical composition. We do not report any structural classification of the resulting NAs, but we have calculated common neighbour analysis signatures and patterns.

2.1.1 Layer-by-layer analysis. We perform a layer-by-layer (LBL) analysis in the direction perpendicular to the coalescence axis, defined as the axis where the centre of mass of the individual seed lies. The analysis code is an updated version of the Sapphire code, soon available. First, a rotation of all atomic coordinates is applied to align the coalescence axis with a Cartesian axis, *e.g.*, the z -axis. Then, the NP is sliced into layers and binned in 2.25 Å, about the inter-layer distance of (111) planes in a Pd bulk. We monitor the number of layers *versus* time, $m_l(t)$, and we count the number of atoms per each chemical species α in each layer, $N_l^\alpha(t)$. The neck is the poorly populated region between the two initial seeds. It is expected when the total number of atoms per layer, $N_l(t) = N_l^{\text{Au}}(t) + N_l^{\text{Pd}}(t)$,

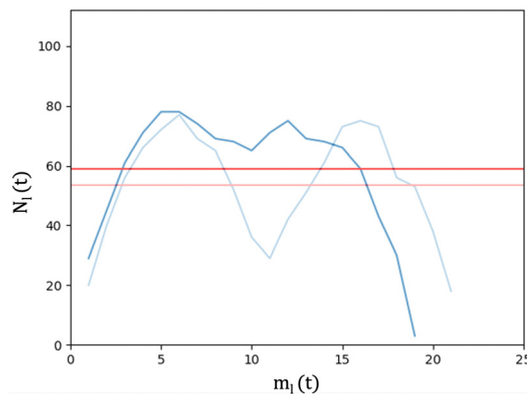


Fig. 2 LBL analysis during the coalescence of $\text{Au}_{561}^{\text{lh}}-\text{Pd}_{561}^{\text{lh}}$ in a vacuum. The straight light red and red lines are the $N_l^{\text{expected}}(t)$ values at the beginning and after 0.07 ns, respectively. The profile of the layer population $N_l(t=0)$ and $N_l(t=0.07 \text{ ns})$ are in light blue and blue, respectively. $N_l(t=0)$ shows a neck corresponding to the depression below its N_l^{expected} . After 0.07 ns, the distribution profile N_l shows an almost flat distribution in the middle region with the neck disappearing.

has a non-monotonic behaviour with respect to the layering. A neck is still present until the minimum of $N_l(t)$ falls within an intermediate position between the two centres of mass, and it is lower than the expected number of atoms per layer, N_l^{expected} . Assuming a uniform distribution of atoms in the NA, $N_l^{\text{expected}}(t) = \frac{N}{m_l(t)}$ where N is the total number of atoms and $m_l(t)$ is the number of layers at that time. If $N_l(t)$ crosses $N_l^{\text{expected}}(t)$ more than twice, we conclude that a neck is still present, as visible from Fig. 2. We define the *neck lifetime* (LTN) as the first instant since $N_l^{\text{expected}}(t)$ is crossed only twice.

2.1.2 Chemical ordering. We use various descriptors to analyse the chemical ordering in the NA on-the-fly. To characterise an atom's local environment, we start counting the number of its nearest neighbours as follows: we consider a cut-off region, described by a sphere with a radius equal to the bulk nearest neighbour distance and centred around the atom, and enumerate the other atoms that fall in such a region. Starting from this quantity, we can calculate the mixing parameter $\mu(t)$,

$$\mu(t) = \frac{\sum_{\alpha} \text{CN}_{\alpha}^{\text{homo}}(t) - \text{CN}_{\text{hetero}}(t)}{\sum_{\alpha} \text{CN}_{\alpha}^{\text{homo}}(t) + \text{CN}_{\text{hetero}}(t)} \quad (3)$$

as widely used in the literature.¹⁰ $\mu(t)$ is the ratio between the number of nearest-neighbour homo-pairs $\text{CN}_{\alpha\alpha}^{\text{homo}}$, and the total number of nearest-neighbour hetero-pairs $\text{CN}_{\alpha\beta}^{\text{hetero}}$, per each configuration at time t . $\mu(t)$ ranges between $[-1; 1]$ where -1 is for a fully alloyed, and $+1$ a complete phase separated system.

