

Cite this: DOI:00.0000/xxxxxxxxxx

Born to be different: the formation process of Cu-nanoparticles tunes the size-trend of the activity for CO₂ to CH₄ conversion

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Received Date

Accepted Date

DOI:00.0000/xxxxxxxxxx

We investigate the impact of the formation process on the distribution of adsorption sites of Cu nanoparticles and hence on their activity. Using molecular dynamics, we model formation pathways characteristic of physical synthesis routes as the annealing of a liquid droplet, the growth proceeding via the addition of single atoms, and the coalescence of individual nanoparticles. Each formation process leads to different and characteristic size-dependent distributions of their adsorption sites, catalogued and monitored on-the-fly by means of a suitable geometrical descriptor. Annealed or coalesced nanoparticles present a rather homogeneous distribution in the kind and relative abundance of non-equivalent adsorption sites. Atom-by-atom grown nanoparticles, instead, display a more marked occurrence of adsorption sites corresponding to adatoms and small island on (111) and (100) facets. Regardless of the formation process highly coordinated sites are more likely in larger nanoparticles while the abundance of low-coordination sites depends on the formation process and on the nanoparticle-size. Furthermore, we show how each characteristic distribution of adsorption sites reflects in different size-trends for the Cu-nanoparticles activity, taking as an example the electro-reduction of CO₂ into CH₄. To this end we employ a multi-scale method and observe that the faceted but highly defected structures obtained during the atom-by-atom growth become more and more active with increasing size, with a mild dependence on the original seed. On the contrary, the activity of Cu-nanoparticles obtained by annealing decreases with their size while coalesced nanoparticles' activity shows a non-monotonic behaviour.

1 Introduction

Reaching the net-zero CO₂ emissions by 2050 requires us to develop new devices to capture and convert CO₂ efficiently. As CO₂ is thermodynamically stable and kinetically inert^{1,2}, its efficient conversion requires the use of catalysts at the electrodes capable of selectively reducing CO₂ to valuable (e.g. methane (CH₄), oxygenated and/or C₂ products) hydrocarbon products at low overpotentials. Out of the polycrystalline metals, copper (Cu) is the only heterogeneous catalyst that has a potential to produce valuable hydrocarbons, aldehydes and alcohols from the electrochemical CO₂ reduction reaction (CO₂RR)¹. Even if copper has already been largely used as a catalyst to reduce CO₂^{1–19}, it is still a challenge to get optimal efficiency and selectivity.²⁰ Size-dependent

selectivity for ethylene in Cu nanocubes exposing mostly (100) facets has been recently reported.²¹ At the same time, high-selectivity for methane in Cu nanowires,²² spheres²³, and octahedra, mostly exposing (111) facets,²⁴ have been also observed. Alloying Cu and other metals also may increase nanoparticles and nanocrystals activities and selectivity^{25,26}. The key to the ultimate design of better Cu-nanocatalysts requires a systematic understanding of CO₂ reduction on the variety of adsorption sites peculiar of individual nanoparticle, and how structural instability and rearrangements affect their activity.

To propose robust design-rules for nanocatalysts -e.g. size and shape trends of their activity; effect of chemical composition and ordering- avoiding a costly and ineffective trial-and-error procedure, two factors should be simultaneously taken into account: (i) the variety of adsorption sites per isomer^{27–32}, which influences both activity and selectivity, and (ii) the coexistence of different geometries, eventually related to their formation process^{33–35}.

In this work, we find that the size-distribution of adsorption sites depends strongly on the formation process, ultimately affecting the size-trend of the mass activity of Cu-NPs. Furthermore, because such distribution of adsorption-sites is characteristic of the selected formation process, our works opens a way of tuning

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† Electronic Supplementary Information (ESI) available: initial configurations geometries - Molecular Dynamics Simulation set-ups - Geometric characterization routines - NanoCHE details for calculating NPs activity - Considerations on the potential energies of NPs sampled during different formation processes. See DOI: 00.0000/00000000.

the catalytic activity by choosing a certain formation protocol. We model the formation of individual Cu-NPs as it is supposed to occur inside inter-gas aggregation sources

36? ? ? -39.

We observe that annealing and coalescence lead to a rather homogeneous distribution of the type and occurrence of adsorption sites as classified accordingly to their generalised coordination number. Atom-by-atom growth, instead, results in characteristic distributions with peaks corresponding to the formation of adatoms and small island on (111) and (100) facets. In particular, the occurrence of low coordinated atoms peaks at different NPs sizes, depending on the formation process. Low coordinated atoms are more abundant in small frozen Cu-NPs, coalesced ones of intermediate size (300-600 atoms), and in larger atom-by-atom grown Cu-NPs (600-1000 atoms). For the case of highly coordinated sites (i.e. the surface atoms presenting a coordination equal or greater to the one on a (111) facet), their occurrence increases with the size of the Cu-NPs, regardless of the formation process.

To calculate the mass activity of Cu-NPs for CO₂ conversion to CH₄ we adopt our recent NanoCHE scheme,⁴⁰ a multiscale method to estimate the catalytic activity of NPs as a function of their structure and structural evolution. By exploiting a scaling relationship previously proposed in the literature,¹⁴ we unveil non-trivial size trends of the NP-activity depending on the the formation process. Using a volcano-plot relationship between coordination and activity which peaks at extremely low coordinated sites we observe that, at small sizes (below 300 atoms), NPs obtained from annealing are the most active in converting CO₂ to CH₄. At intermediate sizes (300-600 atoms) coalesced Cu-NPs are instead more favourable. Their size-dependent activity also shows a non-monotonic trend for the sizes and sampling time here considered. At larger sizes (600-1000 atoms) instead atom-by-atom grown NPs appear more active. We further note that, while the activity on atom-by-atom grown NPs positively correlates with size, we also find that the latter trend is oscillatory in nature.

2 Methodology

The workflow of our multi-scale approach is visually represented in Figure 1 and is based on the following steps:

2.1 Modelling the formation process

We consider Cu-NPs of different sizes and shapes between 110 and 976 atoms, i.e. with a diameter of 1-3 nm. We label each structure as a function of the number of atoms it is made of, e.g. the nomenclature Cu 147 labels a Cu NPs of 147 atoms. NPs initial geometries are: liquid droplets, FCC polyhedra as Octahedra (Oh) or their regular truncation (To), Marks-Decahedra (MDh), Icosahedra (Ih) and defected Icosahedra (dIh), as detailed in Section I of the Supplementary Information. Far from exhaustive of all available isomers in the considered size range, the ensemble of morphology under scrutiny provides, in our opinion, a sufficient structural variety to test the effectiveness of our approach. In fact there is a strong experimental and theoretical evidence which backs up this choice. Icosahedra and decahedra, geometries with 5-fold twinning planes, and a FCC-polyhedra are of

