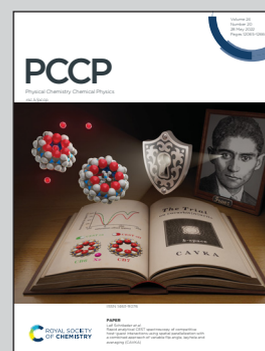


Showcasing work from the Theoretical and Computational Chemistry Lab, CNR-ICCOM, Pisa, Italy and the Department of Nanocatalysis of the J. Heyrovský Institute of Physical Chemistry of the Czech Academy of Sciences, Prague, Czech Republic.

Exploring the materials space in the smallest particle size range: from heterogeneous catalysis to electrocatalysis and photocatalysis

We review recent progress in the theory and characterization of the properties of ultras small subnanometer clusters. We show how atom-by-atom tunability of electronic structure and performance can in principle be achieved *via* a knowledge-based, physico-chemical approach, leading to rational design and potential applications in catalysis, photocatalysis, electrocatalysis, optical and chemical sensors.

As featured in:



See Alessandro Fortunelli, Štefan Vajda *et al.*, *Phys. Chem. Chem. Phys.*, 2022, **24**, 12083.



Cite this: *Phys. Chem. Chem. Phys.*,
2022, 24, 12083

Exploring the materials space in the smallest particle size range: from heterogeneous catalysis to electrocatalysis and photocatalysis

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Ultrasml clusters of subnanometer size can possess unique and even unexpected physical and chemical propensities which make them interesting in various fields of basic science and for potential applications, such as catalysis, photocatalysis, electrocatalysis, and optical and chemical sensors, just to name a few examples. These small particles often offer the tunability of their performance in an atom-by-atom fashion and an economic atom-efficient use of the metal loading. In this paper we review recent progress in the characterization and theory of well-defined subnanometer clusters in catalytic processes, and discuss their optical properties and stability, along with the potential of the size-selected clusters for the understanding of catalytic processes and for the development of new classes of catalysts.

Received 12th December 2021,
Accepted 13th April 2022

DOI: 10.1039/d1cp05677h

rsc.li/pccp

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Introduction

Designing heterogeneous catalytic systems¹ with the desired performance is one of the grand challenges of XXI-century chemistry.² This challenge has become especially urgent due to the need of transitioning from a linear to a circular economy,³ *i.e.*, to an economy based on a sustainable energy cycle, a circular chemicals production and (re-)use, and an



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environmentally-benign chemistry. Presently available catalysts do not satisfy these societal specifications in all important cases, despite century-long extensive explorations. We thus need to rapidly discover new catalysts/catalytic-sites enabling fundamental industrial processes (from small molecule activation to fine chemistry) under mild conditions, being less energy-consuming, and with a high activity and selectivity.²

In this context, we believe that a physical chemistry approach to monitoring catalytic reactions employing a combined, synergic experimental and theoretical approach is key to make real progress not only in understanding but also in controlling and rationally designing new catalysts. In addition, we also believe that the present-day challenges of delivering societally-complying chemical processes can be successfully addressed by exploiting the knowledge drawn from studying highly sophisticated systems, discovered, realized, and characterized in depth: this fundamental knowledge can then open the way to a physical-chemistry-informed rational design of novel nanocatalytic systems. Our focus in this review is on one class of these highly sophisticated systems, that of subnanometer supported cluster systems (reference to larger particles will be introduced as needed). Supported subnanometer clusters present several potential merits and advantages in the catalysis field. As we will see later, their hybrid character, intermediate between organometallic complexes and several-nanometer particles, makes their chemistry in many way unique, *i.e.*, different and tunable at a higher degree than in larger systems. Their economic atom-efficient use of the metal loading is also extremely appealing, especially in view of elemental scarcity and sustainable exploitation of natural resources. Finally, by creating materials in which the metal cluster component

is uniquely defined in both size and composition, we streamline the material phase diagram, thus boosting experimental and theoretical characterization, and catalytic testing (coincidence of spatially averaged and local techniques). However, despite these expectations and promises, the actual catalytic potential of ultra-small metal catalysts is presently very far from been exploited, with the lack of fundamental knowledge on structure and properties at this ultra-nano scale being a major reason hindering progress. Subnanometer clusters therefore represent an ideal playground in which a physical chemistry approach to monitoring catalytic reactions can make a breakthrough difference in deriving a fundamental understanding of catalytic processes and simultaneously developing new classes of catalysts. We present selected examples of experimental and theoretical characterization and reaction monitoring, and try to show how theory and experiment, sometimes in a synergistic thrust, sometimes in parallel, complement each other in achieving as atomistically detailed as possible knowledge on the chemical behavior of these systems. While the first era of intense cluster studies was entirely focused on free clusters in vacuum, it was the pioneering studies by Fayet and Wöste *et al* in the eighties⁴ and Cox and Kaldor *et al.* in the nineties⁵ who built the first bridges towards the interrogation of clusters deposited on surfaces, the first steps in establishing the basics for the expansion into a wide spectrum of research fields and applications of size-selected clusters.

Indeed, this shift of interest from gas-phase to supported small clusters is actually a *deja vu*. The first pioneering experimental studies of subnanometer metal catalysts in fact dealt with clusters deposited on porous silica or alumina, date back to the years 1970 and 1980, and were conducted by Sinfelt and coworkers at Exxon.⁶ In these studies, empirical evidence of the potentialities of the field was reported to the best of our knowledge for the first time. Later, these studies were confirmed and the role of subnanometer metal clusters in important reactions was unambiguously demonstrated by EXAFS data showing that in active and efficient nano-catalysts the coordination number of the metals did not exceed 5–6.⁷

In catalysis, rational design is traditionally associated with “volcano curve” modeling.⁸ In this approach, the translation of the Sabatier principle⁹ combined with the Brønsted–Evans–Polanyi (BEP) relationships,^{10,11} one selects a physical quantity as a “descriptor”, and, assuming that all energy differences and barriers for the given process scale linearly with the descriptor, one searches for the optimal value of the descriptor giving a minimum in the effective barrier (and a corresponding “volcano” maximum in reaction rate).⁸ Unfortunately, the reaction mechanism of typical industrially-relevant catalytic process is complicated, with many potential rate-determining steps, and can rarely be simplified *via* a single descriptor volcano curve. In such typical cases, detailed knowledge of atomistic static and dynamic structure of catalytic intermediates and their interconversion as derived from the studies reviewed here seems a pre-condition to make decisive progress to optimal design.

The plan of the review is as follows. We first describe the tools aimed at clarifying the static and dynamic structure of the



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ERA Chair and is the Head of the in 2019 established Department of Nanocatalysis. He has initiated the studies of subnanometer size- and composition selected clusters under realistic conditions of pressure and has developed methods to the testing and characterization of these catalysts under working conditions. His main focus is on the studies of subnanometer and nanometer size clusters in heterogenous catalytic reactions, in collaborations also in electro- and photocatalytic processes.



supported clusters (Section 2), as the basis to draw structure/catalytic-property relationships. Key to the practical use of these strongly under-coordinated species is their stability, addressed in Section 3. Also due to the importance of the field, an increasing number of studies deal with electrochemical rather than gas/solid heterogeneous catalysts, to which Section 4 is dedicated. With a perspective onto photocatalysis, we finally devote Section 5 to the optical properties of subnanometer substrate-stabilized clusters. Section 6 summarizes conclusions. Needless to say that this minireview is not intended to be nor can be all-inclusive. Rather, we deliberately chose to introduce topics such as photocatalysis, electrocatalysis, optical and chemical sensors that are only in their infancy or at a pioneering stage in the subnanometer-cluster field. One of our goals here is in fact to trigger interest on and promote developing new areas that have been scarcely or not at all explored so far, but that we believe are extremely promising in perspective.

Structure and reshaping

Structure

The electronic structure of clusters and the geometry of clusters on a support, including the accompanying cluster-support interaction, and the evolution of these properties as a function of reaction conditions are the factors determining catalytic activity. Clearly, the relative role played by these factors vary with the size of the clusters/particles, with the electronic structure of small clusters especially affected by cluster-support interactions, and adjustable through atomic control of the cluster size and of the hybridization between the electronic states of the adsorbed reactant molecules and of the cluster in order to realize a quantum-controlled catalyst.^{12–15} Among the various experimental methods to characterize the structure and composition of metal clusters we will first concentrate on microscopy (both electron and tunnel), X-ray and mass spectroscopy as *ex situ* techniques (Section 2.1), and environmental and scanning microscopy, synchrotron-based, and probe spectroscopy as *in situ* techniques (Section 2.2).

STEM

Aberration-corrected electron microscopy, especially aberration-corrected high-resolution scanning tunnel electron microscope (AC-HR-STEM),^{16,17} has made important contributions to characterizing supported metal particles down to single atoms. However, accurately determining the structure of supported subnanometer metal clusters in practical catalysts has been a grand challenge. Due to electron-beam induced effects, it is a non-trivial task to keep the atoms and clusters in their pristine configurations and thus provide useful information on the “intact structure” of supported metal atoms and clusters. Electron-beam induced effects and strategies to alleviate them were discussed in ref. 18 and 19.

A general view of the characterization of metallic alloy nanoparticles in catalysis using aberration-corrected STEM at atomic resolution is presented in ref. 20, where a clear

discussion is provided of the atomic distribution in the metallic nanoparticles affecting the catalytic activity of the nanoalloys, and the capability of high-angle annular darkfield scanning transmission electron microscopy (HAADF-STEM) methods to clarify these features. Down to subnanometer clusters, one of the first applications of AC-HR-STEM with direct atomic-resolution imaging of calcined Au₂₄Pd₁ clusters supported on multiwall carbon nanotubes using aberration-corrected STEM was given.²¹ Theoretical calculations supported the experimental evidence of Au cage structures surrounding the core Pd atom with possible implications for catalytic activity as Pd may act as electron promoter to the surrounding Au atoms when they are involved in catalytic reactions.

Fully exposed Pt₃ clusters fabricated on a defective nanodiamond at graphene (ND@G) by the assistance of atomically dispersed Sn promoters were characterized by HAADF-STEM. Results indicate that the size and structure of Pt species are related to the geometric partitioning effect of Sn atoms, indicating that mono-dispersed Sn species facilitate the formation of atomically dispersed Pt clusters. When the atomic ratio Sn/Pt was greater than 1.7, almost all Pt atoms existed in the form of atomically dispersed Pt clusters.²²

Fast dynamic STEM combined with a spatio-temporal image denoising algorithm was employed to unveil the dynamic structure of subnanometer Pt clusters anchored to a carbon support.²³ The analysis unexpectedly revealed for 30–50 atom Pt clusters a transition from amorphous-like quasi-2D arrangements at room temperature to ordered crystal-like (fcc or hcp) motifs.

For larger particles, size, three-dimensional shape, orientation and atomic arrangement of size-selected gold nanoclusters generated in the gas phase and soft-landed on an amorphous carbon substrate was determined using aberration-corrected STEM.²⁴ The structures of gold nanoclusters containing 309 ± 6 atoms were identified with either Ino-decahedral, cuboctahedral or icosahedral geometries and compared with theoretical modelling. In a further recent development, it has been possible to measure directly the population of competing structural motifs,²⁵ thus providing a quantity which is essential to validate/tune the accuracy of theoretical predictions.

The contribution of theory in this topic has so far concentrated on predicting the relative stability of different isomers and structural families for isolated medium-size and large particles with good success,^{26–30} although even more sophisticated simulations capturing the effects of electron beam on the cluster structure and thus precisely mimicking the experiment are in principle possible. The field of ultrasmall clusters presents advantages for theory as the limited size of these systems allows one to employ rigorous first-principles modeling, but also specific challenges connected with the need of performing systematic searches *via* stochastic methods,¹³ and technical issues such as how to accurately modeling quantum effects³¹ or cluster/support interaction energetics including *e.g.* dispersion terms.³² The important role of theory is also to single out energy terms which ultimately determine the cluster structure, such as many-body and directional effects leading *e.g.* to exotic



motifs. To provide simply one example, a new structural family of gold clusters in the size range between 23 and 42 atoms on the MgO(001) surface, the open pyramidal hollow cages, which has no counterpart in gas-phase clusters was proposed by DFT calculations.³³

Oxide systems can be investigated as well. For example, organic ligand-free, isolated tantalum oxide complexes (Ta_1) and small clusters ($Ta_{n>1}$) on flat silicate supports were prepared by ultra-high vacuum (UHV) techniques followed by oxidation in air. HAADF-STEM analysis of the resulting $Ta_{n>1}$ complexes showed narrow distribution with an size of 7 atoms per cluster, predominantly Ta^{5+} from X-ray Photoelectron Spectroscopy (XPS) analysis, and binding energies slightly shifted to higher values with respect to the bulk oxide values.³⁴ As another example, preparation of size-controlled tin oxide clusters in mesoporous silica using the dendrimer templating method, employing dendritic phenylazomethine with a tetraphenylmethane core (TPMG4) as the template molecule was reported in ref. 35. HAADF-STEM imaging and XPS analysis showed the presence of both tetravalent Sn(IV) and metastable divalent Sn(II) with a size-dependent composition.

Similar importance bear zeolite supports which are capable of hosting and stabilizing single atoms to well-defined ultra-small clusters and have been characterized by a variety of techniques,^{36–48} including zeolites in 2D⁴⁹ and 1D⁵⁰ form.

STM

Scanning Tunnel Microscopy (STM) has also been one of the most important tools to achieve information on cluster structures and catalysis.^{51–61} For this technique, a consolidated synergy between theory and experiment exists, grounded on various decades of combined studies *e.g.* in the proximal field of ultrathin oxide-on-metal systems.^{62–65} Functionalized STM tip allows for systematic atomic structure determination with STM of nanoparticles deposited on metal surfaces, with attention to be paid to tip-convolution effects as demonstrated in ref. 66 by a combined experimental/theoretical study of Co clusters (with average diameter 2.5 nm) deposited under ultra-high vacuum onto a thin insulating NaCl film on Au(111) (Fig. 1a).