Counting homo and hetero pairs follows the introduction of a cut-off distance, R_{cut} , as the first minimum of the pair-distance distribution function, PDDF, as in Sapphire. We build the adjacency matrix from that cutoff, $A(r_{ij})$. $A(r_{ij})$ is set to 1 if the distance between atoms is less than the R_{cut} and 0 if it is greater. The atomic coordination number CN_i , is then the sum



over a row/column of $A(r_{ij})$

$$\begin{aligned} \text{CN}_i &= \sum_{j \neq i} A(r_{ij}) \\ \text{CN}_{\alpha\alpha}^{\text{homo}} &= \sum_{i,j \in \alpha, i \neq j} A(r_{ij}) \\ \text{CN}_{\alpha\beta}^{\text{hetero}} &= \sum_{i \in \alpha, j \in \beta} A(r_{ij}) \end{aligned} \quad (4)$$

where α and β label two different chemical species.

The PDDF is a crucial quantity to characterise the geometry of an NP, not only because it enables to define R_{cut} but also because a second peak of the PDDF in correspondence of the bulk lattice parameter (a_0) stems for a geometrical order.⁴⁶ We note that the maximum of the PDDF, D_{max} , provides an estimate of the NP size.

The local atomic environment (LAE) counts the total number of hetero-bonds the atom i forms. $\text{LAE} = 0$ indicates no hetero-bonds; $1 \geq \text{LAE} \leq 6$ shows a mild mixing tendency. Finally, $\text{LAE} \geq 9$ indicates that β -atoms are encapsulated into a α -matrix. Counting the occurrence of each LAE provides a clear representation of the overall chemical of the NA. For example, a core-shell nanoalloy with a one-layer thick shell of α -atoms is characterised by a negligible value of $\text{LAE}_\alpha = 0$ but a high occurrence of $1 \leq \text{LAE} \leq 6$, for both chemical species.

We evaluate the chemical radius of gyration R_{gyr}^α , the radius of gyration referred to each chemical species. R_{gyr}^α quantifies by which extent atoms are spread around their centre of mass (CoM_w);

$$R_{\text{gyr}}^\alpha = \sqrt{\frac{1}{N_\alpha} \sum_{i \in \alpha} (r_i - \text{CoM}_\alpha)^2} \quad (5)$$

where α is either gold or palladium, N_α is the total number of atoms of that chemical species, and CoM_α is the centre of the mass of N_α only. The chemical radius of gyration, combined with the distance between the centre of mass of the Au and Pd regions, ΔCoM , provides an indication of the chemical order. For example, the condition $R_{\text{gyr}}^{\text{Au}} - R_{\text{gyr}}^{\text{Pd}}$ and $\text{CoM}_{\text{Au}} - \text{CoM}_{\text{Pd}}$ occurs only in a mixed alloyed system. On the other hand, $\text{CoM}_{\text{Au}} - \text{CoM}_{\text{Pd}}$ but $R_{\text{gyr}}^{\text{Au}} > R_{\text{gyr}}^{\text{Pd}}$ stands for a core-shell with Au-rich external shell.

2.1.3 Surface identification. We identify surface atoms when their atop generalised coordination number is less than 10, as implemented in Sapphire.⁴⁵ Let $N_A^S(t)$ and $N_B^S(t)$ be the number of surface atoms of type A and B. Obviously, the total number of surface atoms is simply the sum of the two contributions, $N^S(t) = N_A^S(t) + N_B^S(t)$. The *absolute percentage* of α -atoms at the surface is given as $P_{\alpha_s, \text{abs}} = \frac{N_\alpha^S}{N_{\text{tot}}^S}$. In the ESI,[†] we provide the behaviour of the *relative percentage* $P_{\alpha_s, \text{rel}}$, defined as the percentage of atoms of a given type with respect to its total number.

3 Results

We compare the initial stages of the coalescence in the vacuum and in a certain MEI. We clarify how the neck dynamics and the shrinking process are affected by the surroundings. In particular,

we report whether the nanoalloy still presents an elongated shape or not after the observation time of 20 ns. The top row of Fig. 3 shows the evolution of the number of layers, $m_i(t)$, and their population, namely the number of atoms per each layer, $N_i(t)$, for a typical run for $\text{Au}_{55}^{\text{th}}\text{-Pd}_{55}^{\text{th}}$ and $\text{Au}_{561}^{\text{th}}\text{-Pd}_{561}^{\text{th}}$. Time is set to zero when the two seeds collide, and the layers are counted from the Pd-seed. $m_i(t)$ decreases with time, and the effect is stronger at small sizes, where in just a few frames $m_i(t)$ almost halves. On the other hand, at 1122 atoms, only a few layers are lost over 20 ns. The resulting shape is still elongated, with a peak of the most populated layers located in the Pd region. In any event, we observe a smooth population behaviour per layer.