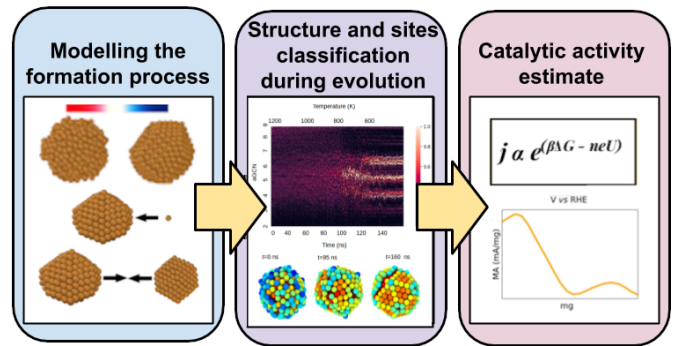


Fig. 1 Graphical representation of the multiscale numerical strategy adopted here. The modelling of the formation processes (annealing, growth, coalescence) is by means of molecular dynamics (MD) staring from a plethora of Cu-NPs of different shapes and sizes. We perform an accurate structural analysis of each trajectory and per each time-step the NP is classified into geometrical families and the type and the occurrence of each adsorption site is monitored following the time evolution of the atop generalised number. We then estimate the CH₄ mass activity for each Cu-NP during their formation process by the multiscale model of Ref.⁴⁰

ten observed in experiments. Further, theoretically Ih, MDh, and FCC-polyhedra are the most favourable morphology for sizes between 1-3 nm both for “magic” and “non-magic” sizes⁴¹⁻⁴⁴, in good agreement with experiments.^{39,43,45,46}

Among the many formation processes enabling the making of metallic nanocatalysts, we consider the annealing of a liquid droplet, the atom-by-atom growth from a small seed, and the coalescence of two Cu-NPs. We simulate these formation processes by means of Molecular Dynamics and model the interatomic interactions with a Rosato-Guilpo-Legrand potential with parameters for Cu taken from the literature⁴⁷. The latter parametrisation was fit to reproduce the bulk modulus, elastic constant, and cohesive energy and was found to quantitatively reproduce size-crossovers among structural motifs in Cu and other transition metal nanoparticles.⁴¹ We employ a velocity-Verlet algorithm⁴⁸ to integrate Newton’s Equation of motion with a time step of 5 fs. The latter timestep is possibly larger than the one usually employed in ab initio molecular dynamics runs, but is commonly used in simulations employing semi-empirical potentials. Indeed the stability and faithfulness of dynamics with second-moment tight-binding formulations was verified also for four-times larger timesteps.⁴⁹ We apply an Andersen thermostat to control the temperature with a frequency of 5×10^{11} Hz.

During annealing simulation the structural evolution of a liquid droplet of 147, 250, 561, 585, 891, or 976 atoms is probed over various concatenated canonical NVT runs, where the temperature is lowered from 1200 K to 400 K at a rate of 5 K/ns. In the atom-by-atom growth simulation, we consider three starting seeds of similar size (Cu₁₄₆ (Oh), Cu₁₄₇ (Ih) and Cu₂₀₁ (To)) and assume that the Cu-NPs are grown by the addition of single atoms, deposited every 4 ns with a temperature of 600K.⁵⁰ To study coalescence we investigate the evolution of two nanoparticles moving against each other with a kinetic energy corresponding to a temperature of 600 K. We analyse the first 100 ns of

their evolution. Per each size and formation process, we perform 10 independent runs, and we average results over them. Section II of the Supplementary material contains further details on the simulation methods to model different formation processes.

2.2 Monitoring the adsorption sites distribution on-the-fly:

We employ Common Neighbor Analysis (CNA) to classify the morphology of the NP into geometrical families - namely FCC, icosahedra (Ih) and decahedra (Dh) - as well as to probe defects formation - i.e. re-entrances and elongated concavities - and the formation of twin planes, grain boundaries, and five-fold axes (see SI section III for more details).

We use the atop generalized coordination number (aGCN) to enumerate the number and kind of all adsorption sites in the nanoparticle. The aGCN of an adsorption site i is defined as:^{27,28}

$$aGCN_i = \frac{\sum_j CN_j}{CN_{max}} \quad (1)$$

where $CN_{max}=12$ and CN_j is the nominal coordination number of the j first nearest neighbour atoms of i . CN_j is calculated by enumerating the number of atoms within a cut-off distance r_{cut} , which accounts to 3.145 Angstroms. This distance is found by locating the first minimum after each nearest neighbour peak in the pair-distance distribution function of each system, averaged over all the time steps. For reference, we display a graphical example about the GCN calculation in the top panel of Figure 2. We instead refer the interested reader to the SI for a cut-off distance choice sensitivity analysis.

As depicted in the middle panel of Figure 2, sites corresponding to vertexes and single adatoms have an aGCN less than 4, while atoms lying at the edges of any facet shows an aGCN between 4 and 5. Sites on (111) and (100) surface respectively display a generalized coordination of 7.5 and 6.67. The aGCN further discriminate atoms still belonging to a facet but closer to the edge, with a value between 5 and 6.5. Atoms at convex sites in correspondence of steps, kinks, and re-entrances instead display a coordination larger than 7.5. Atoms that are not at the surface have a aGCN approaching the bulk limit of 12. It is worthy to note that while we focus only on atop adsorption sites, as CO likely adsorbs in that way, we note that our framework could also be adapted to account for bridge and hollow adsorption too.

2.3 Estimate of the NPs' mass activity:

To monitor the activity of Cu-NPs during their formation process and to derive a size-behaviour of the mass activity, we employ the NanoCHE model.⁴⁰ The latter is a multiscale model built upon the Computational Hydrogen Electrode model, pioneered by Norskov and co-workers,⁵¹ which enables to simulate a linear sweep voltammetry experiment for systems characterised by a variety of non-equivalent and competing adsorption sites.

Within the NanoCHE framework, for a temperature T and a certain applied voltage U^* , the mass activity MA_{NP}^* of a nanoparticle is calculated from the current density it produces, J , its mass

M_{NP} , and its area, A_{surf} :⁴⁰

$$MA_{NP}^*(T, U^*) = \frac{A_{surf}}{M_{NP}} J = \frac{A_{surf}}{M_{NP}} \sum_{i \in \{\text{aGCN}\}} C \frac{\Omega(i)}{N_{sites}} i e^{\beta(\Delta G(i) - neU^*)} \quad (2)$$

In Equation, 2 M_{NP} is estimated from the number of atoms, N , in the NP, times their atomic mass, i.e. 63.546 a.u. for Cu. A_{surf} is instead calculated as a function of the aGCN⁴⁰, $A_{surf} = \sum_N^N 4\pi r_{\text{atomic}}^2 \left(1 - \frac{\text{aGCN}(n)}{12}\right)$ where the sum is over all atoms

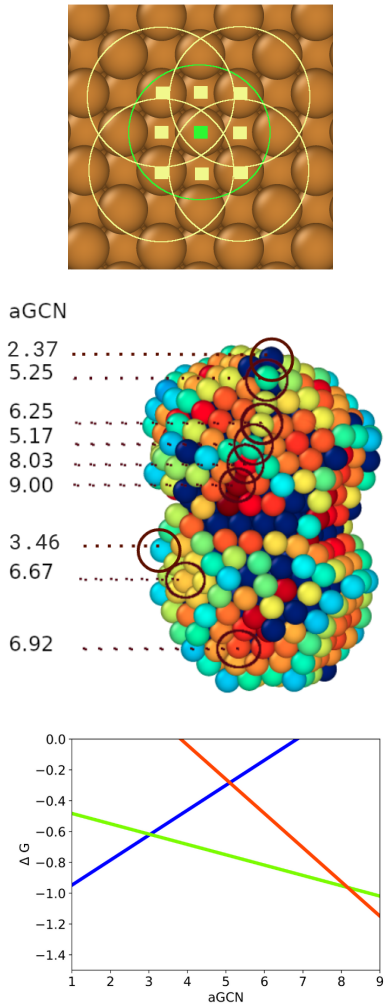
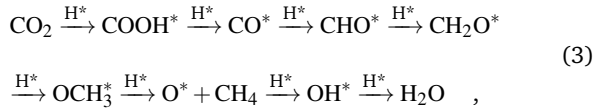