Despite the challenges associated with the subnanometer size of the particles, several examples exist in the ultranano cluster field. The morphology of size-selected Pt_n ($n = 5–13$) clusters on a $CeO_2(111)$ surface using STM was investigated at room temperature.⁶⁷ Two-dimensional morphology of Pt_n clusters was preferable for $n = 5$ and a three-dimensional (3D) morphology for $n \geq 6$. The preference for a 3D tri-layer structure when $n \geq 10$ was observed. The interplay between intra-cluster Pt–Pt and extra-cluster Pt–O interactions (*i.e.* between the cluster and the $CeO_2(111)$ surface) was invoked to rationalize the size-dependent morphology of these $Pt_n/CeO_2(111)$ clusters.

Atomic-resolution images of size-selected clusters on surfaces enabling the identification of atomic alignments in the clusters on the surface were obtained in ref. 68 (Fig. 1b). The size-selected Pt_n ($n = 4, 7–10, 15$) clusters on $TiO_2(110)$ surfaces

were scanned using an ultrahigh-vacuum STM with a carbon nanotube tip. The clusters smaller than Pt_7 laid flat on the surface with a planar structure. However, a planar-to-three-dimensional transition occurred for Pt_8 and both Pt_8 and Pt_9 had two types of geometric structures.

Sub-molecular resolution in STM imaging was achieved on a 5 nm Ag_{374} nanoparticle monolayer-protected by *tert*-butyl benzene thiol.⁶⁹ The experimental data were compared with simulated topography images from DFT through a pattern recognition algorithm.

Tetrahedral (FCC segment) pyramid structure of Au_{20} clusters was confirmed using real-space, atomic-resolution images obtained with HAADF-STEM (Fig. 1c). The time-lapse HAADF-STEM imaging shows that the Au_{20} clusters continuously fluctuate between different structural configurations. The tetrahedral pyramid structure is one of the recurring isomers present in the time-lapse image sequences of the Au_{20} clusters although lower symmetry structures are more commonly observed.⁷⁰

In a pioneering experiment, variable-temperature STM was used to study the effect of kinetic cluster energy and rare-gas buffer layers on the deposition process of size-selected silver nanoclusters on a platinum(111) surface.⁷¹ At impact energies below 1 eV, the clusters landed non-destructively on the substrate, whereas at higher kinetic energies fragmentation and substrate damage were observed. Argon buffer layer on the platinum substrate efficiently dissipated the kinetic energy allowing for soft-landing at elevated impact energy. The opposite case of a hard landing also provides interesting potential for the nanostructuring of surfaces, allowing otherwise thermally unstable structures to be stabilized by the use of defects that were created in the violent deposition process.⁷²

Size-selected Pt cluster ions (size range 5–40 atoms) were deposited on a silicon(111) surface. The surface was characterized using STM under UHV conditions at 77 K. STM images of clusters with different sizes showed that the clusters were flattened and stuck to the surface with a chemical-bond akin to platinum silicide. Platinum atoms occupied preferentially the most reactive sites distributed within a diameter of ~ 2 nm on the silicon surface up to a cluster size of 20 atoms, while, above this size, the diameter of the cluster increased with size. The sticking probability of an incoming cluster ion on the surface increased with the cluster size and reached nearly unity at a size larger than 20 atoms.⁷³

X-ray

X-ray and specifically synchrotron techniques represent invaluable tools to furnish information on the structure of cluster catalysts. Small-angle X-ray scattering (SAXS), extended X-ray absorption fine structure (EXAFS), X-ray absorption near-edge structure spectroscopy (XANES), can give precise information about the size and shape of clusters as well as the chemical nature of their metallic components under operating conditions. Agglomeration of the deposited clusters leads to changes in the particle properties and structure that can be different from that in self-standing nanoparticles of similar sizes. *In situ* studies at



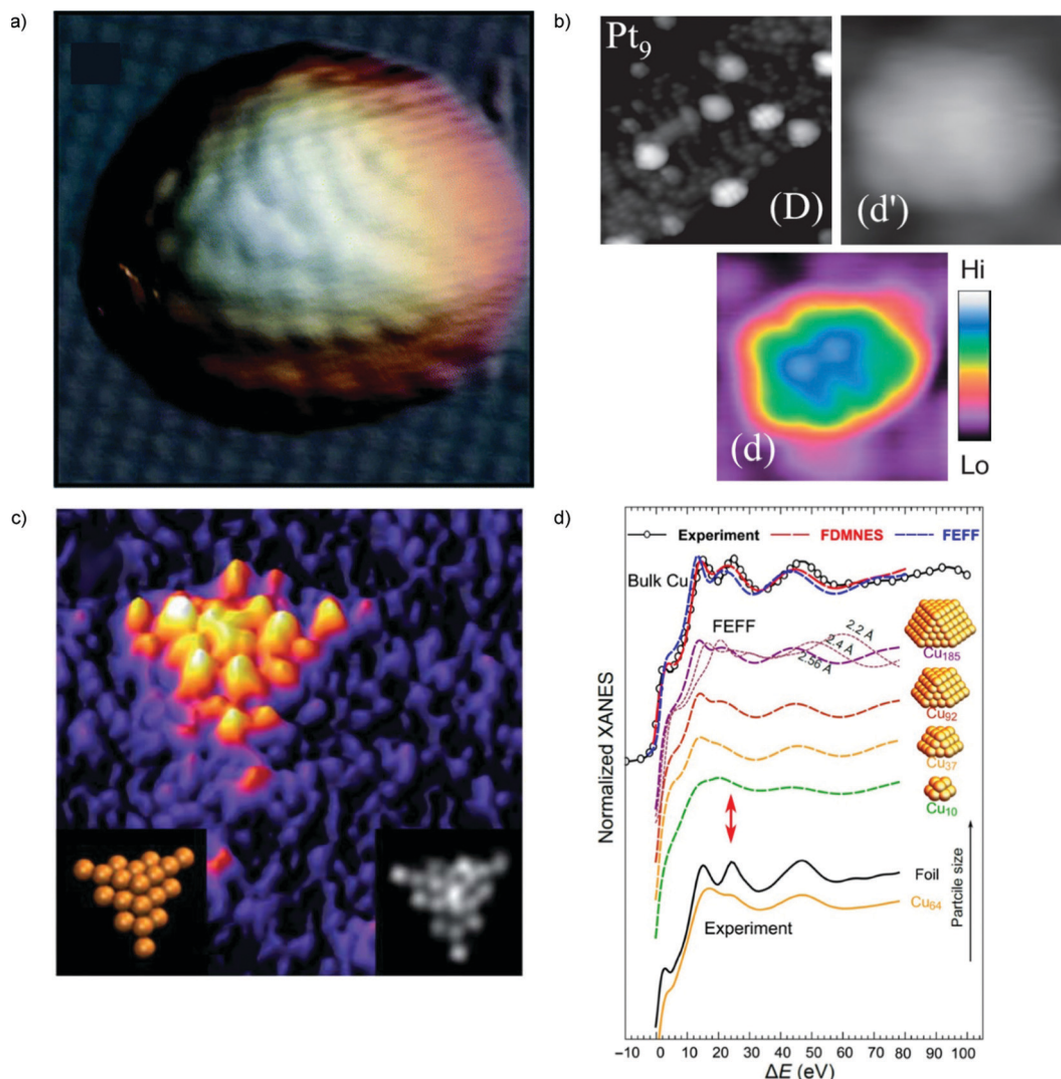


Fig. 1 (a) Atomic resolution STM topography image of Co cluster on NaCl–Au(111). Figure size is $7 \times 7 \text{ nm}^2$. (b) STM images of $\text{TiO}_2(110)$ surface after deposition of Pt_9^+ cluster ions. Image (D) is $20 \times 20 \text{ nm}^2$ and (d), (d') are $3.5 \times 3.5 \text{ nm}^2$ views of one cluster on the same surface. A color scale indicates heights for (d'). (c) HAADF-STEM image ($2.8 \times 2.8 \text{ nm}^2$) of a Au_{20} cluster (left inset shows the orientation of the Au_{20} cluster; the right inset is the simulated STEM image). (d) Experimental Cu K-edge XANES and theoretical spectra calculated with two *ab initio* XANES codes (FDMNES and FEFF) for bulk Cu and Cu clusters of different sizes. For the Cu_{185} cluster, the effect of interatomic distance is also demonstrated based on FEFF simulations. The red arrows mark the feature in XANES spectra that is very sensitive to particle size. (a) is adapted from ref. 66 with permission from the Royal Society of Chemistry, (b) is adapted from ref. 68 with the permission of AIP Publishing, (c) is adapted from ref. 70 with permission from the Royal Society of Chemistry, (d) is adapted with permission from ref. 74. Copyright 2018 American Chemical Society.

different length scales are required to monitor the formation of such complex materials. Combination of SAXS, XANES, *ab initio* simulations, and machine learning (artificial neural network) techniques was reported in ref. 74. Significant differences between the sizes of particle agglomerates, as probed by SAXS, and the sizes of locally ordered regions, as seen by XANES, were detected. These differences were reported as an evidence for the fractal, grape-cluster-like structure of the agglomerates. From the methodological point of view, it should be noted that this work reports one of the first applications of inverse modeling of experimental spectra (*e.g.*, XANES) to extract atomistic structures. This needs an efficient code to do direct modeling (*i.e.*, a code which, given an atomistic structure, predicts the expected spectrum in – say – less

than 1 hour cpu time), which is not the case of first-principles methods, at least so far (Fig. 1d). In multicomponent alloy catalysts, a combination of machine learning and multivariate curve resolution–alternating least square (MCR-ALS) analysis was used to determine the structure of the active site in multicomponent systems.⁷⁵

Another evidence that XANES data are sensitive to size and shape of metallic nanoclusters and neural network can be used to extract structural information from *in situ* XANES data is provided in ref. 76. Neural network assisted XANES method was used to extract information about the size, shape and interatomic distances in silver clusters, and to monitor their changes during the temperature-controlled particle aggregation



showing that particle shape is preserved during the aggregation of Ag clusters in ionic liquid.

A series of size-controlled Pd/hydroxyapatite (HAP) catalysts ranging from single sites (Pd clusters) to nanoparticles ($d_{\text{Pd}} \sim 30$ nm) was prepared and examined in ref. 77. In the subnanosize range, the Pd particles composed of 0D (Pd(IE)), 2D (0.5–2.0 wt% Pd) and 3D (0.5–5.0 wt% Pd) were observed, as identified by XRD, HRTEM, and EXAFS. As it was demonstrated by XPS, *in situ* diffuse reflectance infrared Fourier-transformed spectroscopy (DRIFTS) of CO, and O₂ temperature-programmed desorption (TPD), the electronic structure of the Pd surface can be tailored by varying the particle size. Higher ratios of surface Pd atoms have been found to be positively charged and to form PdOx/Pd interfaces on subnanosized particles.

The geometrical structure of mass-selected Au_n clusters deposited on a silicon substrate prepared by soft-landing conditions was investigated in ref. 78 using grazing incidence small angle X-ray scattering (GISAXS) in combination with a model-based approach. Here the link between structure and observed spectroscopic features is ensured by the use of a simplified (analytic) model, allowing one to obtain the probable morphological information.

Subnanometer Rh clusters derived from Rh-doped SrTiO₃ were produced and monitored by *in situ* XRD and XAFS measurements.⁷⁹ The *in situ* DRIFTS measurements were analyzed and suggested

that the high CO selectivity in CO₂ hydrogenation was attributed to the lower CO binding strength brought about by strong interactions (*e.g.*, charge transfer) between Rh atoms and the defected oxide support see Fig. 2.

From the preparation point of view, an approach to synthesize atomically dispersed uniform clusters *via* a cage-separated precursor preselection and pyrolysis strategy was reported in ref. 80 and illustrated in the production of uniform Ru₃ clusters stabilized by nitrogen species (Ru₃/CN) using well-defined Ru₃(CO)₁₂ separated as a precursor by suitable molecular-scale cages of zeolitic imidazolate frameworks (ZIFs). The structure of the Ru₃/CN clusters was characterized by HAADF-STEM and XAFS and their oxidation activity of 2-aminobenzyl alcohol investigated *via* first-principles calculations. Zeolites,^{36–48} which have been pioneeringly explored by Meurig as a support for subnanometer clusters *via* decomposition of carbonyl complex precursors,⁸¹ including zeolites in 2D⁴⁹ and 1D⁵⁰ form, represent another support which is of current considerable interest, despite the great challenges of their structural characterization.