The neck lifetime, LTN, middle row of 3, averages at 0.02 ns and always less than 0.06 ns for 110 atoms, while it ranges between 0.05 and 0.2 ns for 1122 atoms. At $\text{Au}_{561}^{\text{th}}\text{-Pd}_{561}^{\text{th}}$, a mild effect due to the environment occurs with respect to the vacuum. A few MEIs, set24, set34, set134, increase the LTN but others, set1, set3, fasten it. See the ESI[†] for details on those MEIs.

To further characterise the shrinking process, we compare the difference between the number of layers for a certain MEI, $m_i(t)^{\text{MEI}}$, and the same quantity in a vacuum, $m_i(t)^{\text{v}}$, $\Delta_{\text{layers}} = m_i(t)^{\text{MEI}} - m_i(t)^{\text{v}}$, shown in Fig. 3 at the start (grey circles) of the coalescence and after 10 ns for $\text{Au}_{55}^{\text{th}}\text{-Pd}_{55}^{\text{th}}$ and 20 ns for $\text{Au}_{561}^{\text{th}}\text{-Pd}_{561}^{\text{th}}$ (black stars). We note that the environment influence is negligible when the two seeds come in contact, suggesting that our choice of the MEI mainly affects the surface diffusion and hence the dynamics of the *shrinking process*.

The dynamics of the aggregation of $\text{Au}_{55}^{\text{th}}$ and $\text{Pd}_{55}^{\text{th}}$ remains almost unchanged, with Δ_{layers} less than half of a layer. On the other hand, the shrinking process of $\text{Au}_{561}^{\text{th}}\text{-Pd}_{561}^{\text{th}}$ depends on the surroundings, and it can be either fastened up or slowed, down. The interaction between Pd atoms and the environment becomes the critical factor, while the Au-environment interaction poorly affects the process. If Pd is strongly interacting, e.g. set34, set134, it is more likely to observe elongated shapes. However, if Pd interacts weakly ($\rho = 0.5$, set1, set2, set3, set4), the shape of the NA is more compact than in a vacuum after 20 ns.

To investigate whether the MEI affects the chemical ordering and surface composition, we compare various descriptors. Fig. 4 and 5 show the behaviour at 110 and 1122 atoms, respectively, in a vacuum and within the set34, a MEI-AP with Pd strongly interacting and Au weakly.

First, the chemical R_{gyr}^α values are spread, mainly for Au. We further observe an increment of the radius of gyration per each chemical species after the collision. In a vacuum, the Pd-spread is tiny, and at small sizes, the chemical radii of gyration are similar to those in the MEI-AP. When $R_{\text{gyr}}^{\text{Pd}} \sim R_{\text{gyr}}^{\text{Au}}$, or even longer, it is suggested that Pd atoms populate the external layer. For $\text{Au}_{561}^{\text{th}}\text{-Pd}_{561}^{\text{th}}$, $R_{\text{gyr}}^{\text{Pd}}$ evolves at the same pace as $R_{\text{gyr}}^{\text{Au}}$, indicating that both chemical species diffuse at the surface, while they were almost constant in a vacuum. The behaviour of $R_{\text{gyr}}^{\text{Pd}}$ together with the drop of ΔCoM suggests the presence of a partial core-shell chemical ordering in the AuPd in a vacuum. At the same time, we expect an intermixed surface layer in the MEI-AP.