Fig. 2 Upper panel: Visual example of how the GCN for an atom (labelled with a green square) on a (100) surface is evaluated. We identify the atoms (pale yellow square) in its first neighbour shell (green circle), and calculate their coordination number, by counting the atoms in their first nearest neighbours (yellow circle). We finally evaluate the GCN of the atom labelled in green following Eq 1. Middle panel: a Cu nanoparticle where surface atoms are coloured according to their aGCN, following the colour gradient 2 → 9. The aGCN is also spelt out for example sites. Any atom with more than 11 nearest-neighbour, are coloured in dark/black blue and they are not able to act as anchor for CO-adsorption. Lower panel: the volcano relationship of Eq. 4, which correlates the site aGCN to the $\text{CO}_2 \rightarrow \text{CH}_4$ reaction free energy, adapted from Ref.¹⁴. The rate limiting step for sites with coordination lower than 3.1 is H_2O desorption (blue line), for sites with aGCN between 3.1 and 8.4 CO protonation to form CHO (green line), and for sites above 8.4 the formation of COOH from CO_2 (red line).

N in the NP, and r_{atomic} is the atomic radius of Cu, 0.128 nm. $aGCN$ is the list of all the atop generalized coordination number that a NP shows, and it is constantly updated during the evolution of the NP. $\frac{\Omega(i)}{N_{sites}}$ counts the number of adsorption sites with $aGCN = i$ over the total of surface sites. The constant C is fitted to reproduce experimental activities. As discussed in detail Sect. IV of the supplementary information, C is found equal to $-3.01 \cdot 10^{14}$ mA/cm². β is a short-hand notation for the Maxwell-Boltzmann factor, $\frac{1}{K_B T}$. To scrutinise on equal footing structures sampled within different ensembles, T is kept at 300 K, the temperature at which catalytic activities are usually probed experimentally.^{52,53} We note that while Cu-NPs may be formed at higher temperatures, they do not undergo major structural rearrangements if rapidly annealed to room temperature,⁵⁴ hence justifying the choice of probing all NPs activities at room temperature. Finally, ΔG is reaction free energy of the rate limiting step of the chemical reaction under study, calculated as a function of the atop generalised coordination number of the site where the reaction occurs. While the NanoCHE was first developed to predict the activity of Pt nanoparticles over oxygen reduction, we show here that it could be extended easily to other electrochemical reactions. From this point of view, the NanoCHE model appears as a quite general approach to estimate the activity of metallic nanoparticles. In this view, we critically assess further improvements.

Under the assumption of Broensted-Evans-Polyani relationships holding for the process of interest,⁵⁵ the difference in binding energies between the two adsorbates determining the rate limiting step conversion dictates the activity of the site at which the reaction occurs. Although the pathway from CO₂ to CH₄ on Cu sometimes shows bifurcations (i.e. CHO can attach via C or O)⁵⁶, the reaction steps established by Peterson et al.^{13,57} were employed to calculate the reaction free energy:



where * stands for adsorbed molecules and the rate limiting step is, on the majority of sites, the protonation of adsorbed CO to make adsorbed CHO¹⁵. Following ab-initio calculations,¹⁴ it was found that the binding energy of CO₂ conversion to CH₄ intermediates display a linear relationship with the generalized coordination of the site at which the reaction occurs. Conversely the reaction free energy ΔG of the rate limiting step of the reaction can be written as a function of the atop generalized coordination number of each surface atom n at which CO can be adsorbed:

$$\Delta G = \begin{cases} +0.162 aGCN_n - 1.11 & \text{if } aGCN_n < 3.1 \\ -0.067 aGCN_n - 0.416 & \text{if } 3.1 < aGCN_n < 8.4 \\ -0.222 aGCN_n + 0.894 & \text{if } aGCN_n > 8.4 \end{cases} . \quad (4)$$

For completeness, in Figure 2 middle and lower panels we report the volcano shaped structure-activity relationships and show adsorption sites with different generalized coordination to exemplify their coordination environments. We note that such relationsips are extracted from GGA DFT calculations employing a re-

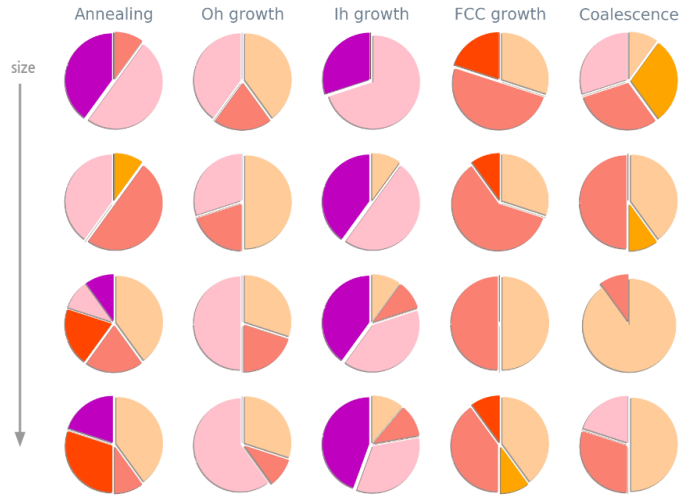


Fig. 3 The pie charts represents the statistical shape-distribution from 10 independent simulations. Each row corresponds to a certain size (from top to bottom: 240-250, 581-585, 719 and 966-976 atoms). The colours refers to the formation of icosahedra (purple), defected Ih (pink), FCC (red), defected FCC (light red), i.e. FCC with a twinning plane, decahedra (orange), defected (light orange) dDh. Such classification is done on the basis of CNA-signatures – such as (5,5,5), (4,2,1) and (4,2,2) – which enable to distinguish Ih, Dh, and FCC. A shape is classified into a certain family when more than the 70% matches the reference values.

vised Perdew-Burke-Ernzerhof functional and no dispersion correction. The choice of functional may affect the trends observed in numerical investigations.⁵⁸

3 Results and discussion

3.1 Formation process and NP structure

Depending on the formation process, a variety of geometries is observed in the size range between 1-3 nm. These are visually resumed in Figure 3 where, for each formation process, we report the proportion of trajectories which result in the formation of an Icosahedron, a Decahedron, a fcc structure, or their defected counterparts.

In agreement with previous study, we observe a structural crossover with size.⁴¹⁻⁴⁴ At small ones, the Ih archetype and its defected counterpart are more likely to be observed. They represent the majority of all structures but for the case of growths started from a truncated Octahedron. Increasing the size of the nanoparticle FCC like structures and defected decahedra are more probable, until they become the most likely to be observed. Such structural crossover sizes appear formation process dependent.

We remark, however, that this analysis offers only a qualitative indication of the true distributions of morphology resulting in each of the formation processes under consideration. It is likely that a larger number of independent runs are needed to obtain finely converged statistics. Nevertheless, our multi-scale model to calculate the activity of Cu-nanoparticles assumes an explicit dependence on the distribution of non-equivalent adsorption sites only. The statistics to converge the latter converge for a lesser number of runs, vide infra.