Link with akin systems: ligand-protected metal clusters

The field of supported size-selected metal catalysts presents specifics and advantages that make them unique (see the introduction). However, structural characterization still present

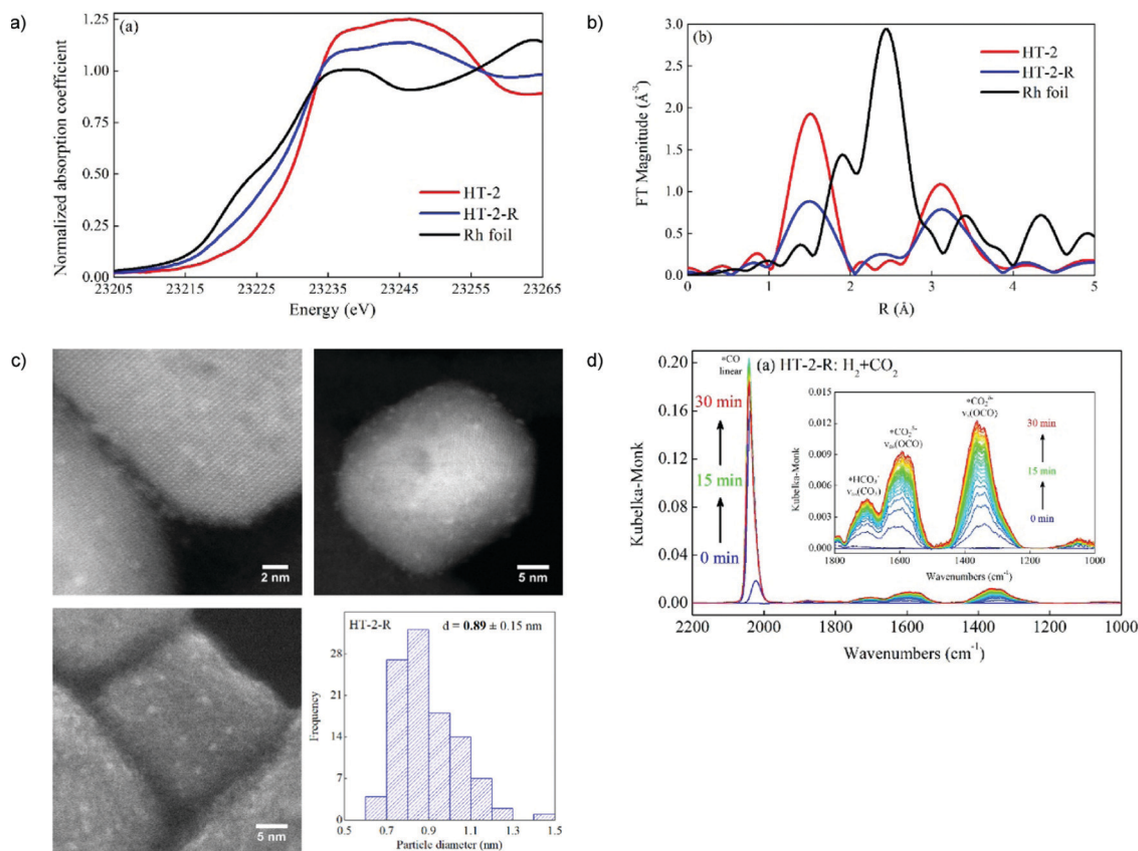


Fig. 2 *In situ* XAFS data of Rh foil, the HT-2 and HT-2-R samples: (a) XANES region, and (b) Fourier transform magnitudes of the k^2 -weighted EXAFS data. (c) Representative annular dark-field (ADF)-STEM images (top), and high-resolution SEM images (bottom left), and the size distribution of Rh nanoparticles (bottom right) of the HT-2-R sample. (d) *In situ* DRIFT spectra of CO₂ hydrogenation after H₂ reduction over HT-2-R. Adapted from ref. 79, Copyright 2018, with permission from Elsevier.



problems and challenges, as discussed above, and has not yet reached the targeted ideal, absolute precision. For these and also for synergistic reasons, we believe that the study of these materials would strongly benefit by a closer interaction and comparison with the techniques, results and materials obtained in the related sub-field of size- and composition-selected ligand-protected metal clusters (also called monolayer-protected metal clusters, MPC). The MPC field has been initiated more than 2 decades ago,⁸² but has witnessed an explosive growth more recently since the first compositionally unique crystalline samples were achieved and their atomistic structure determined *via* X-ray.⁸³ The crystallization and X-ray structure determination of a *p*-mercaptobenzoic acid (*p*-MBA)-protected gold nanoparticle, which comprises 102 gold atoms and 44 *p*-MBAs was reported in ref. 83. The central gold atoms were packed in a Marks decahedron, surrounded by additional layers of gold atoms in unanticipated geometries. The *p*-MBAs formed a rigid surface layer due to interaction not only with the gold but also with one another. Since then, nearly 200 distinct MPC compounds have been characterized. The appeal of these materials comes from the possibility of obtaining atomically precise stoichiometry and well-defined crystal structures. Knowledge of stoichiometry and static structure together with the use of advanced experimental and theoretical characterization tools has allowed researchers to achieve an extraordinary accuracy in fundamental studies cross-validating experiment and theory,⁸⁴ and an in-depth understanding of a wealth of physical phenomena, including *e.g.* optical absorption and plasmonics.⁸⁵ We believe that this knowledge can be usefully brought about into the subnanometer size- and composition-selected supported catalyst field, apart from the appealing possibility of using MPC both as precursors for preparing supported species⁸⁶ and to create hybrid systems combining supported and MPC species *e.g.* as internal standards in characterization studies (the latter possibility is so far unexplored, to the best of our knowledge). Hereafter and later in the optics section we thus briefly recall a few studies in the MPC field to trigger interest and possible scientific common investigations.

Size-dependent optical properties of ligand-protected gold nanoparticles have thus been studied extensively,⁸⁷ with the catalytic properties more recently also investigated.^{88,89} Ideal model catalysts should be atomically precise AuNPs with well-defined structures and uncoordinated surface sites as *in situ* active centers, however, the need to remove ligands can introduce uncertainties in both the structures and sizes of ligand-protected AuNPs for catalytic applications. The discovery of the highly stable tetrahedral (T_d) Au₂₀ cluster in the gas phase has inspired exploratory efforts toward its large-scale solution syntheses. In order to keep the tetrahedral structure of this gold pyramid, which has the bulk atomic arrangement with all its atoms on the surface, phosphine ligands were selected.⁹⁰

An all-amidinate-protected gold nanocluster [Au₂₈(Ph-form)₁₂](OTf)₂ (Ph-form = *N,N'*-diphenylformamidinate) (Au₂₈) has been synthesized and structurally resolved. Single crystal X-ray diffraction reveals that Au₂₈ has a compact Au₄@Au₂₄ tetrahedral core-shell structure of T symmetry, which is fully protected by

12 bridging formamidinate ligands. This cluster is quite robust as indicated by the fact that it can stay intact in solution at 80 °C for 6 days. A superatomic electron configuration of 1S²1P⁶2S²1D⁴ has been clarified by DFT computations, and the strong gold–ligand binding and geometric shell closure account for the superior stability of Au₂₈.⁹¹

Mass spectroscopy (MS), XPS, HR-TEM, FT-IR, UV-vis, and ultrafast spectroscopy techniques are commonly used in this field to unveil composition, morphology, and optical properties of the clusters, see *e.g.* ref. 92 It is notable that progresses in this field have enabled investigation of monolayer-protected clusters (MPC) up to fairly large sizes. For example, the properties of the ~300 kDa gold–hexanethiolate compound, extracted from the Faradaurate series of smaller and larger homologues were measured by mass spectrometry, optical spectroscopy, electron microscopy, X-ray scattering, and diffraction.⁹³ A monocrystalline metallic core of ~3.1 nm minimum dimension supports a clear plasmonic optical response, along with a diffuse exterior shell.

The recent rapid evolution of a multitude of characterization techniques including in realistic reaction environment and often in combination with state-of-the art computational capabilities, allow now tackle the structure–function relationships down at a single atomic level, thus paving the road to the atomic level understanding of and tailoring the properties of matter at the (sub)nanometer scale.

Dynamics, reshaping

Clearly, the atomistic structure of a catalyst is not a static feature, and cluster dynamics and restructuring (reshaping) is an important topic for a deeper understanding of the catalytic event. Especially for ultras-small systems, fluxional behavior has long been advanced as an important feature¹³ and evidence that dynamics plays a crucial role in many reactive processes and under many conditions has now accumulated. In the following, we will review this past experience, trying wherever possible to frame it into a set of basic concepts.

In particular, we would like to recall the concept of ligand/cluster/support catalytic complex. The theoretical understanding of catalysis by size-selected subnanometer (or ultranano) metal clusters deposited onto an oxide surface is based on three founding concepts: (1) the fluxional character of ultranano-clusters and the consequent need for systematically sampling the potential energy surface of the system, (2) the influence of the underlying support (including possible defects) in terms of geometric relationships, electrostatic and electronic interactions, and the possible active role of the support in the reaction mechanisms, (3) the evolution of the energetics, structure and dynamics of ligand adsorption as a function of coverage, including many-body interactions among ligands. These three concepts can be synthesized into a single one: the formation of ligand/cluster/support catalytically active complex. These species are aggregates involving the metal and support atoms and those of the intermediates produced by the catalytic process, and form *in situ* with a stoichiometry depending upon the given reaction mechanism and reaction conditions, and are the real actors of



the catalytic process. It is, therefore, crucial for assessing efficiency but also the robustness of the catalyst (*e.g.*, in terms of surface diffusion and cluster disaggregation) to single out which is the catalytically active species and which are its features.¹³

The first aspect, *i.e.*, a tool for systematically sampling the potential energy surface of ultrasmall catalytic systems using a first-principles theoretical approach to heterogeneous catalysis, was provided in ref. 94. There, propylene partial oxidation by Ag₃ supported on MgO(100) was studied theoretically. Computational sampling of the reaction path was achieved *via* a global search in the phase space of structures and stoichiometry combined with filtering, which takes into account the given experimental conditions (catalytically relevant temperature and reactant pressure), and corresponds to an incremental exploration of the disconnectivity diagram of the system. An alternative approach, dubbed global activity sampling method, that directly and globally optimizes the configuration of the supported cluster corresponding to the transition states for the catalytic reaction without requiring prior knowledge on stable cluster structures, was later reported with the example of C–H bond dissociation on Pt₇ and Pt₈ supported clusters and by an explicit sampling of cluster configurations at C–H dissociation transition states.⁹⁵

As for the second aspect, *i.e.*, the influence of support, in ref. 96, it was also shown how the presence of an oxide support induces extreme change of the potential energy landscape of the system with respect to the gas phase, favoring configurations that interact positively with the electrostatic field generated by the surface.⁹⁷

Finally, the reaction energy barriers for the various mechanisms are crucial in the competition between thermodynamically and kinetically favored reaction products. The co-adsorption of O₂ and other ligands on small metal clusters have strong cooperative effects (many-body or synergic effects).

STEM

The atom-precise and fully scalable synthesis of platinum clusters on a milligram scale from tiara-like platinum complexes with various ring numbers ($n = 5$ –13) was shown in ref. 98. Atomic resolution STEM showed, unexpectedly, that a Pt₁₀/Ketjenblack catalyst retained its original size features after 2 hours of hydrogenation reaction without obvious aggregation and decomposition. In contrast, in reactions under oxidative conditions, cluster degradation were more pronounced. Formation of single atoms was the mechanism leading to cluster disaggregation and reaggregation.

In another study, attention concentrated on the possibility of stabilizing metallic copper clusters under oxidizing reaction conditions by adjusting the atomicity of subnanometer copper clusters.⁹⁹ By a combination of the XPS and single molecule spectroscopy (SERS) data under the different atmospheres, it was concluded that Cu₅ clusters interact weakly with oxygen species and are easier to reduce and more difficult to oxidize than the Cu₈ analogues. In contrast, Cu₂₀ nanoparticles were less reducible, while they could be easily reoxidized at 25 °C. DFT simulations supported these findings.

The effect of the size of Cu clusters on the binding strengths of reactants and reaction intermediates, as well as the activation barriers, for the elementary reaction steps underlying CO₂ hydrogenation was studied,¹⁰⁰ typically finding a linear scaling relationship of reaction barriers with CO and O adsorption energies. Medium-sized Cu₁₉ clusters were predicted to exhibit the highest CO₂ hydrogenation activity, ascribed to a compromise between moderate CO₂ saturation coverage and a low CO₂ dissociation barrier, but all clusters below 55 Cu atoms were predicted to exhibit strong variations of CO and O adsorption energy while the reactivity of larger clusters and nanoparticles was associated with surface atoms with low coordination number.

STEM and XPS characterization before and after reaction to assess the stability of single Pt atoms and Pt clusters on a Si wafer support and the question of heterogeneity *versus* homogeneity in the catalytic process was addressed in ref. 101. XPS before and after catalysis showed changes in the oxidation states of the clusters, aggregation, and a loss of metal into solution. Characterization of the corresponding Pt clusters supported on TEM grids by means of HAADF-STEM, however, revealed stable clusters before and after exposure to reaction conditions, which was explained by the difference in the support between the silicate TEM grids and the wafers, resulting in different cluster–support interactions.

STM was used to study a Pt_{1–6}/Fe₃O₄ model catalyst exposed to CO, H₂, O₂, and mixtures thereof at 550 K. The proposed mechanism foresees that CO extracts lattice oxygen atoms at the cluster perimeter to form CO₂, creating holes in the metal oxide surface. H₂ and O₂ dissociate on the metal clusters and spill over onto the support, the former creating surface hydroxyl groups, ultimately leading to the desorption of water, while the latter reacting with Fe from the support to create new Fe₃O₄(001) islands. The presence of the Pt is crucial because it catalyzes reactions that already occur on the bare iron oxide surface, but only at higher temperatures¹⁰² (Fig. 7b).

The stability of pinned size-selected palladium clusters under reaction conditions (catalytic oxidation of cyclohexane) was investigated by STM measurement both before and after reaction. Pd_N clusters ($N = 10, 20, 55$ and 120) were found to be rather stable against sintering under reaction conditions, in a temperature range of 25–300 °C and at a pressure of 800 torr¹⁰³ (Fig. 3).