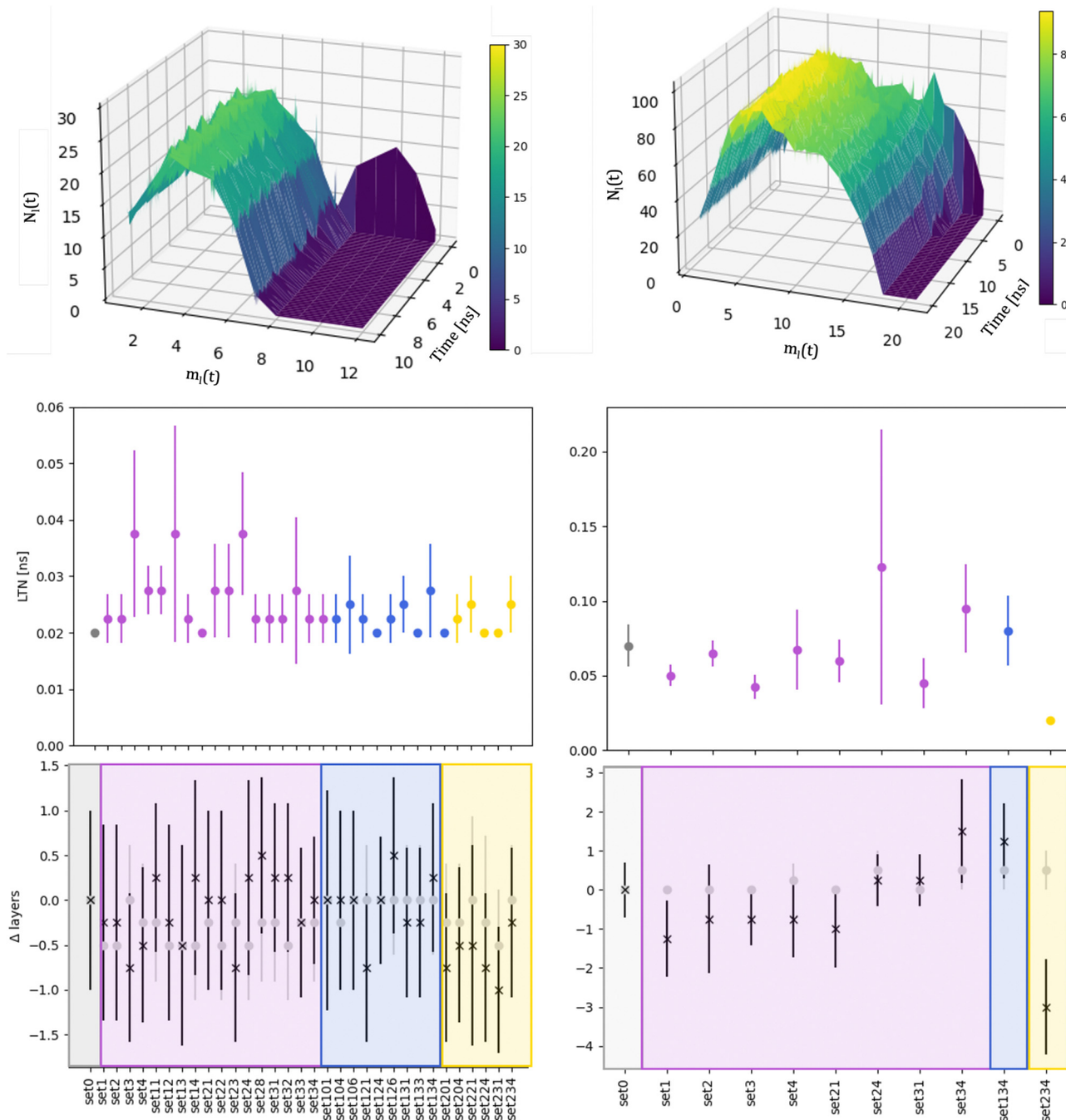


Fig. 3 Left column refers to $\text{Au}_{55}^{\text{h}}-\text{Pd}_{55}^{\text{h}}$ and right column to $\text{Au}_{561}^{\text{h}}-\text{Pd}_{561}^{\text{h}}$. Top: Distribution of the number of atoms per layer with $\rho_{\text{Au}} = 0.5$, $\rho_{\text{Pd}} = 1.5$, $\eta_{\text{Au}} = 0.02$ eV and $\eta_{\text{Pd}} = 0.8$ eV, aka set 34. The colour gradient represents $N_i(t)$. Middle: Mean value of the LTN and its standard deviation over independent simulations for each MEI. Vacuum, labelled as set0, is in grey. Bottom: Difference between the number of layers $N_i(t)$ in an MEI and the vacuum (Δ_{layers}) averaged over the independent simulations and their standard deviation at the initial time (grey full circles) and after 10 ns for $\text{Au}_{55}^{\text{h}}-\text{Pd}_{55}^{\text{h}}$ and 20 ns for $\text{Au}_{561}^{\text{h}}-\text{Pd}_{561}^{\text{h}}$ (black stars). For the middle and bottom rows, the x-axis labels the environment; MEIs are coloured in accordance with chemical species interactions: only Pd is blue (MEI-P); only Au is yellow (MEI-A); both Au and Pd interacting (MEI-AP) are violet. For a full description of the MEI, see Table S1 of the ESI.†

Such a picture is confirmed by the relative abundance of surface atoms; see bottom rows of Fig. 4 and 5. For coalescence in a vacuum, the surface is mainly composed of Au atoms. Introducing a strongly interacting environment with Pd only, this chemical species tends to be at the surface, occupying

about 50% of the surface area, Fig. 4, and a $\mu(t)$ of just 0.4, much less than 0.6 in a vacuum. The more we decrease the values of ρ and η , the more the μ tends to its value in a vacuum. Those results suggest that mildly interacting environments, both in terms of the type of interaction and the strength of