The heterogeneity in morphology distributions stems from ki-

netic factors and from the intrinsic complexity of the potential and free energy landscape of metallic NPs. Pathways connecting different isomers may appear either as shallow, resulting in fast and fluxional rearrangements among different geometries, or characterized by high energetic barriers, determining kinetic trapping in one isomer, and the appearance of surface defects.⁵⁴ Section V of the supplementary material reports a detailed discussion on the energetic landscapes of the systems under investigations. Our study on the formation process of 1-3 nm Cu nanoparticles confirms that:

- The nucleation into a morphology or another is the most likely cause for the morphological heterogeneity during the annealing.⁵⁹
- The interplay between initial seed geometry and kinetic effects determines the formation of either icosahedral-like or asymmetric-FCC geometries.⁶⁰
- Nanoparticles from the coalescence are likely to assume non-spherical geometries.⁶¹

3.2 Shape and Non equivalent site distribution evolution during each formation process

The heterogeneity in the shapes sampled during annealing, growth, and coalescence parallels the one in the non-equivalent adsorption sites distributions. Figure 4 reports paradigmatic evolution of the occurrence of the number and kind of NEAS of nanoparticles during each formation process. In the top panel, there is an example annealing trajectory. At the beginning, the aGCN distribution is widely spread for liquid droplets and it narrows down to specific signatures when the NP solidifies. Such a transition in the aGCN distribution emerges at the phase transition temperature. Another general consideration about the differences between solid and liquid nanoparticles pertains the average coordination number observed in the two phases. Nanodroplets are by definition more disordered and this reflects in a lesser coordination of atoms at their surface. We indeed observe a decrease of the surface area during the annealing process, dropping from values between $1.2\text{-}5.4 \cdot 10^{-15} \text{ cm}^2$ to $1.3\text{-}8 \cdot 10^{-15} \text{ cm}^2$, respectively. Together with Figure 4 top panel, the interested reader is referred to the Supplementary Information Figure S5 upper panel for other examples corroborating this observation.

When analyzing the evolution of the aGCN distributions during a growth process as a function of time (and thus size) we note that the single addition of an adatom on a rather close-shell geometry results in the formation of low coordinated sites. For reference, we report in Figure 4 middle panel (see also Supplementary Information S5 middle panel) the time evolution of the aGCN distribution of surface sites during a growth from an Oh. For the case of the growth from an Ih seed, the latter proceeds in a layer-by-layer fashion while always maintaining an icosahedral center. On the contrary, Oh (and the To) evolve either: 1) in structures with many (111), (100) and (211) facets and a strong anisotropy, or 2) undergo a kinetic rearrangements towards more favourable morphology (i.e. dIh for the case of the Oh or dDh for

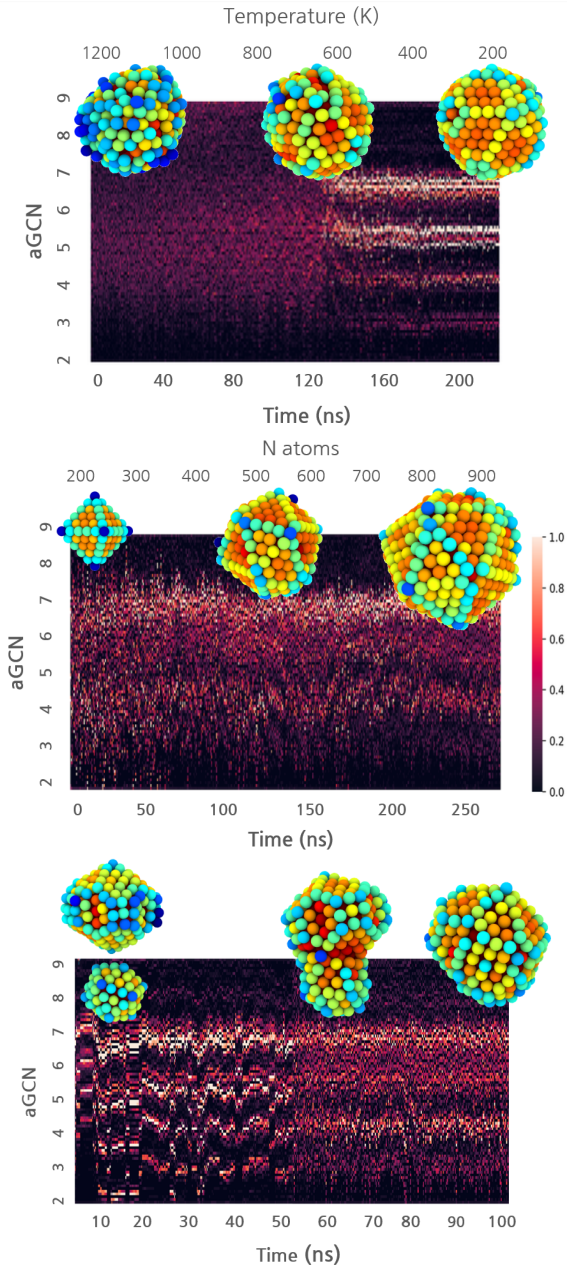


Fig. 4 An example of the different formation processes' kinetics represented through a heatmap. The aGCN occurrence is normalised and plotted against time for annealing, growth and coalescence. From left to right, the annealing of a cluster of 585 atoms, the growth of an Oh from 146 to 976 atoms and the coalescence of Cu₁₄₇ with Cu₄₃₄. Snapshots of the geometries are taken at relevant time steps. The atoms are coloured following the palette , where blue corresponds to an aGCN of 2 and red of 9.

the To). These solid-solid transitions are exemplified in Figure S5 middle left panel.

When looking at the aGCN distribution evolutions during coalescence, exemplified in Figure 4 lower panel (and S5 lower panel), we note three stages. Before the collision, the system is made up of two ordered individual nanoparticles; hence the aGCN occurrence for the whole system is, simply, the sum of the two. During the collision, the distribution of aGCN changes con-

siderably, indicative of a highly defected and disordered atomic arrangement. Conversely, the Cu-NPs can't be described neither as icosahedral, decahedral, or FCC cuts. This is also related to the rather pronounced anisotropy they display. The aGCN list and occurrence are very disordered and spread out, but still reminiscent of the initially preferred aGCN values, around 5 and 6.5, which corresponds to the sites in the initial structures which are not affected by the collision. We further highlight how the neck generated by the first moment of the collision is always highly coordinated. After the collision, atomic rearrangements at the surface succeed one another. This may also result in collective rearrangements (e.g. time=0.01 ns). Depending on the size of the system, the chosen observation time-scale either enables (small sizes) or forbids (large sizes) a complete rearrangements towards a defected but lesser anisotropic morphology.

Previously, it has been shown by numerical calculations^{54,62} and experiments^{39,45,63-66} that kinetic trapping in high-energy isomers take place even if the latter are quite unfavourable w.r.t. the global minimum at the same size. We further verified that the NEAS distribution of each metastable isomer is stable within the 10²-10³ ns timescale (see SI Section VI). As a final note, we remark that we have analysed NEAS distributions in Cu-nanoshapes obtained from three out-of-the-equilibrium formation routes representative of, e.g., inert-gas aggregation sources. Nevertheless, the proposed framework is transferable to estimate the NEAS distributions and hence the mass activity of any NP structure. Nanoparticles shapes can be predicted via global minimum searches algorithms, e.g. basin hopping^{67,68} or genetic optimization⁶⁹. Similarly, the probability to remain in a certain structural basin can be evaluated by accounting for energetic and entropic contributions^{70,71}.