Sintering of supported Au nanoparticles is significantly different under reaction conditions compared to a vacuum environment.¹⁰⁴ The sintering in UHV of Au particles with a size of 2–5 nm occurs above 600 K as it was shown by STM. CO exposure has no apparent effect on the morphology of Au/TiO₂(110), whereas significant changes occur after exposure to O₂ or CO–O₂ at or near room temperature. The Au particle density was greatly reduced in CO–O₂, atmosphere because of sintering. Indeed, in ref. 96, it was theoretically demonstrated that the tendency of ultrasmall Au clusters to disaggregation was highly enhanced by the simultaneous adsorption of CO and O₂ species, an effect that was mitigated by alloying Au with Ag. An interesting comparison of the catalytic activities



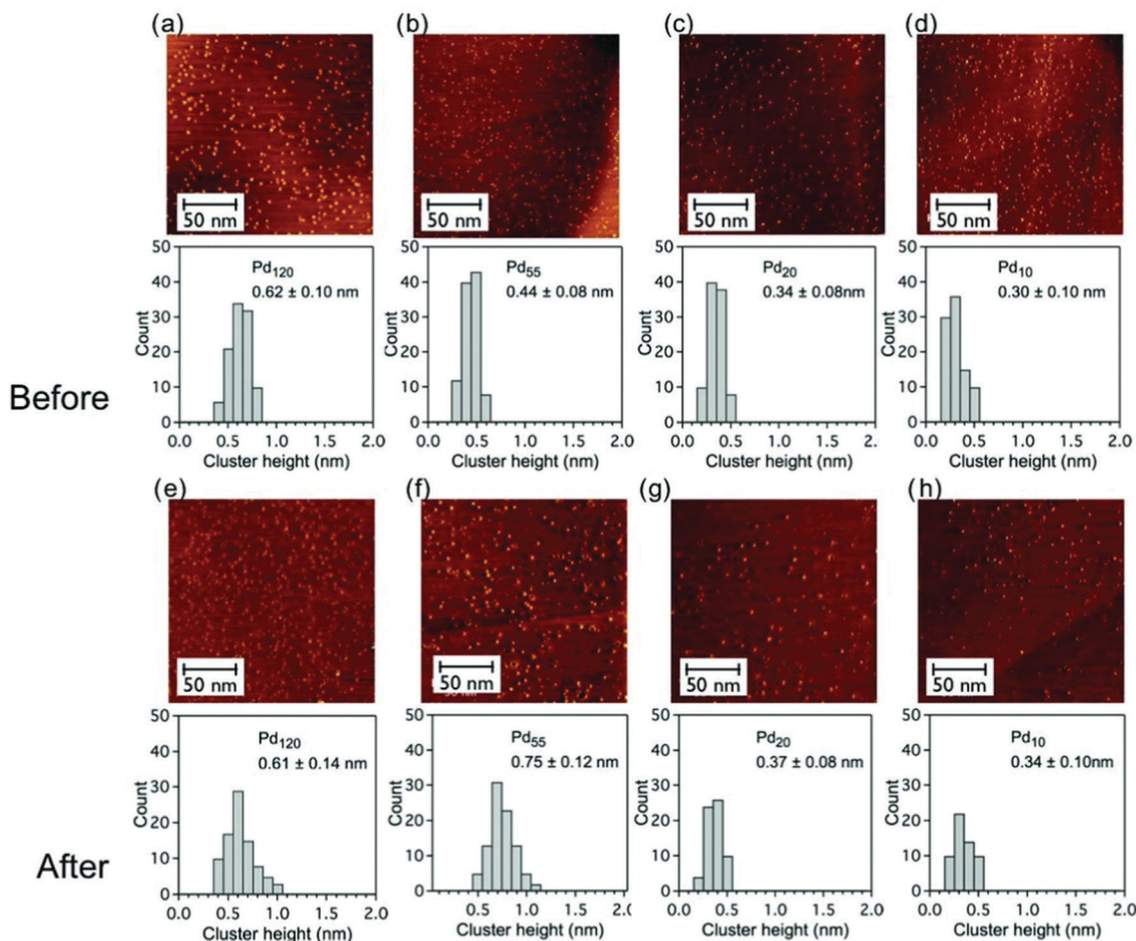


Fig. 3 Constant current STM images ($200 \times 200 \text{ nm}^2$, 0.4 nA, 0.4 V) of graphite-supported size-selected Pd_N clusters ($N = 120, 55, 20$ and 10 , low coverage) and histograms of the cluster height distributions before and after catalytic reaction at reactant ratio of $\text{C}_6\text{H}_{12}:\text{O}_2 = 1:10$; (a–d) as-deposited clusters before catalytic measurements and (e–h) after reaction catalytic measurements. The mean height of the cluster is given in each case. Adapted from ref. 103 with permission from Taylor & Francis.

reported for platinum group metals and Au catalysts of different sizes in CO oxidation was given in ref. 105. In terms of CO oxidation activity, Au NPs showed remarkable catalytic efficiency at low temperatures, superior to that of the single-atom Au catalyst. Single Pt atoms are superior to Pt clusters at high temperatures above $\sim 100^\circ\text{C}$. Nevertheless, although NPs show catalytic activity for CO oxidation, single Pt atoms do not show activity below 100°C .¹⁰⁵ In their pioneering works, Goodman and coworkers demonstrated the role of Au particle size or the role of gold in dispersing single Pd atoms in highly efficient CO oxidation^{104,106,107} or the acetylation of ethylene.¹⁰⁸

Thermal stability of atomic platinum clusters supported on $\text{Al}_2\text{O}_3/\text{SiO}_2/\text{Si}(100)$ as a function of the thickness of the alumina film and presence of hydrogen was studied by *in situ* GISAX in ref. 109. Extremely high thermal stability of Pt_{7-10} clusters *in vacuo* as well as in the presence of hydrogen was observed on $\text{SiO}_2/\text{Si}(100)$ coated with six cycles of Al_2O_3 film prepared by an atomic layer deposition technique.

Ligation and decomposition of 1,6-hexanedithiol on Cu clusters have been studied using TPD and XPS. Copper cluster

anions were size selected and soft landed into a frozen matrix of 1,6-hexanedithiol on HOPG maintained at 100 K. After warming up to 298 K, the sample was characterized by a combination of TPD and XPS. Two different binding motifs leading to different decomposition products were observed: the dangling motif is with one sulfur atom binding to a copper cluster, and the bidentate motif is with both sulfur atoms binding to a copper cluster. XPS measurements at varied temperatures confirmed the formation of dithiol ligated copper clusters through Cu–S bond formation, and the decomposition of them *via* C–S bond scission, consistently with TPD.¹¹⁰

Subnanometer ($d = 0.8 \pm 0.2 \text{ nm}$) gold particles homogeneously dispersed on amino-functionalized silica catalyze Glaser-type alkyne coupling, providing corresponding 1,3-diyne under mild conditions. The catalyst can be recycled at least five times, giving consistently high isolated yields and maintaining the size and distribution of gold clusters¹¹¹ (Fig. 5d).

Environmental TEM

Environmental TEM is a technique in which TEM characterization is performed under gaseous pressures up to realistic



values, currently typical upper limit being few tens torr but current efforts aim at reaching 1 bar.¹¹²

E-TEM has already provided information valuable in catalytic studies, and seems especially powerful when combined with concurrent synchrotron characterization techniques, and also in the future with a proper selection of adsorbate gases going beyond typical hydrogen, oxygen and CO molecules. For example, in one of the first *in situ* studies of nanoparticle dynamical reshaping, the combined use of *in situ* E-HRTEM, GISAXS, SEM, TPD, and theoretical simulations, led to the idea that the shape can depend on the reaction dynamics/conditions,¹¹³ in particular for model size-selected silver large nanoparticle catalysts deposited on amorphous alumina films in the partial oxidation of propylene (Fig. 5b). Notably, the calculated surface energies resulted *via* the Wulff construction in particle shapes resembling the shapes experimentally observed under realistic reaction conditions of partially oxidized silver clusters, implying that the flow of reactants corresponded to an intermediate

oxidation state of the surfaces, a concept now generalized as “steady-state Wulff construction under reaction conditions”.¹¹⁴

A mechanism of oxidation/reduction processes of Pd nanoparticles supported on various substrates, including CeO₂, g/d-Al₂O₃, ZrO₂ and SiO₂ was studied by means of environmental TEM.¹¹⁵ The observations showed that Pd nanoparticles were transformed to PdO during an exposure to ~1 mbar of O₂ at 400 °C *via* the formation of surface oxide, and subsequent nucleation and growth of PdO and planar front transformation (Fig. 4).

The evolution and stabilization of subnanometric Pt species confined in MCM-22 zeolite has been studied by *in situ* TEM under oxidation–reduction and reaction conditions in ref. 46. Dynamic and reversible transformation between single atoms, clusters and nanoparticles has been observed under CO + O₂ reaction conditions at different temperature. The subnanometric Pt clusters can be stabilized in MCM-22 crystallites during NO reduction with CO and H₂.

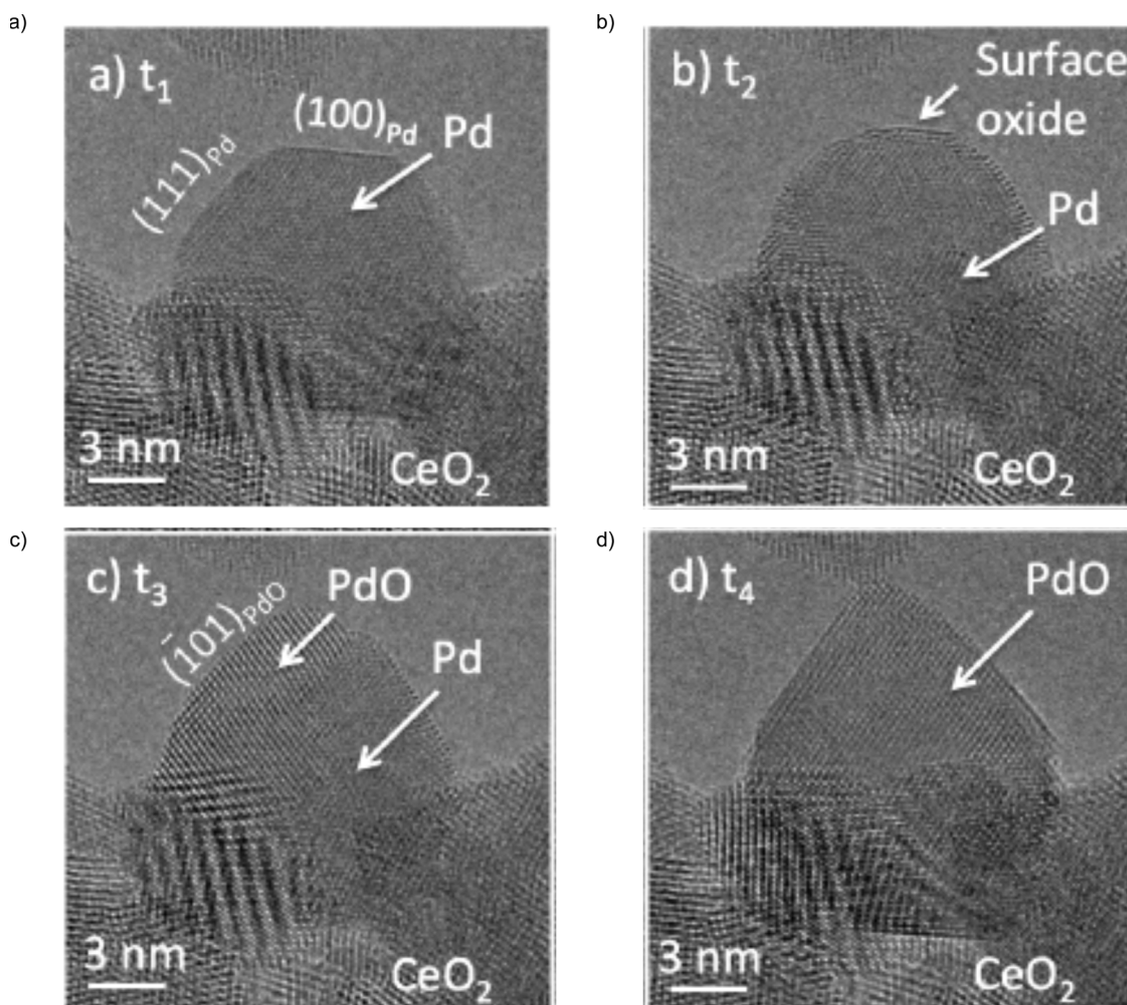


Fig. 4 High resolution ETEM observation of Pd to PdO transformation at oxygen partial pressure of ~1 mbar and temperature of 400 °C. (a) Initial Pd nanoparticle supported on CeO₂. (b) Transition period corresponding to the formation of surface oxide. (c) Transition period corresponding to growth PdO. (d) Final stage of transformation corresponding to the formation of PdO. Adapted with permission from ref. 115.