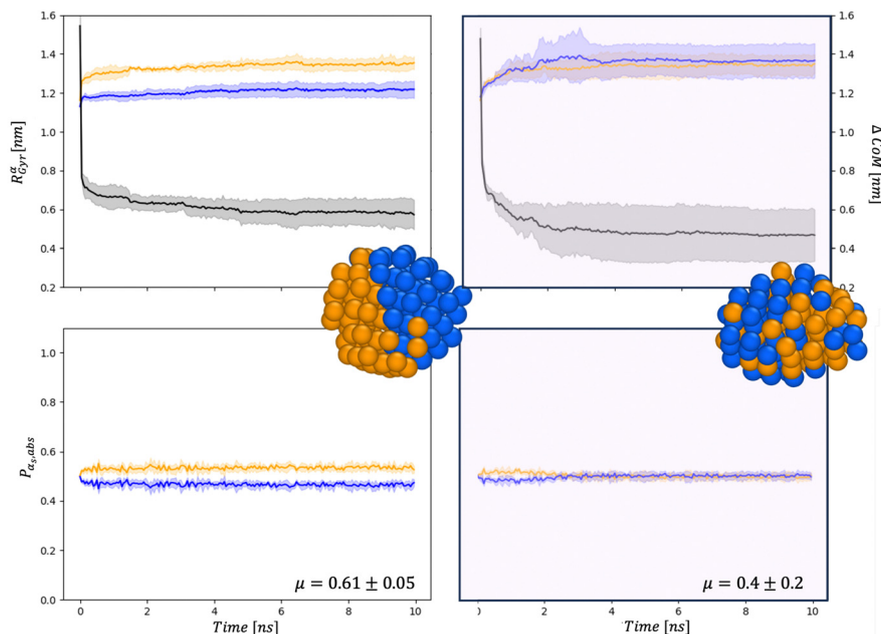


Fig. 4 Chemical distribution in $\text{Au}_{55}^{\text{Ih}}-\text{Pd}_{55}^{\text{Ih}}$ in a vacuum (left) and in set34 (right). Paradigmatic snapshots taken after 10 ns, where Au atoms are represented in orange and Pd in blue. Top panel: Referring to the left y-axis: time evolution of R_{gyr}^z in nm. The solid line is the averaged value over the independent simulations, Au in orange and Pd in blue, respectively. The shadow represents their standard deviation. The time evolution distance between the centres of mass of the two chemical species ΔCoM , referring to the right axis, is represented in black. Bottom panel: Time evolution of the percentage of surface atoms of type α with respect to the number of surface atoms, $P_{\alpha,\text{abs}}$. Colour as in the above panels. Also, the μ at 10 ns averaged over the four independent simulations, with its standard deviation is reported in these figures.

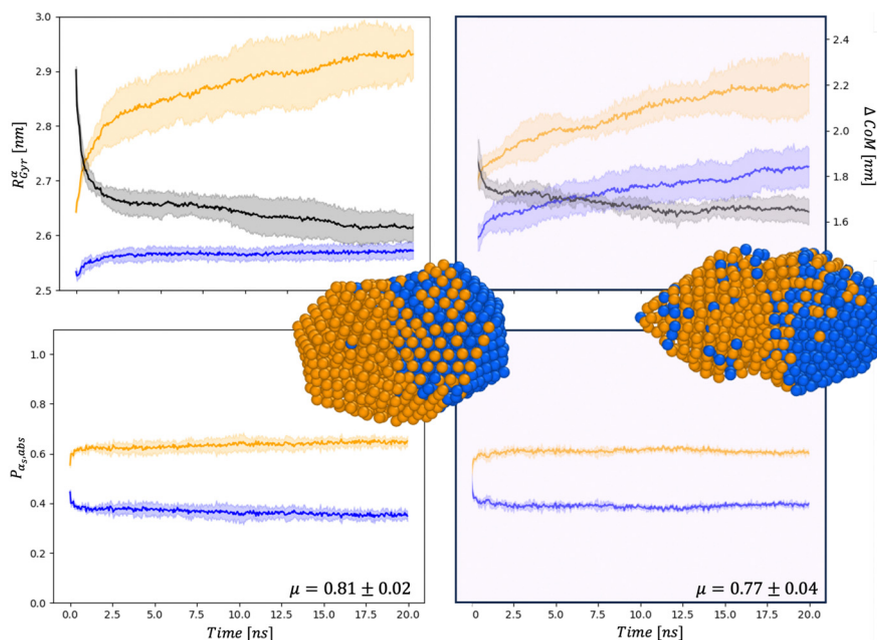


Fig. 5 Chemical distribution for $\text{Au}_{561}^{\text{Ih}}-\text{Pd}_{561}^{\text{Ih}}$ in a vacuum (left) and in set34 (right) for R_{gyr}^z in nm and ΔCoM in nm (top), and $P_{\alpha,\text{abs}}$ (bottom). Snapshots are taken after 20 ns with the same colour-coding as in Fig. 4.

the interaction, will poorly affect the dynamics of aggregation and the overall chemical stability of NAs. At larger sizes, the MEI affects the chemical ordering, although less pronounced. We

believe this is only a kinetic effect, as shown by a more elongated shape. Full details of the $\mu(t)$ analysis are reported in the ESI† but they confirm a tendency to mix in the considered MEI-AP.