3.3 Formation process, NP size, and resulting non equivalent site distributions

In Figure 5 we report the aGCN distributions for size-selected geometries sampled during the three formation processes under consideration.

For the case of annealing (top panel), the number and type of non-equivalent adsorption sites change drastically with the nanoparticle size. The aGCN distribution of solidify Cu₂₅₀ nanoparticles show peaks at around 5.5 and 7, suggesting an abundance of small (111)-like facets. Cu₅₈₅ solid structures display aGCN zeniths at around 5.5, 6.5 and 7.5, which correspond to a more ordered and faceted geometry delimited by (211), (100) and (111) facets. Finally, the aGCN distributions for Cu₇₁₉ and Cu₉₇₆ show a high occurrence of aGCN-sites between 8 and 9. This corresponds to an higher probability of finding the surface decorated by re-entrances, island, and steps. The occurrence of sites of this nature, in turn, appears to correlate with the NP size. Viceversa the relative abundance of low-coordinated sites is marked for small Cu-NPs (below 300 atoms) and anti-correlates with size.

For the case of atom-by-atom grown NPs, (three central panels) though the genome distribution is more uniform for smaller structures, it evolves towards specific signatures as the size grows.

Notwithstanding the structural diversity of the initial seed and final products, the aGCN distributions appears rather similar for the three sets of four independent trajectories considered. Regardless of the initial seed and final morphology we in fact observe peaks in the aGCN occurrence at values of around 7.5, corresponding to (111) facets, the most energetically favourable surface for FCC metals, and particularly so for the case of Cu. The number of sites corresponding to atoms at steps and re-entrances (generalized coordination above 7.5) is instead small. The occurrence of sites along (100) surfaces (aGCN = 6.67) and corresponding to vertex and adatoms (coordination less than 5) is also noticeable as a feature in the aGCN distributions. The formation of anisotropic shapes results in a lesser number of low-coordinated sites during the course of the growth.

Looking at the aGCN distributions of coalesced nanoparticles, Figure 5 lower panel, we observe they are homogeneously distributed for Cu-NPs of small sizes. Features of the original individual NPs are instead partly preserved in systems larger than 300 atoms. At the same time, in the neck region several steps and defects are formed leading to a large variety of atop generalised coordination values, namely above 7.5 or below 4.

3.4 Formation process, NP size, and NP activity

The heterogeneity of the type and occurrence of the adsorption sites of Cu NPs, which is related to their formation processes, leads to different size-trend of the NP catalytic properties. Under the assumption that structure-property relationship expressed in Eq. 4 contains all the needed ingredients in determining the activity of a NP, we are equipped to predict the catalytic activity of Cu-NPs obtained through diverse formation processes in the conversion of CO₂ into CH₄.

As a rule of thumb, because under-coordinated atoms are the most active according to the structure-property relationships in Equation 4, we expect that defects enhance the catalytic activity towards methane conversion. Using Equation 2, we calculate the current density and mass activity of Cu NPs catalysts driving the conversion of CO₂ in CH₄ at an applied potential close to V vs RHE = -1.1 V. We then report on the size-dependence of the current density for CO₂ conversion to CH₄ as a function of the formation process through which the Cu-NPs are synthesized. These estimates are reported in Figure 6. The MA prediction statistics are converged to a satisfactory degree according to a bootstrapping analysis (supplementary information, Section VII).

In agreement with the aGCN distributions reported in Fig. 5, respectively for annealing, atom-by-atom growth and coalescence, we observe the activity trend depends strongly on the formation-process and nanoparticle size. While the calculated current densities and MA* range between 0.1 and 1 mA/cm² and 5-11 mA/mg respectively, we observe that: (i) Annealed Cu-NPs display a decreasing mass activity with size, their MA moves from 9.6 mA/mg in NPs of 147 atoms down to 5.2 mA/mg in NPs of 976 atoms. Indeed the current densities, regardless of size, span only between 0.1 and 0.2 mA/cm². (ii) Atom-by-atom grown NPs find, on average, an enhancement of their MA activity from 5.7 to 11.2 mA/mg, although strong oscillations occur, mainly due

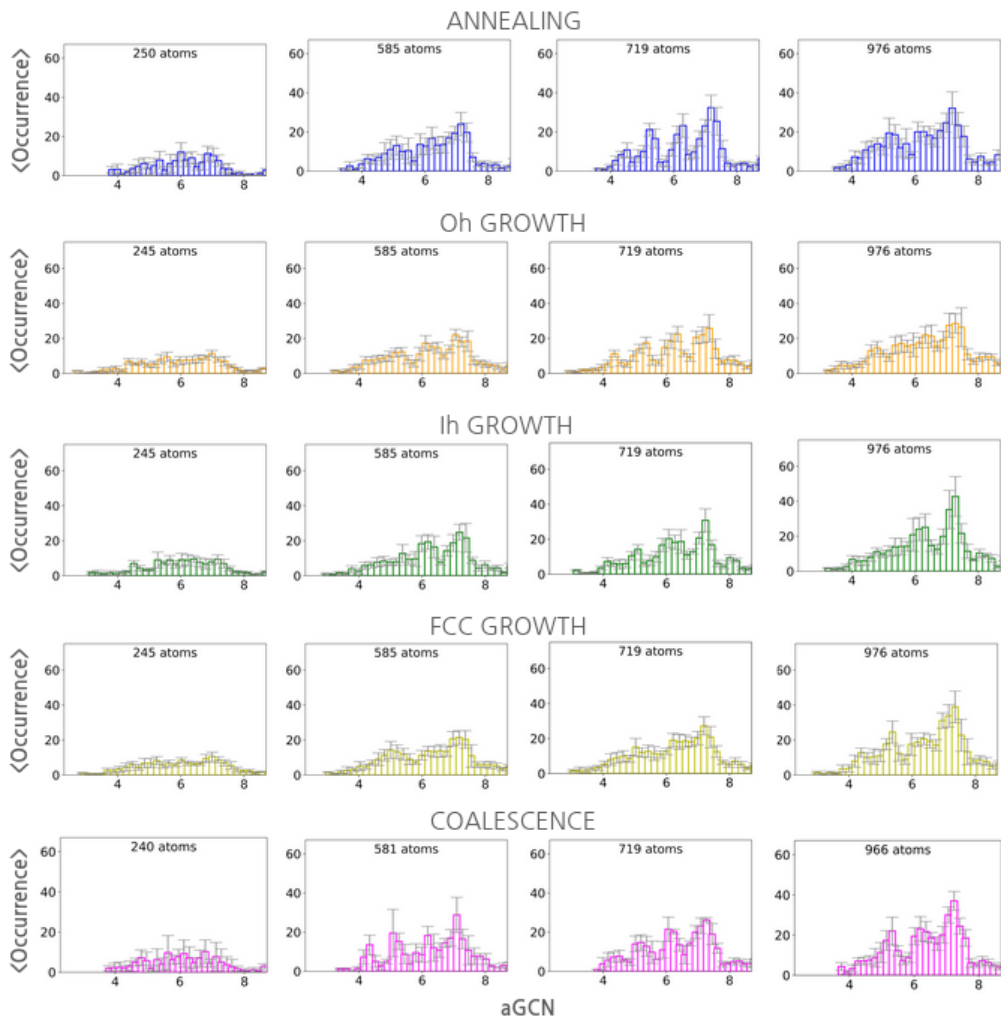


Fig. 5 Averaged aGCN distributions for the set of final configurations observed when analyzing the annealing (upper panel), growth with different seed (panels two to five), and coalescence (lower panel) of Cu-NPs of different sizes.

to local surface rearrangements, and depending on the initial core. The current density magnitude positively correlates with nanoparticle size.; (iii) Coalesced NPs show a rather smooth but non-monotonic size-behaviour with their MA lying in the 8.1-10.3 mA/mg range. Their current densities at -1.1V vs RHE extend between -0.2 and -0.3, with the largest observed for the case of NPs of largest sizes. Between 200-600 atoms, the NPs with the largest mass activity are obtained from the coalescence of smaller NPs of 1 nm in diameter, with a gain of about 4 mA/mg NP with respect to NPs formed following a different processes. On the other end, Cu-NPs grown atom-by-atom become the most active at larger sizes, above 800 atoms.