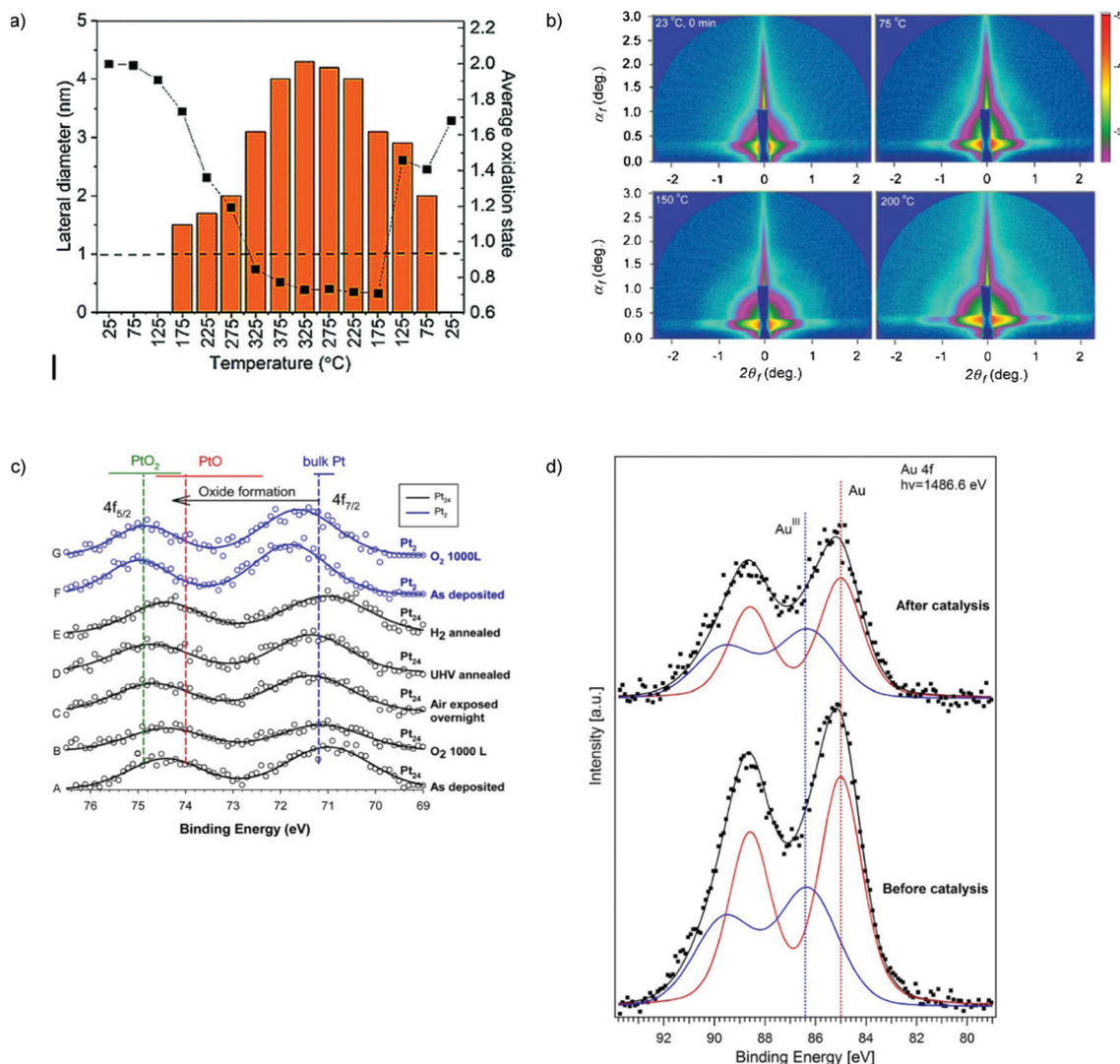


Fig. 5 (a) *In situ* characterization for the Cu₄ cluster sample under CO₂, H₂, and He in a 1 : 3 : 1 ratio, supplied at a rate of 5 sccm with 1000 ppm oxygen present. Plot of the change in the oxidation state of copper with the evolution of the particle size of the assembly. The dashed line drawn at 1 nm indicates the cut-off below which the used GISAXS geometry did not allow the resolution of the particle size. The estimated fitting error in the cluster composition obtained from XANES is between 5–10% (relative). (b) GISAXS images on ~9 nm silver particles. Change of the scattering pattern during ramping the temperature from 23 to 200 °C. (c) (A–E) Pt 4f XPS for Pt₂₄/SiO₂/Si: (A) as-prepared in UHV; (B) with 1000 L RT O₂ exposure; (C) after overnight air exposure; (D) after annealing to 150 °C in UHV; (E) after annealing at 150 °C in 10⁻⁴ torr of H₂; (F and G) Pt 4f XPS for Pt₂₄/SiO₂/Si, as deposited and after O₂ exposure. Horizontal bars show the range of 4f_{7/2} BEs reported for bulk Pt and oxides. Vertical lines show mean BE values. (d) XPS spectra of subnanometer ($d = 0.8 \pm 0.2$ nm) gold particles supported on 3-aminopropyl-functionalized silicate SBA-15 before and after its use in catalysis. (a) is adapted from ref. 118 with permission from the Royal Society of Chemistry, (b) is adapted from ref. 113, Copyright 2010, with permission from Elsevier, (c) is adapted with permission from ref. 121, Copyright 2016 American Chemical Society, (d) is adapted with permission from ref. 111, Copyright 2017 American Chemical Society.

X-ray

Combined *in situ* grazing incidence small angle X-ray scattering (GISAXS), grazing incidence X-ray absorption near-edge structure (GIXANES) and temperature programmed reaction (TPRx) experiment under a total pressure of 800 mbar was used to monitor the sintering resistance, oxidation state and catalytic performance of the Co₄ and Co₂₇ clusters.¹¹⁶ The GISAXS data showed no evidence of agglomeration of the clusters during the course of the reaction. The XANES spectra of the clusters are indicative of Co(II), whereas there is no

XANES evidence of Co(III). The spectra of the as-prepared samples suggest that Co clusters occupy various sites on the amorphous substrates.

Platinum–titania catalyst was studied by *in situ* electron microscopy, *in situ* X-ray photoemission spectroscopy and *in situ* X-ray diffraction, aided by density functional theory calculations, providing a real time view on how strong metal–support interaction occurs. The migration of reduced titanium oxide, limited in thickness, and the formation of an alloy are competing mechanisms during high temperature reduction.



Subsequent exposure to oxygen segregates the titanium from the alloy, and a thicker titania overlayer forms.¹¹⁷

An *in situ* characterization of Cu nanoassemblies formed transiently from size-selected subnanometer Cu₄ clusters supported on amorphous OH-terminated alumina during conversion of CO₂ into methanol and hydrocarbons under near-atmospheric pressure reveals that the clusters self-assemble into 2D nanoassemblies at higher temperatures which then disintegrate upon cooling down to room temperature. DFT calculations suggest a formation mechanism of these nanoassemblies by hydrogen-bond bridges between the clusters and H₂O molecules, which keep the building blocks together while preventing their coalescence¹¹⁸ (Fig. 5a).

XPS and STM were used for investigation of structures of Ag_n cluster anions ($n = 3-16$) deposited on sputter-damaged HOPG surfaces using a soft-landing technique at room temperature.¹¹⁹ Results suggested that clusters stayed as individual entities with minor agglomeration upon deposition. The deposited Ag clusters were oxidized using atomic oxygen and attempts were made to reduce them with CO, however, such a reduction was not found. The inertness of the small oxidized Ag nanoparticles on HOPG has been attributed to Ag carbonate formation.¹²⁰

A series of size-selected Pt_n/SiO₂ samples was analyzed using XPS, low energy ion scattering, GISAXS, and XANES in ref. 121.

The oxidation state and morphology were characterized both as-deposited in UHV, and after air/O₂ exposure and annealing in H₂. The clusters were found to be stable during deposition and upon air exposure, but to sinter when heated above ~150 °C (Fig. 5c).

A growth of Cu clusters on the TiO₂ support, including sintering and oxidation, was modeled using DFT.¹²² *Ab initio* thermodynamics calculations showed that O₂ readily oxidized most clusters, while H₂O was a mild oxidant, and CO₂ did not oxidize any clusters at relevant temperatures.

In a pioneering study, aberration-corrected STEM was used to analyze several iron oxide-supported Au cluster catalyst samples, ranging from those with little or no activity to others with high activities. High catalytic activity for carbon monoxide oxidation was correlated with the presence of bilayer clusters that are ~0.5 nanometer in diameter and contain only ~10 gold atoms¹²³ (Fig. 6c and d).

Temperature-programmed reaction of CO with O₂ catalyzed by Pd_n clusters ($n = 1, 2, 4, 7, 10, 16, 20, \text{ and } 25$) deposited on rutile TiO₂(110) was studied in combination with XPS.¹²⁴ The Pd 3d binding energy varied non-monotonically with cluster size and the changes correlated with strong size variations in CO oxidation activity. Low activity was correlated with higher-than-expected Pd 3d binding energy, which is attributed to a

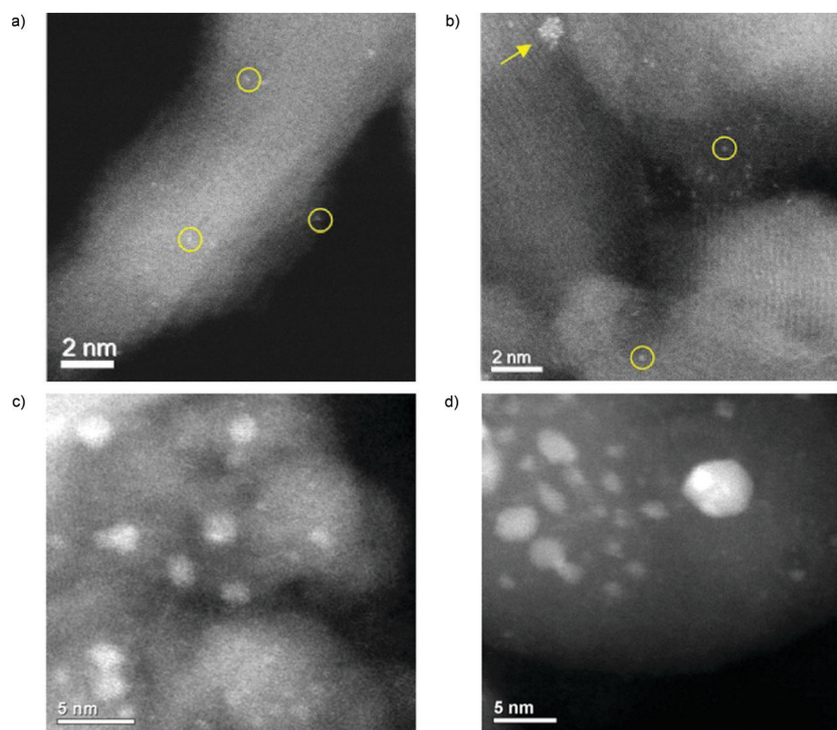


Fig. 6 Representative STEM images of Pt/ γ -Al₂O₃ samples through the catalysis experiments. 0.3 wt% Pt/Al₂O₃ catalyst after calcination (a), and after the full experiment (b) average size of the clusters (such as that pointed by the arrow): 1.0 ± 0.3 nm. Single Pt^{m+} atoms ($m \geq 2$), resulting from the standard impregnation–calcination procedure of SAC preparation, are poorly active. However, they gradually and irreversibly convert into highly active ~1 nm sized Pt^{δ+} clusters ($\delta < 2$) throughout the heating–cooling reaction cycles, even under highly oxidizing conditions favorable to atomic dispersion. Aberration-corrected STEM-HAADF images of the active 2.9 atomic% Au/FeOx catalyst 2 calcined for 3 hours at 400 °C (c), and 550 °C (d). The heat-treatment procedures have substantially decreased the population of subnanometer Au clusters relative to the highly active, dried catalyst, while at the same time they have increased the population of particles in the 1 to 3 nm range. (a and b) are adapted with permission from ref. 213, Copyright 2019 American Chemical Society, (c and d) are adapted with permission of American Association for the Advancement of Science, from ref. 123.



particularly stable valence electronic structure; electron transfer from the TiO₂ support to the Pd clusters also occurs. Ion scattering showed that small clusters form single-layer islands on the surface and that formation of a second layer begins to occur for clusters larger than Pd₁₀.

A liquid-phase-synthesis method to prepare isolated gold atoms supported on functionalized carbon nanotubes was presented and their performance in the oxidation of thiophenol with O₂ was studied.¹²⁵ It was shown that the single gold atoms are not active, but they aggregate under reaction conditions into gold clusters of low atomicity that exhibit a high catalytic activity. When clusters grow into larger nanoparticles, catalyst activity drops to zero.

Paramagnetic clusters possessing a well-defined structure of six equivalent Ag nuclei were prepared by reduction of oxidized silver–sodium containing zeolite LTA using hydrogen. An absorption of O₂ and/or C₂H₄ was inspected by means of continuous wave and pulse electron paramagnetic resonance (EPR) and *in situ* EXAFS. The unpaired spin density distribution was strongly affected by adsorption of C₂H₄ at room temperature, and the initially symmetric structure of the reduced Ag₆⁺ cluster turned less symmetric or less compact, lifting the equivalence of the silver atoms. The multiple-shell fitting distinguished a shell that is split into two components of slightly different distances within the limit of EXAFS. The overall coordination number is 3.90 ± 0.07 for the reduced silver cluster consisting of six Ag atoms. The EXAFS results showed that the Ag–Ag coordination number does not change on C₂H₄ adsorption.¹²⁶

EXAFS spectroscopy of titania-supported Pd-doped Au nanoparticle catalysts, in the as-prepared and post-catalysis state, shown that the distribution of Pd in Au depends on the chemical history of the samples: as-prepared samples contain highly-dispersed Pd close to the nanoparticle surface, whereas Pd is segregated as larger moieties in the post-catalysis samples. Pd restructures among Au as a result of exposure to the reaction conditions.¹²⁷

An interesting phenomenon has been unveiled in ref. 128 and 129, *i.e.*, the possibility that clusters change their adhesion to the support as a function of environmental conditions, such as temperature. In the theoretical paper,¹²⁸ it was shown how increasing temperature of an amorphous silica support leads to de-hydration of the silica surface, in agreement with experimental evidence,^{130,131} with the consequent formation of stronger adsorption sites for deposited species. In particular, if ultrasmall Au₆ clusters are deposited onto such silica support, an orientation transition from clusters singly pinned to the support and therefore perpendicular to it at low T, to clusters doubly anchored to the support and its stronger adsorption sites and therefore parallel to it at higher T is predicted, in excellent agreement with GISAXS measurements on the same system.¹²⁹