At 110 atoms, for set0 or the vacuum, we observe a predominance of ball-cup chemical ordering, where the Pd region is only partially covered by an Au-monolayer.

To be more quantitative, we analyse the local atomic environment (LAE) of each chemical species; see Fig. 6. For $\text{Au}_{55}^{\text{Ih}}\text{Pd}_{55}^{\text{Ih}}$ after 10 ns in a vacuum, $\text{LAE}_{\text{Au}} = 0\text{--}40\% < \text{LAE}_{\text{Au}} \in (3\text{--}6) \sim 55\%$ and $\text{LAE}_{\text{Pd}} = 0 \sim \text{LAE}_{\text{Pd}} \in (3\text{--}6) \sim 45\%$, while there is a negligible occurrence of Au and Pd atoms in a thoroughly mixed local environment ($\text{LAE} \geq 9$). We note that $\text{LAE}_{\text{Au}} = 0$ is always less or equal to $\text{LAE}_{\text{Pd}} = 0$, supporting the idea that Au atoms diffuse over Pd. Furthermore, a feature of $\text{LAE}_{\text{Pd}} = 0$ about 50% suggests that the Pd-seed is at least partially preserved. At the same time, but in a few cases, we observe individual Pd atoms embedded in an Au-matrix ($\text{LAE}_{\text{Pd}} \geq 9$, see the inset panel of 6). For example, set34, set134, set 124, set224, set 231, set 234 suggest a different chemical ordering than the vacuum highlighted by a drop of each $\text{LAE} = 0$ together with an increment of the $\text{LAE} \in (3\text{--}6)$. In particular, the LAE signatures for set234, namely $\text{LAE} = 0$ and $\text{LAE} \geq 9$ both lower than 10% and $\text{LAE} \in (3\text{--}6)$ above 80% for both chemical species suggest the formation of an Au-shell over a Pd-core. On the other hand, set34, set134, predict a stronger mixing with some Pd atoms dispersed in the Au matrix and a low $\text{LAE} = 0$ and a high $\text{LAE} \in (3\text{--}6)$ for both chemical species.

At 1122, the effect of the MEI seems less evident because the simulation time is relatively shorter than at the 110 atoms. Nonetheless, there is a clear indication that set34, set134 behave differently than the vacuum. In particular, $\text{LAE}_{\text{Pd}} \geq 9$ is almost 2%, and there is an increasing trend of mixing both at the surface and in the inside.

For both sizes and all the MEIs, a greater structural order emerges at the end of the simulation (10 ns for $\text{Au}_{55}^{\text{Ih}}\text{Pd}_{55}^{\text{Ih}}$ and 20 ns for $\text{Au}_{561}^{\text{Ih}}\text{Pd}_{561}^{\text{Ih}}$) to the time zero, set as the contact time of the two seeds, see PDDFs profiles in Fig. S4–S27 and S28 (ESI[†]).

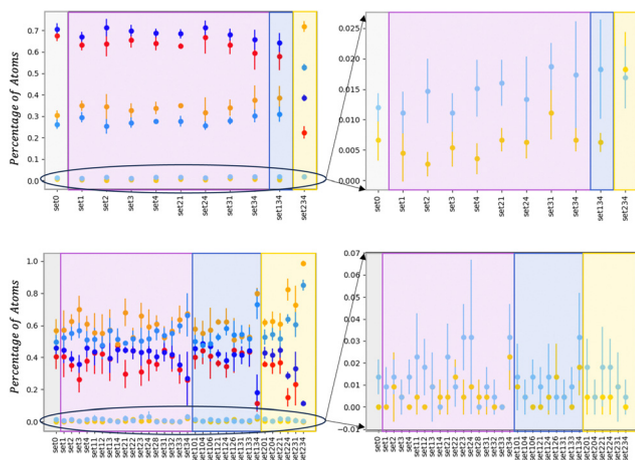


Fig. 6 Percentage of the LAEs defined in the text for each chemical species, namely Au: $\text{LAE}_{\text{Au}} = 0$ in red, $\text{LAE}_{\text{Au}} \in (3\text{--}6)$ in orange, ≥ 9 in yellow. Pd chemical LAEs are in blue, light-blue, sky, respectively. The right column is a zoom for the LAEs ≥ 9 . They refer to the last configuration and averaged over all independent simulations for $\text{Au}_{561}^{\text{Ih}}\text{Pd}_{561}^{\text{Ih}}$ (top) and $\text{Au}_{55}^{\text{Ih}}\text{Pd}_{55}^{\text{Ih}}$ (bottom).