In comparison with experimental data (beyond the ones used to fit the constant in Eq.2), we observe that the magnitude of current densities, 0.1-1 mA/cm², finds a good agreement with the one for CO₂ conversion to CH₄ reported in the literature for Cu catalysts, e.g., Cu foils and pristine surfaces¹.

Beside such quantitative agreement, other comparisons are not trivial because of the complex multi-products outcome in the reduction of CO₂. We note hydrogen evolution reaction (HER) is

known to compete with CO₂ reduction, with an increased activity for the latter observed only in catalysts of larger sizes than the ones here considered for the case of spherical nanoparticles.²³ Also, it is has been reported that (100) facets are less active than (111) for CH₄^{1,72} at U* vs RHE = 1.1V, which is the applied potential we use in Eq. 2. We note that towards accounting in Eq. 2 for the transformation of CO₂ into molecules other than CH₄ one can introduce one or multiple terms simulating the competition between different products. This may then enable to recover the above experimental observations.

By the same token in the literature it is reported that an increased tendency to form either CH₄ or C₂ products is due to the presence of active sites at steps close to facets (steps at (111) facets for methane, steps at (100) for C₂ products). These adsorption sites display a lower coordination than the one of atoms on (111) surfaces.^{1,72-75} Employing a volcano relationship between coordination and activity, which peaks at low aGCN, values may in turn also help in the design of Cu catalysts active for CO₂ conversion into C₂ products.

As final consideration, let us stress that the workflow in Fig.

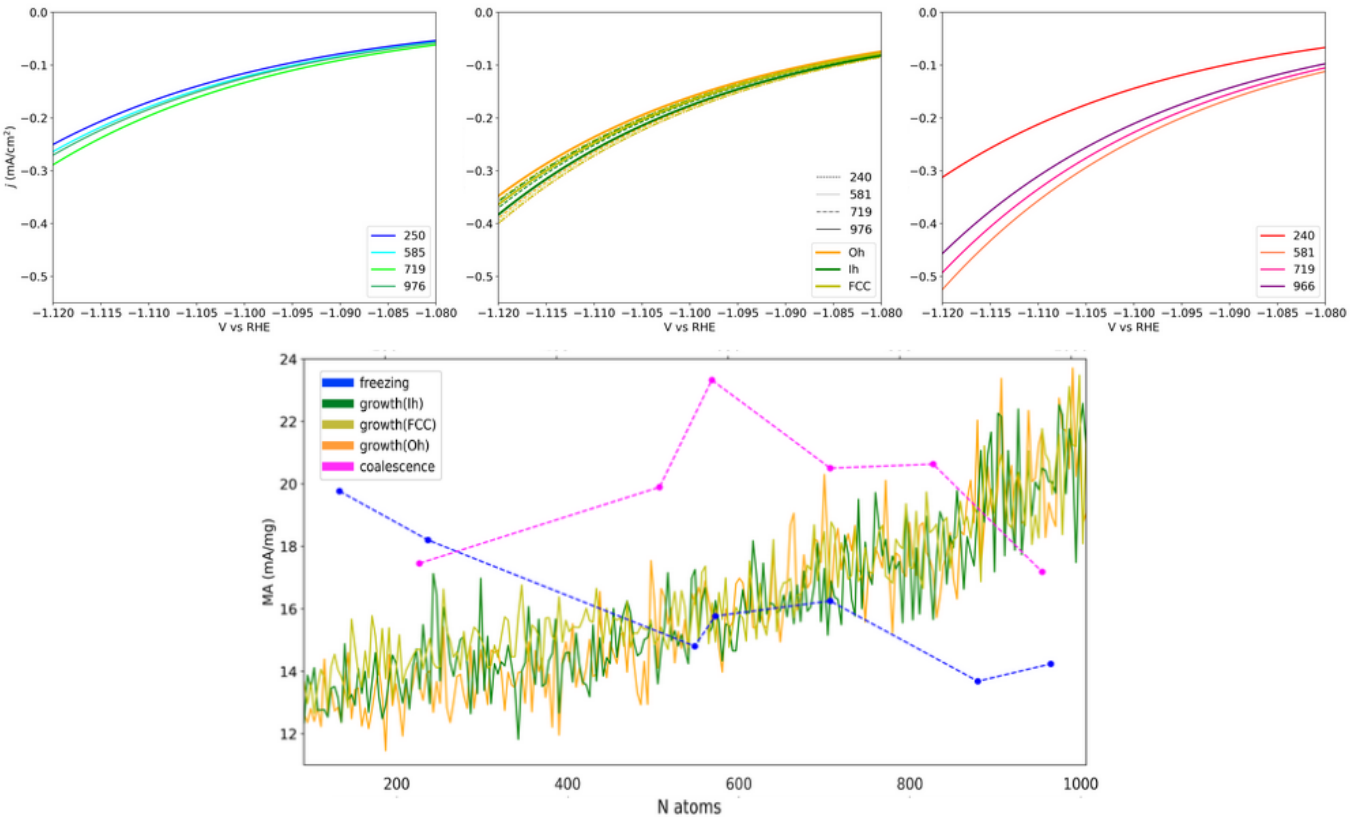


Fig. 6 Upper row displays the current densities for $\text{CO}_2 \rightarrow \text{CH}_4$ conversion around $U=-1.1\text{V}$ vs RHE for Cu-NPs of different sizes. Left panel represents Cu-NPs from the annealing of a liquid droplet (top-left, blue), the middle panel from the atom-by-atom growth (top-centre, orange is for Oh seed, dark-green for Ih and olive for FCC initial seeds, respectively). Right panel reports coalesced Cu-NPs (top-right, pink). The bottom row shows the mass activity for CH_4 conversion at $U=-1.1\text{V}$ vs RHE plotted versus the size of Cu-NPs sampled over the three formation processes (blue annealing; green-orange growth; magenta coalescence). Dashed lines for annealing and coalescence are a guide to an eye. Growth lines look continuous as data are available at each size. Those data points are filtered via a Gaussian Kernel for readability. The non-filtered data display oscillations within $\pm 5\text{mA}/\text{mg}$, due to the instantaneous appearance/disappearance of surface defects with aGCN values close to the structural relationship's maximum (Fig. 2).