Subnanometer Ag_xAu_{3-x}/MgO(100) clusters⁹⁶ transform, after ligand adsorption into reaction complexes which catalyze CO oxidation through a variety of different mechanisms, occurring *via* both Langmuir–Hinshelwood and Eley–Rideal paths

and in some cases directly involving the oxide support. It was concluded that the reaction path for the reaction of an adsorbed O adatom with incoming CO would follow a Langmuir–Hinshelwood mechanism *vs.* an Eley–Rideal one depending on whether CO adsorption energy on a neighboring site on the cluster was large enough to overcome the loss of the free-energy of gas-phase CO and create a [–CO, –O] co-adsorbed species or whether adsorption was unfavorable with respect to a reactive scattering mechanism of CO “picking up” the O adatom from the cluster (that presented a small energy barrier). This conclusion was claimed to be general: a Eley–Rideal mechanism (*e.g.*, gas-phase CO picking up an O adatom or gas-phase O₂ reacting with adsorbed CO to produce CO₂ plus an O adatom) should hold in all cases in which (roughly) the absolute values of the entropic free-energy of the gas-phase molecule minus the adsorption energy is larger than the energy barrier for the scattering event (*i.e.*, roughly, the transition state of the adsorption process is higher than the transition state of the scattering event). This claim was supported by a later study of CO oxidation on Ag₉Pt₂₋₃ clusters supported on amorphous alumina in which it has been shown that reaction rates predicted along the above assumption are consistent with experiment,¹³² and in general by the need of assuming adsorption barriers to reconcile predicted and observed catalytic rates for small inert molecule activation at high temperature, see *e.g.* N₂ activation in ref. 133. Notably, the ligand/cluster/support catalytic complex in the case of Ag₃ was singled out as a carbonate Ag₃(CO₃) species. In this case, the formation of the carbonate does not poison the catalyst, on the contrary, it is beneficial for stabilizing the cluster *i.e.*, increasing the barriers to disaggregation and diffusion as demonstrated in successive work.^{134–136} The alloyed Ag₂Au₁ cluster was proposed as the best catalyst in terms of efficiency and robustness.⁹⁶ It is interesting to underline that in successive work it has been shown how the Ag₃(CO₃) species, which are the ligand/cluster/support catalytic complex for CO oxidation, transform into Ag₃(CO₃)(NO₃)₂ species under NO oxidation conditions, which are the ligand/cluster/support catalytic complexes for this reaction, still remaining able to act as a good catalyst,¹³⁵ a process resembling the catalyst preparation of “real-world” catalyst.

Clearly, dynamics effects can be very important to determine reaction kinetics and therefore to achieve a predictive modeling which can then be turned into rational design. It should be noted that the classical implementation of the Sabatier principle as “volcano curve” modeling⁸ typically focuses on descriptors based on electronic energy, usually neglecting entropic contributions or assuming they are constant with respect to a reference catalyst. However, as recalled above for the case of CO oxidation on Au_xAg_{3-x} and Ag₉Pt₂₋₃ clusters, recent research has demonstrated that entropic effects can give substantial contributions to overall reaction free-energy barriers, *e.g.*, stemming from adsorption free-energy barriers of gas-phase molecules at high temperature (such as CO¹³²). A quantitatively predictive modeling must take into account such effects, as done in several recent theoretical work, see *e.g.* ref. 133, 137 and 138. For example, a recent claim suggests that Brønsted–Evans–Polanyi (BEP) relationships can be broken through fluxional



dynamics of the catalyst: indeed, subnanometer gas-phase and graphene-deposited Pt_n cluster catalysts were shown to exhibit poor correlation between binding energies of the intermediates, O, OH, and OOH, involved in the scaling relationships for oxygen reduction reaction.¹³⁸ An advanced view of the issues facing quantitative modeling of entropic phenomena has been recently presented and we refer to this work for more discussion.¹³⁹

IR and UV-vis

Traditionally, IR and UV-vis spectroscopies have given an important contribution in catalytic studies.¹⁴⁰ To quote one clear example, kinetic, isotopic, and infrared studies on well-defined dispersed Pt clusters, in combination with first-principle theoretical methods, on model cluster surfaces, were performed to probe the mechanism and structural requirements for CO oxidation catalysis at conditions typical of its industrial practice.¹⁴¹ In the next section we will report several examples of a more recent implementation of these studies that correspond to *operando* modality.

Cluster stability

The issue of stability is one of the most important in catalysis science. To address the increasingly stricter environmental and sustainability requirements that modern society poses to the field, catalytic processes should be as atom-economical as possible and leach out as little as possible of their active species, which are often precious and sometimes potentially poisonous metals. The issues of catalyst disaggregation and diffusion leading to sintering are thus crucial for the future of the field. It is clear that, also under this respect, the catalyst support cannot be considered as a passive element, but plays a decisive role in maintaining the catalysis in a highly dispersed, intrinsically meta-stable state. Very many groups have therefore investigated the topic of catalyst stability and how to monitor it using physical chemistry techniques. Here we will present first the results of our own investigations, and then complete with reviewing other illustrative cases.

For subnanometer cluster catalysts, we start by recalling the concept of ligand/cluster/support catalytic complex.¹³ As anticipated above, both ligands and support and their interaction, are the factors determining the adhesion of the complex and its energy barriers to diffusion. We have already recalled as an example that the Ag₃(CO₃) species, generated during COox on Ag₃/MgO(100) catalysts, exploits its electrostatic interaction with the substrate to greatly increase its diffusion barriers.⁹⁶ A further major result in this respect when dealing with oxide, charge separated supports is that, especially at low temperature where *e.g.* oxide support is hydroxylated,^{130,131} the structure of the support likely corresponds to an amorphous one. In detail, one of the first *in situ* studies showing that amorphous support is essential for the stabilization of subnanometer clusters in HUNC is presented in ref. 142. Subnanometer Pt clusters on a crystalline ultrathin alumina film can easily diffuse on the support and sinter already at room temperature, whereas, once deposited onto a similarly ultrathin but amorphous alumina substrate, the ultrasmall Pt clusters are stable up to fairly high

temperatures. Even more importantly, in this work, it was demonstrated that the crystalline alumina support transform into an amorphous one under COox realistic reaction conditions, thus bridging the gap between UHV and real-world catalytic systems (Fig. 7a).

Concerning diffusion, we have conducted a series of systematic studies of this phenomenon¹⁴³ as the main effect controlling aggregation, Ostwald ripening and sintering. For model systems, an excellent agreement was achieved between the predicted kinetics of diffusion and aggregation and the experimentally observed one as a function of temperature,¹⁴⁴ thus validating the theoretical approach.

A competing pathways theory for sintering, which, at the atomistic level, describes the found size-specific sintering behavior, was proposed in ref. 145.

The analogy between homogeneous catalysis and heterogeneous catalysis by very small, subnanometer-size metal clusters, as proposed in ref. 134, seems promising to widen the spectrum of heterogeneous catalyst design and realize synergistic investigations with accumulated knowledge on homogeneous systems. The ligand/cluster/support complexes generated *in situ* under reaction conditions may be rather different from the original preparation, but it is their properties, and how to tune them, that must be the focus of operative research. Indeed, the ligand/cluster/support catalytic complex generated by CO oxidation over silver trimers deposited on the regular MgO(100) surface – *i.e.* a Ag₃/carbonate or Ag₃(CO₃)/MgO(100) species – can be used as a catalyst in a different reaction.¹³⁵ The Ag₃(CO₃)/MgO(100) species, however, transforms under NOox conditions into an even more complex aggregate, a mixed carbonate/double-nitrite Ag₃(CO₃)(NO₂)₂/MgO(100) species, which can then act as an efficient catalyst of NOox. Under NOox reaction conditions a different ligand/cluster/support catalytic complex is formed with respect to the original COox one. It is such diversity of the catalytic chemistry of ultranano metal clusters deposited on oxide substrates, the formation of many different ligand/cluster/support aggregates, and the vast amount of combinatorial possibilities thus opening, which makes detailed physico-chemical studies, including stochastic structural and stoichiometric computational searches, indispensable in order to cope with such a multiform diversity.

Dopant atoms may act as anchoring centers

The high activity and stability of Ag–Pt sub-nanometer clusters, as heterogeneous catalysts in the CO → CO₂ reaction (COox), are ascribed to a synergic role of Ag and Pt in ultranano-aggregates.¹³² It is notable that Pt anchors the clusters to the support and binds and activates two CO molecules, while Ag binds and activates O₂, and Ag/Pt surface proximity disfavors poisoning by CO or oxidized species. We recall once more that this Ag₃Pt₂₋₃ case study has been important to elucidate the important role of entropic effects (such as adsorption energy barrier) on catalysis, a role particularly crucial in the activation of small inert molecules, that require high temperatures to favor un-coverage of catalytic sites as well as bond breaking, however thus facing the issue of overcoming entropic barriers.



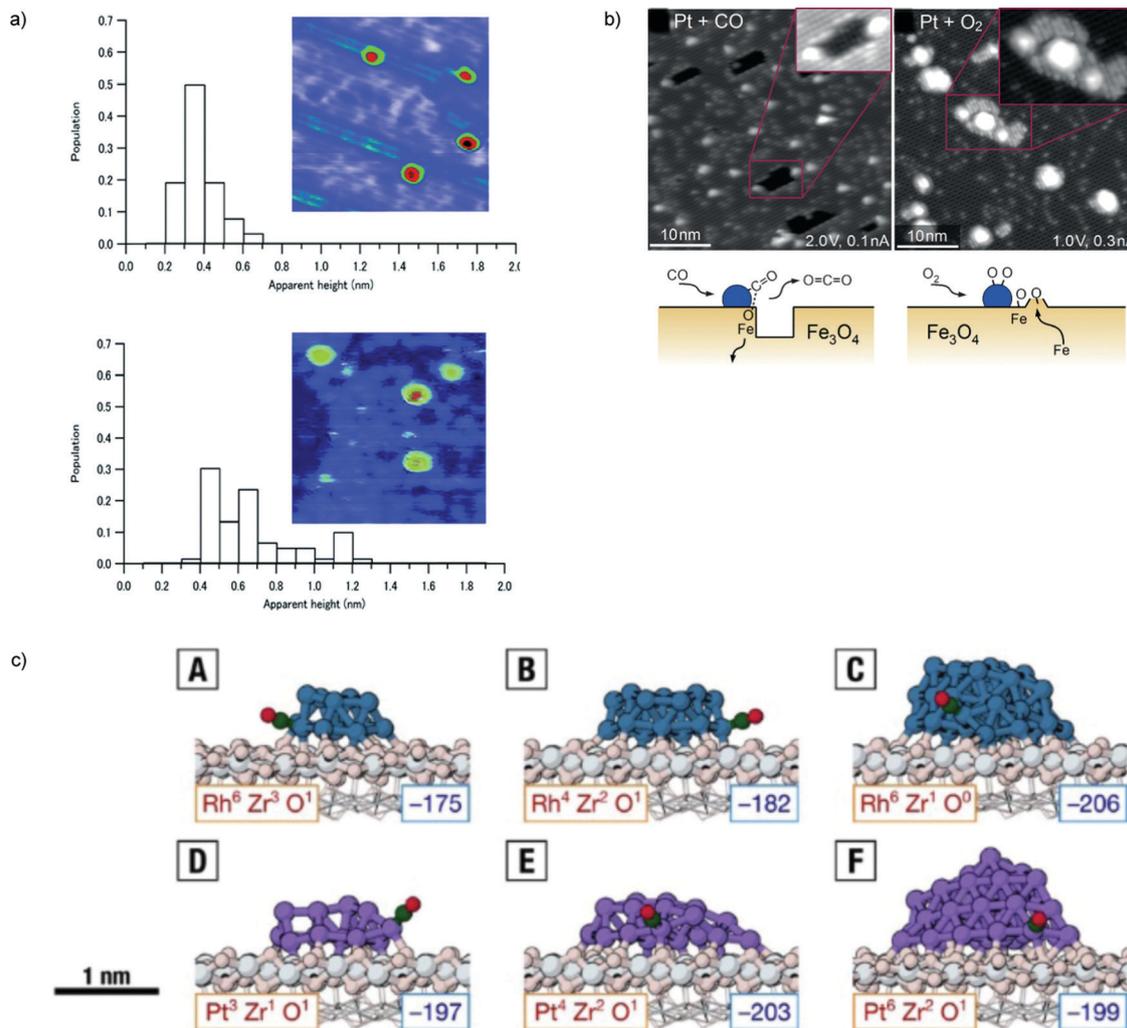


Fig. 7 (a) Histograms of apparent height and STM topographic images ($24 \times 24 \text{ nm}^2$, 3.5 V_s, 100 pA) of as-deposited size-selected Pt₁₀ clusters under vacuum at 27 °C (upper part) and after CO and O₂ exposure at 27 °C (lower part). (b) STM images acquired following exposure of the Pt/Fe₃O₄(001) model catalyst to 1×10^{-7} mbar of CO (left), or O₂ (right) at 550 K. Following exposure to CO (60 min), approximately 50% of the Pt clusters sit at the edge of or inside monolayer holes in the Fe₃O₄(001) terrace. The rows (rotated by 90°) of the next Fe_{oct}-O layer can be seen inside the holes (inset). The illustration shows how CO extracts O_{lattice} atoms from the cluster perimeter and how CO₂ desorbs from the surface. Undercoordinated Fe atoms diffuse into the Fe₃O₄ bulk. Following exposure to O₂ (20 min), all Pt clusters reside on top of islands with a step height of 2 Å. On the larger islands, the rows (rotated by 90°) of the next Fe₃O₄(001) layer are clearly visible. The islands result from the spillover of oxygen atoms onto the Fe₃O₄ support, which react with Fe that diffuses out from the bulk. (c) Most favorable adsorption of a CO molecule onto the perimeter sites of (A) Rh₁₃/ZrO₂, (B) Rh₁₉/ZrO₂, (C) Rh₄₃/ZrO₂, (D) Pt₁₃/ZrO₂, (E) Pt₁₉/ZrO₂, and (F) Pt₄₃/ZrO₂. The corresponding adsorption energies (in kJ mol⁻¹) are given in the bottom right corners of the panels. The coordination of the corresponding perimeter sites in the absence of CO is given in the bottom left corner of the panels. Color scheme: Rh in teal, Pt in purple, Zr in light gray, oxide O in pink, and CO in green and red. The fixed layers of the supporting oxide are shown in wireframe. (a) is adapted from ref. 142 with permission from the Royal Society of Chemistry, (b) is adapted with permission from ref. 102, Copyright 2015 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, (c) is adapted with permission from ref. 160.