A deeper look indicates that the Pd region displays a geometrical order, even at small sizes, while the PDDF of the Au-part does not often show a peak at the lattice distance, Fig. S7–S9 (ESI[†]). In any event, the distance between the two seeds is similar and tends to be close to 1.6.

For $\text{Au}_{561}^{\text{Ih}}\text{Pd}_{561}^{\text{Ih}}$ in a vacuum, the Pd region tends to remain as the original icosahedral seed, with few Au atoms hopping over its surface. In contrast, in the presence of a strong MEI-P or a MEI-AP, an intermixing of Pd and Au is observed at the surface. As Pd strongly interacts with the surroundings, Pd atoms spread randomly around the surface layer, eventually clustering along low coordinated sites, *i.e.* edges, to balance the interaction with the implicit environment.

3.1 Effect of the initial seed-shape

Because the symmetry of nanoparticles is a critical parameter, we consider another initial seed for Au and Pd, namely a cuboctahedron (Co) of 55 atoms. The choice is motivated by the fact that Co has the same geometrical shell closure sequence as Ih. We therefore analyse the coalescence of $\text{Au}_{55}^{\text{Ih}}\text{Pd}_{55}^{\text{Ih}}$, $\text{Au}_{55}^{\text{Co}}\text{Pd}_{55}^{\text{Ih}}$, and $\text{Au}_{55}^{\text{Co}}\text{Pd}_{55}^{\text{Co}}$. Fig. 7 summarises and compares all our findings. In a vacuum, Dh-nanoparticles are more likely to grow, albeit incompletely and asymmetrically, featuring a Janus-like chemical ordering, identified by a high percentage of both $\text{LAE}_{\text{Au}} = 0$ and $\text{LAE}_{\text{Pd}} = 0$. When only Pd strongly interacts with the environment, the percentage of FCC is not negligible, alongside Dh and Ih-like geometry. The chemical ordering shows a tendency of Pd atoms to diffuse

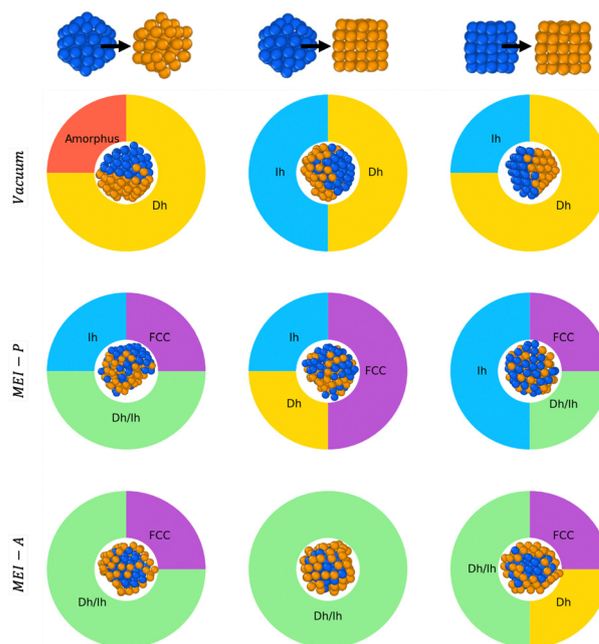


Fig. 7 Effect of the initial seed geometry – Ih and Co – on the coalescence of 55-atom Au and Pd colliding. Structures are collected after 10 ns and classified on the basis of their CNA and LAEs. Paradigmatic examples are shown for each case with Au and Pd coloured in gold, blue respectively. Each row corresponds to a different MEI: from top to bottom vacuum, MEI-P, MEI-A.