1 is fully general and that corrections and modifications to the structure-property relationship in Eq. 4 can be made, although this is out of the aim of this work. In turn, other, more accurate and complex, functions of structural/electronic descriptors accounting for, e.g., multiple elements in the catalyst, and also found, e.g., by means of statistical learning models beyond linear ones, may improve and generalize the framework here discussed. Also, the reaction free energy estimator can encode a potential-dependent term, so to predict consistently CO_2RR activity over a range of applied potential.⁷⁶

4 Conclusions

We systematically analyse Cu nanoparticles' structural properties grown following different routes representative of inert gas aggregation sources. We demonstrate how the chosen formation method could impact, and potentially tailor, the activity of Cu-NPs for the conversion of CO_2 into methane. We show that the formation process leads to different shape distributions, even in the same size range. Each characteristic shape distribution influences the kind and abundance of non-equivalent adsorption sites, distinguished and catalogued according to their atop generalised coordination number. We observe that the occurrence of highly co-

ordinated sites mainly correlates with the nanoparticle's size. Cu-NPs are likely to be delimited by (111) facets, especially during an atom-by-atom growth, as expected by surface energy considerations. On the other hand, the abundance of low-coordinated sites depends strongly on the NP's formation route. Employing an extended version of the nanoCHE model -a multi-scale approach bridging structural features to activity-, we observe and explain non-trivial size-trends during the CO_2 into CH_4 reaction. Our work proposes the manufacturing process of Cu-NPs as a feasible tool to tailor the non-equivalent adsorption sites distribution and hence the activity of copper nanoparticles. Nonetheless, we are aware that a better quantitative prediction requires experimentally validated structure-property relationships. The latter will account for bias-dependent barriers and multiple products from the electrochemical reduction of CO_2 . We hope to stimulate the community's interest in providing a full structural characterisation of Cu-NPs before and after CO_2 conversion of differently grown samples.

5 Data Availability

The LoDiS package to run MD simulation is available at: <https://github.com/kcl-tscm/LoDiS>. The Python3 code used to esti-

mate the mass activity for CO₂ into CH₄ is available at: https://github.com/kcl-tscm/CO2RR_Cu_NPs. Example snapshots sampled during the trajectories discussed in this report are also available at the same page.

6 Acknowledgement

The authors thank Raffaella Buonsanti for fruitful discussions. The authors further thank Alejandro Santana-Bonilla, Matteo Tiberi, Robert Jones, Laia Delgado, Raphael Pinto-Miles, and Armand Auquier for their suggestions in developing the computational tools within the LoDiS software project, supported by the IAA-EPSRC grant EP/R511559/1. FB thanks the EPSRC “Towards an Understanding of Catalysis on Nanoalloys” Critical Mass Grant (EP/J010812/1) as does KR (ER/M506357/1). All the authors are grateful to the Royal Society, grant RG 120207, and the IT support offered by the NMS Faculty at King’s College London.

Notes and references

- 1 C. Hahn, T. Hatsukade, Y.-G. Kim, A. Vailionis, J. Baricuatro, D. Higgins, S. Nitopi, M. Soriaga and T. Jaramillo, *Proceedings of the National Academy of Sciences of the United States of America*, 2017, **114**, 5918–5923.
- 2 X. Kang, Q. Zhu, X. Sun, J. Hu, J. Zhang, Z. Liu and B. Han, *Chem. Sci.*, 2016, **7**, 266–273.
- 3 S. Min, X. Yang, A.-Y. Lu, C. Tseng, M. Hedhili, L. Li and K. Huang, *Nano Energy*, 2016, **27**, 121–129.
- 4 D. Raciti, K. J. Livi and C. Wang, *Nano Letters*, 2015, **15**, 6829–6835.
- 5 S. Nitopi, E. Bertheussen, S. Scott, X. Liu, A. Engstfeld, S. Horch, B. Seger, I. Stephens, K. Chan, C. Hahn, J. Nørskov, T. Jaramillo and I. Chorkendorff, *Chemical reviews*, 2019, **119**, 7610–7672.
- 6 K. Manthiram, B. Beberwyck and A. Alivisatos, *Journal of the American Chemical Society*, 2014, **136**, 13319–13325.
- 7 S. Roberts, K. Kuhl and A. Nilsson, *Angewandte Chemie International Edition*, 2015, **54**, 5268–5271.
- 8 D. Ren, Y. Deng, A. Handoko, C. Chen, S. Malkhandi and B. Yeo, *ACS Catalysis*, 2015, **5**, 2814–2821.
- 9 K. Kuhl, E. Cave, D. Abram and T. Jaramillo, *Energy Environmental Science*, 2012.
- 10 H. Mistry, A. S. Varela Gasque, C. Bonifacio, I. Zegkinoglou, I. Sinev, Y.-W. Choi, K. Kisslinger, E. Stach, J. Yang, P. Strasser and B. Roldan Cuenya, *Nature Communications*, 2016, **7**, 12123.
- 11 C. Li and M. Kanan, *Journal of the American Chemical Society*, 2012, **134**, 7231–4.
- 12 J. Vickers, D. Alfonso and D. Kauffman, *Energy Technology*, 2017, **5**, 775–795.
- 13 A. Peterson, F. Abild-Pedersen, F. Studt, J. Rossmeisl and J. Nørskov, *Energy Environmental Science*, 2010, **3**, 1311–1315.
- 14 Z. Zhao, Z. Chen, X. Zhang and G. Lu, *The Journal of Physical Chemistry C*, 2016, **120**, 28125–28130.
- 15 W. Durand, A. Peterson, F. Studt, F. Abild-Pedersen and J. Nørskov, *Surface Science*, 2011, **605**, 1354–1359.
- 16 X. Zhang, L. Jin-Xun, B. Zijlstra, I. Filot, Z.-Y. Zhou, S. Sun and E. Hensen, *Nano Energy*, 2017, **43**, 200–209.
- 17 W. Tang, A. Peterson, A. S. Varela Gasque, Z. Jovanov, L. Bech, W. Durand, S. Dahl, J. Nørskov and I. Chorkendorff, *Physical chemistry chemical physics : PCCP*, 2011, **14**, 76–81.
- 18 Y. Hori, R. Takahashi, Y. Yoshinami and A. Murata, *Journal of Physical Chemistry B*, 1997, **101**, 7075–7081.
- 19 Y. Hori, A. Murata and R. Takahashi, *J. Chem. Soc., Faraday Trans. 1*, 1989, **85**, 2309–2326.
- 20 J. Qiao, Y. Liu, F. Hong and J. Zhang, *Chem. Soc. Rev.*, 2014, **43**, 631–675.
- 21 R. M. Arán Ais, D. Gao and B. Roldan Cuenya, *Accounts of Chemical Research*, 2018, **51**, 2906–2917.
- 22 D. Gao, R. M. Arán Ais, H. S. Jeon and B. Roldan Cuenya, *Nat. Cat.*, 2019, **2**, 198–210.
- 23 R. Reske, H. Mistry, F. Behafarid, B. RoldanCuenya and P. Strasser, *Jour. Amer. Chem. Soc*, 2014, **136**, 6978–6986.
- 24 Y. Kang, C. Chen, J. Snyder, N. Marković, P. Yang and V. Stamenkovic, *Science*, 2014, **343**, 1339–1343.
- 25 X. Liu, J. Xiao, H. Peng, X. Hong, K. Chan and J. K. Nørskov, *Nature Communications*, 2017, **8**, 15438.
- 26 C. Cui, L. Gan, M. Heggen, S. Rudi and P. Strasser, *Nature Materials*, 2013, **12**, 765–711.
- 27 F. Calle-Vallejo, J. M. García-Lastra, P. Sautet and D. Lofreda, *Angewandte Chemie (International ed. in English)*, 2014, 8316–8319.
- 28 F. Calle-Vallejo, J. Tymoczko, V. Čolić, H. Vu, M. Pohl, K. Morgenstern, D. Loffreda, P. Sautet, W. Schuhmann and A. Bandarenka, *Science*, 2015, **350**, 185–189.
- 29 K. Rossi, G. Asara and F. Baletto, *ChemPhysChem*, 2019, **20**, 3037–3044.
- 30 K. Rossi, G. Asara and F. Baletto, *Physical Chemistry Chemical Physics*, 2019, **21**, 4888–4898.
- 31 K. Reuter, C. Plaisance, H. Oberhofer and M. Andersen, *J. Chem. Phys.*, 2017, **146**, 040901.
- 32 K. Kalz, M. Krahnert, R. Dvoyashkin, R. Dittmeyer, R. Glaser, U. Krewer, K. Reuter and J. a. Grunwaldt, 2017, **9**, 17–29.
- 33 D. Foster, R. Ferrando and R. Palmer, *Nature Communications*, 2018, **9**, 1323.
- 34 H. Zhai and A. Alexandrova, *ACS Catalysis*, 2017, **7**, 1905–1911.
- 35 Z. B. Zhang, Z. and A. Alexandrova, *Acc. Chem. Res.* 2020, **2020**, **53**, 447–458.
- 36 H. Haberland, M. Karrais and M. Mall, *Zeitschrift für Physik D Atoms, Molecules and Clusters*, 1991.
- 37 K. Wegner, P. Piseri, H. V. Tafreshi and P. Milani, *Cluster beam deposition: A tool for nanoscale science and technology*, 2006.
- 38 H. Haberland, *Gas-Phase Synthesis of Nanoparticles*, 2017.
- 39 A. A. Turkin, M. V. Dutka, Y. T. Pei, D. I. Vainshtein and J. T. M. De Hosson, *Journal of Applied Physics*, 2012.
- 40 K. Rossi, G. G. Asara and F. Baletto, *ACS Catalysis*, 2020, **10**, 3911–3920.
- 41 F. Baletto, R. Ferrando, A. Fortunelli, F. Montalenti and C. Mottet, *Journal of Chemical Physics*, 2002.