The interaction between single metal atoms and support, as well as surface properties of supports, control the catalytic behavior of SACs. A systematic investigation of the effects of supports on CO oxidation by single Pt (Pt₁) atoms dispersed on different metal oxides (highly reducible Fe₂O₃, reducible ZnO, and irreducible γ -Al₂O₃) found that Pt₁ atoms on three metal oxides are active for CO oxidation, and the chemical properties of supports determine the catalytic performance of Pt₁ single-atom catalysts (SACs). Both the presence of -OH groups on support surfaces and the addition of H₂O significantly modify

CO oxidation on three SACs and reduce the effects of supports on their catalytic performances.¹⁴⁶ Note however that it has recently been claimed that dopant atoms can increase the reducibility of irreducible supports.^{147,148}

Doping, alloying and support effects on stability and catalyst performance

Modification of ultrasmall clusters by doping, overcoating, changing size and composition with atomic precision or through cluster support interactions provides high-fidelity tuning knobs



to control not only their stability, but their activity and, importantly, their selectivity as well.^{75,116,124,132,149–157}

Investigation of atomically dispersed Pt/TiO₂ catalysts by pre-/postreaction scanning transmission electron microscopy and operando X-ray absorption spectroscopy–mass spectrometry under gas-phase ethanol dehydrogenation conditions under ultraviolet–visible light irradiation shows that the general effect of the reaction on initially fully oxidized (Pt IV) single-atom catalysts (SACs) is Pt reduction and aggregation. This combined phenomenon can be inhibited in two manners. First, when it is supported on high-surface-area anatase, the Pt SAC retains an intermediate oxidation state, and its clustering is limited to Pt dimers and trimers. Second, with the same support, a mild reducing pretreatment generates, through the formation of Pt–Ti bonds, near-neutral and ultradispersed Pt species (from single atoms to nanometric clusters) with the highest stability and activity in photocatalytic hydrogen evolution.¹⁵⁸ Platinum complexes supported in zeolite KLTL remained site isolated in the zeolite pores after oxidation and catalysis of CO oxidation⁴⁴ (Fig. 8a).

The importance of dispersion effects on the cluster/support interaction was studied in ref. 32 where the energetics of an Au atom and Au_N clusters (*N* = 2–6) supported on pristine and reduced MgO(100) surfaces was analyzed using an all-electron full-potential density functional theory approach. For Au_N clusters (*N* > 1), dispersion effects start to make an increasingly important contribution to the adhesion energy.

Dispersion corrected density functional theory (DFT-D) was used to study the effect of commonly used support materials (MgO, C, CeO₂) on small gold particles with up to 19 atoms.¹⁵⁹ The preferred cluster shape and morphology is highly dependent on the support material due to different adsorption strength and structural mismatch between the cluster and the surface material. Depending on the support material, the gold clusters exhibit a positive or negative polarization, which

ultimately has strong implications on the catalytic activity of such particles.

A DFT-based genetic approach to obtain globally optimized nanostructures for Rh and Pt clusters on a ZrO₂ support was employed in ref. 160 and demonstrated that even an inert support can intricately influence the reactivity of interfacial sites. The analysis of the obtained structures showed that Rh clusters had more compact shapes, whereas Pt preferred elongated and low-symmetry structures. It was found that metal–oxide perimeter sites are structurally different, presenting varying Pt and Rh coordinations and CO adsorption energies. The presence of a support always destabilizes CO adsorption at the cluster edge, but the magnitude of destabilization varies substantially from site to site (Fig. 7c).

Small Cu clusters deposited on alumina support are more susceptible to oxidation than those in gas phase.¹⁶¹ This result was explained in terms of strong cluster–support interactions, which differ significantly for the oxidized and elemental clusters. The stable cluster phases also feature novel oxygen stoichiometries. The results suggest that one can tune the degree of oxidation of Cu catalysts by optimizing not just their size, but also the support they are deposited on. Effect of the morphology of the support on nanocatalysts performance in case of CO₂ methanation catalyzed by copper clusters was also pronounced in ref. 162. The Cu₄ supported by nanostructured zirconia (NS ZrOx) was found to be about 30% more active compared to Cu₄ supported by atomic layer deposited zirconia (ALD ZrOx), whereas Cu₁₂/NS ZrOx was 100% more active compared to Cu₁₂/ALD ZrOx. However, the activity of Cu₁₂/NS ZrOx dropped during the repeated cycle, most likely due to the possible migration of the clusters into the pores and voids with more limited access to reactants.

The strong effect of SiO₂ and MgO support on the activity of ethylene hydrogenation catalyzed by size-selected platinum clusters (8–20 atoms per cluster) was observed.¹⁶³ Predicted

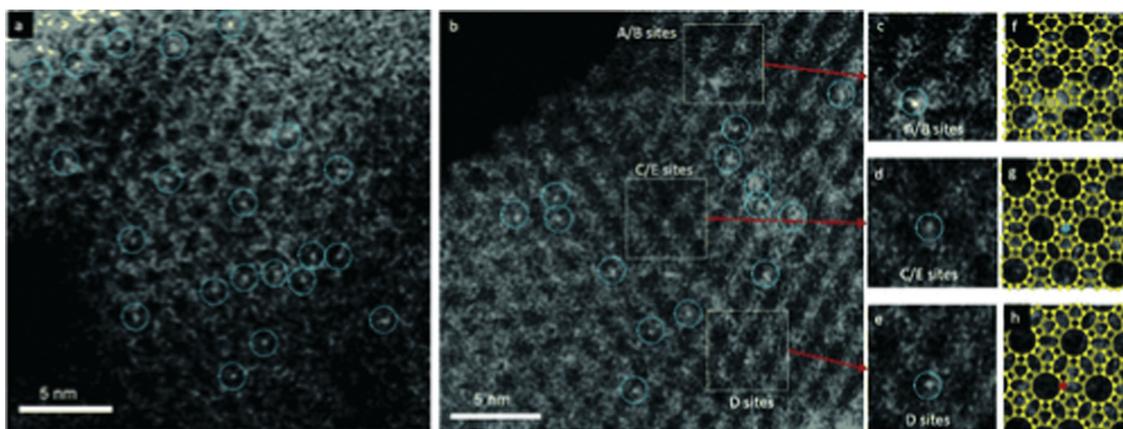


Fig. 8 STEM images showing site-isolated Pt atoms in KLTL zeolite in the (a) oxidized and (b) as-prepared samples. White features in dashed blue circles indicate Pt atoms. Magnified views (c–e) of the highlighted regions in (b), containing one Pt atom each at A/B sites in (c), at C/E sites in (d), and at D sites in (e). Simulations (f and g) of the LTL zeolite in the¹¹⁰ direction superimposed on the magnified views in (c–e), showing Pt atoms (green) at A/B sites in (f), at C/E sites in (g) (purple), and at D sites in (h) (red). Pt atoms are located right at the edge of the 12-membered rings of site D; between the two 12-membered rings of sites C/E; and in the center of three 12-membered rings of sites A/B. Adapted with permission from ref. 44, Copyright 2014 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.



trends based on the acidic (SiO_2) or basic (MgO) properties of the support are subtle, with the influence of the support also being a function of the cluster size. Pt_{13} had the lowest hydrogenation activity of all sizes on amorphous SiO_2 , while being the most active size on MgO . Adding Sn to size-selected Pt clusters deposited on amorphous SiO_2 also dramatically enhances the thermal stability of the clusters against sintering. The binding energy of Pt clusters to the support increases by >1 eV upon adding Sn, and Sn forms strong polar covalent bonds with Pt within the clusters and quenches all of the unpaired spins which tends to stabilize the Pt core of the clusters against sintering.¹⁵⁰

An *operando* generated stable Ni–In kinetic phase was discovered that selectively converts CO_2 to methanol (CTM) at low pressure compared to the state-of-the-art materials. The catalytic nature of a well-known methanation catalyst, nickel, has been tuned with the introduction of inactive indium, which enhances the CTM process. The remarkable change in the mechanistic pathways toward methanol production has been mapped by *operando* diffuse reflectance infrared Fourier transform spectroscopy analysis, corroborated by first-principles calculations. The ordered arrangement and pronounced electronegativity difference between metals are attributed to the complete shift in mechanism.¹⁶⁴

A novel *operando* XAFS technique was developed called the *operando* polarization-dependent total reflection fluorescence (PTRF)-XAFS technique, which can provide information on the valence state (XANES) and three-dimensional (3D) structure (EXAFS) of active metal species dispersed on a well-defined single-crystal surface during catalytic reactions. The technique was applied to a $\text{Pt}/\alpha\text{-Al}_2\text{O}_3(0001)$ model catalyst during the CO oxidation reaction. Pt clusters that favor an icosahedral Pt_{55} structure were formed on the $\alpha\text{-Al}_2\text{O}_3(0001)$ surface after Pt deposition at room temperature, while they were converted to larger cuboctahedral clusters (Pt_{147}) under the CO oxidation reaction at 493 K. The turnover frequency (TOF) of the CO oxidation activity at 493 K was also estimated to be 0.06 s^{-1} from simultaneous QMS and XAFS measurements.¹⁶⁵

High flux density X-ray beam on the catalyst used for *in situ* and *operando* studies can affect the catalyst structure. It was shown that the high flux density causes the reduction of the highly dispersed $\text{RhO}_x/\text{Al}_2\text{O}_3$ catalyst, even at room temperature. Additionally, exposure to the higher flux density X-rays during *in situ* reduction results in significant agglomeration of the Rh clusters. The final size of the Rh nanoparticles reduced at 310°C is equivalent to that of particles formed after the reduction at $600\text{--}650^\circ\text{C}$ in the absence of the beam. Significant beam-induced reduction and agglomeration is also shown for Ni supported on beta zeolite during *in situ* reduction at an intermediate-flux-density, indicating that beam-induced changes in heterogeneous catalysts could be common at intermediate- and high-flux-density beamlines.¹⁶⁶

Zeolites and MOFs

In zeolite-like systems, ultra-small cluster catalysts have been deposited since a while.⁸¹ Zeolite-confined noble metals have

unique confinement effects on a molecular scale, and thus enables spatially confined catalysis akin to enzyme catalysis.⁴⁷ Most of the currently available modeling tools are introduced and illustrated on the most challenging problems in zeolite science in review.⁴⁸

The combined experimental–theoretical approach, encompassing advanced molecular dynamics simulations and *in situ* micro-spectroscopy, gives a profound insight into the role of water in the catalytic process at the molecular and single-particle level in the methanol-to-olefins (MTO) process over H-SAPO-34.¹⁶⁷

Mordenite micropores provide a perfect confined environment for the highly selective stabilization of trinuclear copper-oxo clusters that exhibit a high reactivity towards activation of carbon–hydrogen bonds in methane and its subsequent transformation to methanol.⁴³ Structure determination of the copper-oxo active site was determined by DFT calculations and verified by XAS analysis. The reactivity data point to a uniform nature of the Cu sites in the Cu-MOR materials and, therefore, Cu K-edge spectra may yield direct information of the structure of the active clusters. *In situ* spectroscopy methods (XANES, EXAFS and UV-vis) were used to monitor the formation of the cluster during activation under O_2 and its interaction with CH_4 under reaction conditions.

Subnanometric hybrid $\text{Pd-M}(\text{OH})_2$ ($\text{M} = \text{Ni}, \text{Co}$) clusters encapsulated within purely siliceous zeolites were synthesized *via* a hydrothermal synthesis method.⁴⁰ The hybrid bimetallic nanocatalysts exhibit superior thermal stability at 600°C – 700°C because the bimetallic hybrid clusters are well confined within the zeolite matrix. The subnanometric sizes of the hybrid bimetallic Pd and $\text{Ni}(\text{OH})_2$ clusters were confirmed using STEM and EXAFS analyses.

Metal–organic frameworks (MOFs) have a big potential as platforms for the development of heterogeneous single-site catalysts.¹⁶⁸ Single atoms and few-atom clusters of platinum were uniformly installed on the zirconia nodes of a MOF NU-1000 *via* targeted vapor-phase synthesis.¹⁶⁹ *In situ* IR spectroscopy revealed the presence of both single atoms and few-atom clusters that depended upon synthesis conditions. Operando X-ray absorption spectroscopy and X-ray pair distribution analyses revealed changes in chemical bonding environment and cluster size stability while on stream. Density functional theory calculations elucidated a favorable reaction pathway for ethylene hydrogenation with the novel catalyst (Fig. 11b).

Ceria

Silver single atoms and small clusters supported on a model $\text{Ce}_{21}\text{O}_{42}$ nanoparticle have been studied computationally at a DFT+ U level.¹⁷⁰ It was found that silver atoms, trimers and tetramers are oxidized upon interaction with the (100) and (111) nanofacets of the ceria particle and adsorption of silver species at (100) facet is $\sim 0.5\text{--}1.0$ eV stronger than to (111) one. Ag_3 are adsorbed as upright triangles; Ag_4 prefer tetrahedral configuration. Denucleation of silver clusters to atomic species at (100) facet is the most favored.



Effect of ligands on electronic properties

Ligands can also strongly affect redox properties of clusters as it was shown in theoretical study.¹⁷¹ Attaching phosphine ligands to simple metal, noble metal, semiconducting, metal-oxide, and metal-chalcogen clusters is shown to severely reduce ionization energies in all classes of clusters. Ligation is shown to be an outstanding strategy for the formation of multiple electron donors.