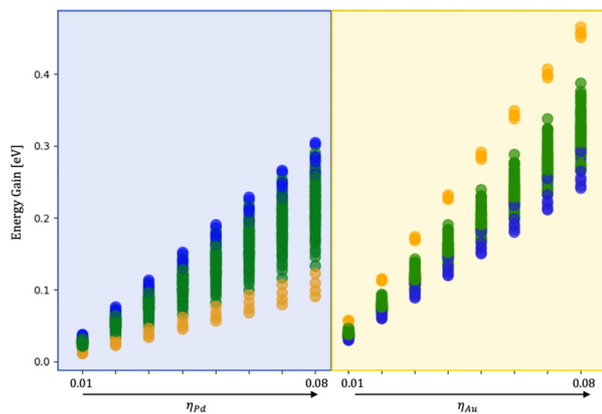


Fig. 8 Energy gain per atom for AuPd in various environments w.r.t. the vacuum varying the strength of Pd (blue shadow)/Au (yellow shadow) interaction (η_x), fixing ρ_x at 1.5. Orange circles represent core-shell initial configurations, mixing configurations are in blue and Janus-like in green. An inversion of the chemical ordering stability occurs as soon as only Pd interacts while AuPd favours core (Pd)-shell (Au) ordering.

over gold, and the NA surface is Pd-rich. Conversely, when only Au interacts with the environment, the chemical ordering is core-shell, with a net tendency for incomplete Dh structures covered by an HCP island.

3.2 Environment effect on the chemical ordering

To quantify the effect of the environment, we have performed additional ionic relaxation of the 120 isomers obtained during the coalescence varying the MEI, namely only Pd and only Au interacting fixing $\rho = 1.5$ and η between 0.01 and 0.08 eV per atom.

Fig. 8 shows the energy gain per atom following the embedding of the nanoalloy into a medium. The core-shell chemical ordering is the most favourable in a vacuum with Pd in the core. Such tendency is also evident during melting simulations starting from various configurations obtained during the coalescence. Indeed, even starting from a Janus-like order before melting, a 110-atom AuPd rearranges into an Au-shell over a Pd-core. However, a liquid nanodroplet always displays a randomly mixed chemical ordering with a mild tendency to have gold atoms at the surface.

4 Conclusions

Through classical molecular dynamics simulations, we model the coalescence of gold and palladium NPs, both in a vacuum and embedded in an environment, through an implicit model.

We examine the coalescence of two 55-atom seeds, simulated at 400 K, and of two seeds of 561 atoms each, simulated at 600 K. We selected icosahedral seeds of the same size but we checked that similar results are available even for colliding two cuboctahedral seeds and Ih and Co. We tune the parameters of the interacting environment, obtaining more than 30 different scenarios. We focus on the initial steps of the coalescence process, and we provide insights into the kinetics of the process

by looking at the different surface chemical compositions after 10–20 ns. We also comment on the temperature stability of various chemical orderings approaching the melting transition.

Our analysis reveals that the neck disappears in 0.06 ns for 110 atoms and 0.2 ns for 1122 atoms, independent of the type of surrounding. We can conclude that the environment poorly affects the neck's lifetime, which is mainly tuned to the seed size. Nonetheless, alongside the size effects, the surrounding interaction is critical concerning the rate of shrinking and the chemical ordering. Notably, when observing the process over 20 ns, specific environments induce a more compact shape than vacuum conditions. In contrast, others slow the process, leading to more elongated shapes than in a vacuum.

At small sizes, we observe a pronounced tendency towards the formation of ball-cup structures, in good agreement with global optimisation studies,⁴⁷ where the innermost layers are mainly of Pd and gold atoms diffuse above the Pd seed to form a partial external shell. Nonetheless, when Pd interacts strongly with the environment, the preferential chemical ordering changes towards a mixed nanoalloy with both Pd and Au equally abundant at the NA surface, as in Fig. 8. We observe that Pd tends to keep its icosahedral shape at large sizes, and Au diffuses upon it. In strong MEI-AP and MEI-P, we reveal the formation of an intermixed external shell, with Au diffusing over Pd atoms and a few Pd atoms inter-diffusing in Au layers.

In conclusion, our research shows clearly that the presence of an interacting environment affects the kinetics of the coalescence process. In particular, we highlight the effect of the shrinking process on the surface chemical composition of AuPd NAs. Our work shows the need to include the environment when modelling the trajectory of NA formation and, hence, their chemophysical properties. Indeed, as the kinetics of shrinking are dependent on the size and environment, it is not obvious that extrapolation to other sizes or chemical compositions is necessary. Since Au-based nanoalloys play pivotal roles in plasmonic devices and nanocatalysis, variations in environmental conditions can lead to changes and rearrangements in their morphology, thereby influencing their optical response and possibly their performance. Additionally, strongly interacting environments with the dopant transition metals, but not Au, will impact the surface chemical composition of the NAs, affecting their catalytic activity.

Data availability

All the MD trajectories obtained are available on Unimi DATAVERSE?.

Conflicts of interest

There are no conflicts to declare.

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