- 42 A. L. Garden, A. Pedersen and H. Jónsson, *Nanoscale*, 2018.
- 43 S. G. Lambie, G. R. Weal, C. E. Blackmore, R. E. Palmer and A. L. Garden, *Nanoscale Advances*, 2019.
- 44 J. M. Rahm and P. Erhart, *Nano Letters*, 2017.
- 45 D. Nelli, A. Krishnadas, R. Ferrando and C. Minnai, *Journal of Physical Chemistry C*, 2020.
- 46 L. D. Marks and L. Peng, *Nanoparticle shape, thermodynamics and kinetics*, 2016.
- 47 https://github.com/orgs/kcl_tscm/teams/lodis, 2019.
- 48 D. Frenkel, B. Smit, J. Tobochnik, S. McKay and W. Christian, *Computers in Physics*, 1997, **11**, 351.
- 49 O. A. Villarreal, Marcos A. Oviedo and E. P. M. Leiva, *J. Chem. Theory Comput.*, 2012, **8**, 1744–1749.
- 50 F. Baletto and R. Ferrando, *Rev. Mod. Phys.*, 2005, **77**, 371–423.
- 51 J. Nørskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. Kitchin, T. Bligaard and H. Jonsson, *Journal of Physical Chemistry B*, 2004, **108**, 17886–17892.
- 52 D. Esrafilzadeh, A. Zavabeti, A. R. Jalili, P. Atkin, J. Choi, B. Carey, R. Brkliča, A. O'Mullane, M. Dickey, D. Officer, D. MacFarlane, T. Daeneke and K. Kalantar-zadeh, *Nature Communications*, 2019, **10**, 865.
- 53 E. García-Suárez and A. Riisager, in *Sustainable Catalysis: Energy-Efficient Reactions and Applications, Introduction to Room-Temperature Catalysis*, 2018.
- 54 K. Rossi, L. Pavan, Y. Soon and F. Baletto, *European Journal of Physics B*, 2018, **91**, 1–8.
- 55 M. Evans and M. Polanyi, *Transactions of the Faraday Society*, 1937, **33**, 448–452.
- 56 F. Calle-Vallejo and M. T. M. Koper, *ACS Catalysis*, 2017, **7**, 7346–7351.
- 57 A. Peterson and J. Nørskov, *The Journal of Physical Chemistry Letters*, 2012, **3**, 251–258.
- 58 J. Davis, F. Baletto and R. Johnston, *Journal of Physical Chemistry C*, 2015, **119**, 9703–9709.
- 59 K. Rossi, L. Bartok-Partay, G. Csanyi and F. Baletto, *Scientific Reports*, 2018, **8**, 1–9.
- 60 F. Baletto, C. Mottet and R. Ferrando, *Phys. Rev. Lett.*, 2000, **84**, 5544.
- 61 D. Nelli, M. Cerbelaud, R. Ferrando and C. Minnai, *Nanoscale Advances*, 2021.
- 62 K. A. Fichthorn and T. Yan, *The Journal of Physical Chemistry C*, 2021.
- 63 D. Pearmain, S. J. Park, A. Abdela, R. E. Palmer and Z. Y. Li, *Nanoscale*, 2015.
- 64 Z. W. Wang and R. E. Palmer, *Physical Review Letters*, 2012.
- 65 E. Ringe, R. P. Van Duyne and L. D. Marks, *Journal of Physical Chemistry C*, 2013.
- 66 L. Martínez, M. Díaz, E. Román, M. Ruano, D. Llamosa P. and Y. Huttel, *Langmuir*, 2012.
- 67 T. V. Bogdan, D. J. Wales and F. Calvo, *Journal of Chemical Physics*, 2006.
- 68 G. G. Rondina and J. L. Da Silva, *Journal of Chemical Information and Modeling*, 2013.
- 69 S. Heiles and R. L. Johnston, *Global optimization of clusters using electronic structure methods*, 2013.
- 70 F. Calvo, J. P. Doye and D. J. Wales, *Chemical Physics Letters*, 2002.
- 71 E. Panizon and R. Ferrando, *Physical Review B - Condensed Matter and Materials Physics*, 2015.
- 72 A. Loijudice, P. Lobaccaro, E. Kamali, T. Thao, B. Huang, J. Ager and R. Buonsanti, *Angewandte Chemie (International ed. in English)*, 2016, **128**, 5789–5792.
- 73 O. Piqué, F. Vines, F. Illas and F. Calle-Vallejo, *ACS Catalysis*, 2020, in press.
- 74 C. Choi, S. Kwon, T. Cheng, M. Xu, P. Tieu, C. Lee, J. Cai, H. M. Lee, X. Pan, X. Duan, I. Goddard, William A. and Y. Huang, *Nature Catalysis*, 2020, in press.
- 75 K. Jiang, Y. Huang, G. Zeng, F. T. Toma, W. A. Goddard III and A. T. Bell, 2020, **5**, 1206–1214.
- 76 A. J. Garza, A. T. Bell and M. Head-Gordon, *ACS Catalysis*, 2018, **8**, 1490–1499.