Clusters in electrochemical applications

An overview of studies performed under both realistic reaction conditions aimed at the interrogation, characterization, and understanding of the performance of supported size-selected clusters in heterogeneous and electrochemical reactions, which address the effects of cluster size, cluster composition, cluster-support interactions, and reaction conditions, the key parameters for the understanding and control of catalyst functionality, was provided in ref. 172. Several ultrasmall systems have been studied in this context. Without pretending to be exhaustive, we will discuss a few cases below which further underline the importance of a multi-probe characterization of subnanometer size clusters along with the atomic-precise tunability of their performance in various electrochemical and electrophotocatalytic processes, such as oxygen or hydrogen evolution reaction, CO₂ reduction or batteries.

Pd₄, Pd₆, or Pd₁₇ clusters soft-landed on a ultrananocrystalline diamond (UNCD) Si-coated electrode show stable electrochemical potentials over several cycles, and synchrotron studies of the electrodes show no evidence for evolution or dissolution of either the electrode material or the clusters¹⁷³ under for the oxygen evolving reaction (OER). Even under the harsh experimental conditions (basic, high potential) typically employed for water oxidation catalysts, UNCD exhibits a very wide potential electrochemical working window and only minor evidence of deterioration, and with Pd clusters decorated, these electrodes reached the highest reported rates of oxygen formation based on metal loading.

Arrays of discrete iridium clusters with identical metal atom number (Ir₂, Ir₄, or Ir₈) were deposited in submonolayer coverage on conductive oxide supports, and the electrochemical properties and activity of each was evaluated.¹⁷⁴ Exceptional electroactivity for OER was observed for all cluster samples in acidic electrolyte. Reproducible cluster-size-dependent trends in redox behavior were observed and rationalized *via* first-principles modeling of the individual discrete-size clusters.

Oxygen reduction (ORR), as well as ethanol oxidation on subnanometer size-selected Pt clusters showed high activity as well as non-linear dependency of activity with cluster size, discussing activity correlated to the electronic structure of the Pt clusters.¹⁷⁵

The possibility of realizing low overpotential electrocatalytic conversion of CO₂ using subnanometer copper clusters deposited on glassy carbon was demonstrated in ref. 176 where the

electrochemical behavior (in the presence of N₂ or CO₂) of size-controlled naked Cu₅ and Cu₂₀ nanoclusters was studied. Cu₂₀ nanoclusters cause anodic redox processes stirring at much lower potential with respect to Cu₅ nanoclusters, which behave relatively similar to much larger Cu particles. However, Cu₅ nanoclusters coordinate effectively CO₂ in solution, in comparison to Cu₂₀ nanoclusters and larger Cu particles. This effect, rather than the redox behavior, is evidently connected to the ability of Cu₅ nanoclusters to reduce CO₂ under cathodic conditions at low overpotential and indicate the tunability of electrochemical performance within the subnanometer size range.

Size-selected cobalt clusters (Co₂₇) deposited on hematite (Fe₂O₃) anodes were tested for water oxidation activity in ref. 177. The Co₂₇ clusters are stable to dissolution even in the harsh water oxidation electrochemical environment. The turnover rates for water oxidation are comparable or higher than those reported for Pd- and Co-based materials or for hematite. Annealing of the samples (with the clusters already deposited) significantly improved the electrocatalytic and the photoelectrocatalytic activity of the Co₂₇/Fe₂O₃ material, another control knob in the design of high-performance subnanometer cluster-based materials.

In the study described, subnanometer sized palladium clusters accommodated into porous ceria were synthesized and showed highly efficient electrocatalytic activity and stability for the hydrogen evolution reaction (HER).¹⁷⁸ The synthesized Pd subnanoclusters have a size of about 0.5 nm as it was shown by characterizations from high-resolution transmission electron microscopy (HRTEM) and extended X-ray absorption fine structure (EXAFS). Density functional theory suggested that the active centers are the subunit of Pd clusters rather than CeO₂ support based on the free energy of absorption for hydrogen atoms (ΔG_{H^*}). Compared to the free Pd nanoclusters, the electron transfer between Pd and O atoms in Pd NC@CeO₂ can effectively modulate the adsorption state of H* on the subunit of the Pd cluster, and this conjoint effect can effectively promote the HER catalytic activity of the composite system.

The degradation of size-selected Pt nanoclusters was studied under electrochemical conditions by HAADF-STEM and confirmed by transmission electron microscopy (TEM).¹⁷⁹ It was demonstrated that even extremely small Pt clusters exhibit a remarkable stability under electrochemical conditions. Furthermore, applying mixed cluster samples of Pt₂₂ and Pt₆₈, no preferential dissolution of Pt₂₂ by Ostwald ripening – usually held responsible to be the main mechanism for activity loss in Pt fuel cell catalysts – was observed.

Another direction of great interest for future developments is the Hydrogen Evolution Reaction (HER). Indeed, it has been demonstrated that subnanometric metal clusters can be used as photocatalysts for H₂ production in the presence of holes or electrons scavengers by water splitting.¹⁸⁰

Copper-platinum (Cu-Pt) dual sites were alloyed with palladium nanorings (Pd NRs) containing 1.5 atom% Pt, using atomically dispersed Cu on ultrathin Pd NRs as seeds. The ultrafine structure of atomically dispersed Cu-Pt dual sites was



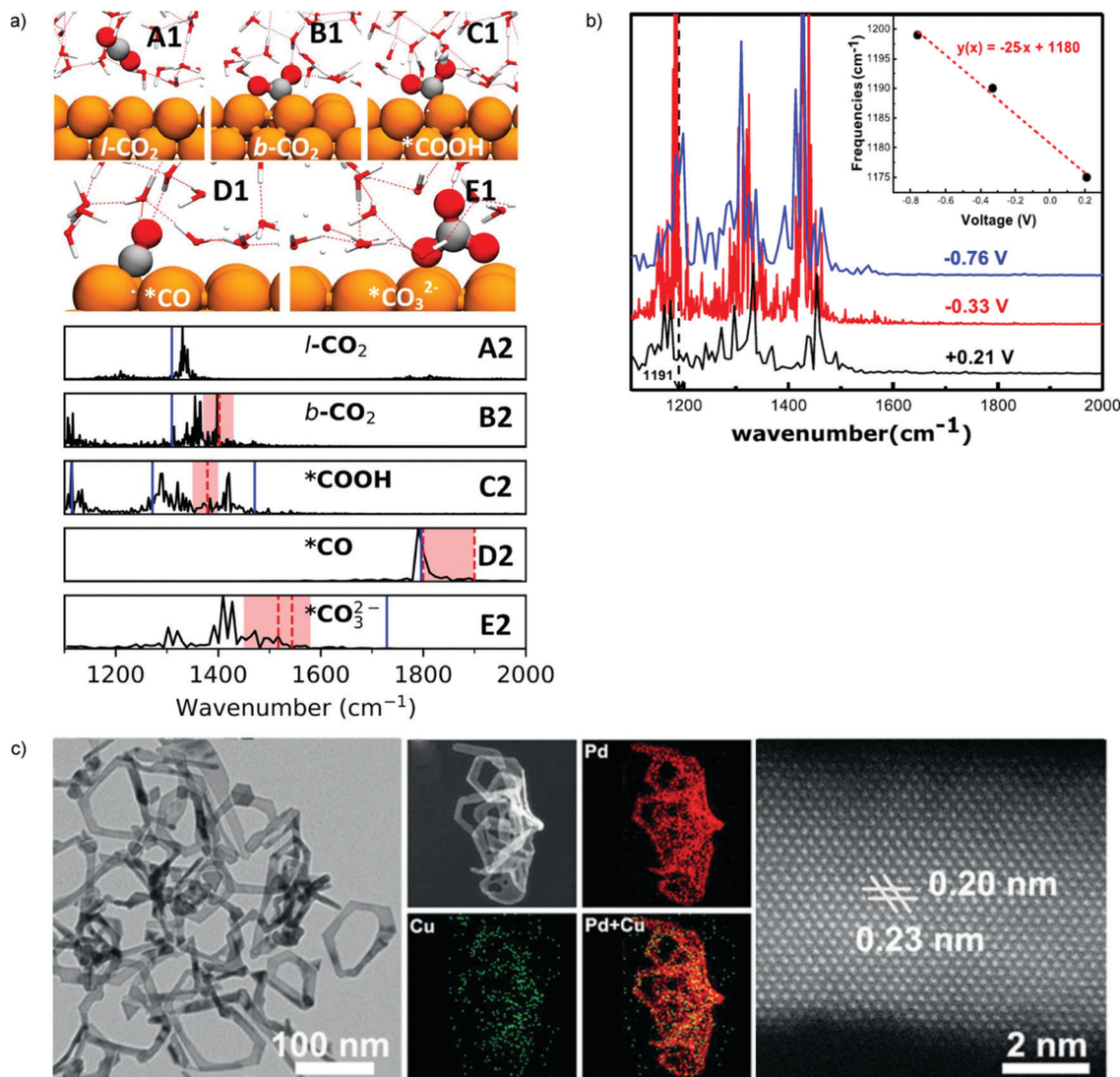


Fig. 9 (a) Snapshots of the structures and v-DoS CO₂ (l-CO₂), chemisorbed CO₂ (b-CO₂), *COOH, *CO, and *CO₃²⁻ from o-QM-MD simulations and 2PT analysis. The color codes are Cu in orange, O in red, C in silver, and H in white. The hydrogen bonds are shown as dashed red lines. In bottom part, v-DoS from o-QM-MD is shown as a solid black line, the experimental frequencies are shown as a red dashed line, and the vibrational frequencies from v-QM optimization are shown as solid blue lines for comparison. (l-CO₂ and b-CO₂ are the same from v-QM, because b-CO₂ is unstable in vacuum and automatically converts to l-CO₂ after optimization.) (b) The simulated potential-dependent v-DoS of *HOC-COH at +0.21 V, -0.33 V, and -0.76 V. Inset shows the blue shift of the 1191 cm⁻¹ peak (a feature of C-O stretch of C-OH) as a function of applied potential. (c) Characterization of Pd/Cu NRs. From left to right: TEM image; HAADF-STEM image and the corresponding STEM-EDX elemental mapping; atomic resolution HAADF-STEM image. (a) and (b) are adapted with permission from ref. 187, (c) is adapted with permission from ref. 181, Copyright 2017 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

confirmed with EXAFS measurements. The Pd/Cu-Pt NRs exhibit excellent HER properties in acidic solution, also supported by DFT calculations¹⁸¹ (Fig. 9c).

Indications for the theoretical design of HER subnanometer catalysts were drawn in ref. 182. Ag-Au subnanometer clusters as catalysts were proposed for the HER and investigated computationally, by conducting an exhaustive first-principles study of the HER process. Interestingly, the hydrogen-saturated resting state under standard conditions was determined as Ag₁₂AuH₁₁, from which HER reaction energies and barriers were predicted also including solvent effects using both implicit and explicit models. It was found that Ag₁₂Au is a good

candidate as a HER catalyst, with good stability and an overall reaction energy barrier of 0.89 eV.

A method for the electrodeposition of an isolated single Pt atom or small cluster, up to 9 atoms, on a bismuth ultramicroelectrode (UME) was demonstrated in ref. 183. The electrochemical characterization was made *via* the hydrogen evolution reaction (HER) that occurs readily on the electrodeposited Pt but not on Bi. Taking the potential at a certain current density as a measure of the relative rate of the HER, it was found that the potential shifted positively as the size increased, with single atoms showing the largest overpotentials compared to bulk Pt.



It should be underlined that a development of great potential in terms of characterization under electrochemical conditions is represented by electrochemical scanning tunneling microscopy (EC-STM),¹⁸⁴ naturally in combination with standard electrochemical measurements such as cyclic voltammetry (CV). To the best of our knowledge, this technique has not yet been applied to nanocatalyst systems, but it is to be expected that its recent progresses will allow researchers to achieve this soon.¹⁸⁵

Another development that has already achieved very interesting progress and is extremely promising in perspective is related to *operando* characterization modality. Recent experimental techniques have in fact been developed that allow one to realize *operando* identification of reaction intermediates using surface infrared, Raman, and XPS spectroscopy. To provide an interpretation of these experimental studies and target for additional studies, theoretical calculations are needed, for example generating vibrational spectra of the various candidate intermediate that can be directly compared with the reported experimental spectroscopy data.¹⁸⁶ One recent finding is that explicit consideration of solvent, electrode–electrolyte interface, and applied potential effects in theoretical/computational modeling is indispensable to achieve quantitative agreement and correct interpretation of experiment. Spectroscopic features (peak positions and widths) can depend substantially on both the applied potential in terms of charge residing on the surface, and on interfacial atomistic structures *e.g.* the network of hydrogen bonds surrounding the adsorbate species under consideration. An example of such *operando* theoretical spectroscopy, in this case fully based on quantum mechanics molecular dynamics, can be found in ref. 187, where the possible reactive intermediates in carbon dioxide reduction over the Cu(100) surface were characterized *via* vibrational spectroscopy and important ones identified in the experimental spectra, uniquely different from those predicted neglecting potential and solvent effects (Fig. 9a and b). Experimentally, the determination of XPS core-level shifts under electrochemical conditions is difficult, due to the perturbing background generated by the environment, but progress is continuously being made since the pioneering studies in this field.¹⁸⁸ XPS prediction also faces technical issues especially in finite-state simulations due to the hole localization in periodic systems, to overcome which a local fragment similarity (LFS) procedure has been recently proposed.¹⁸⁶ In this procedure, accurate and predictive final-state XPS simulations on fragment species are used as a database to complement and correct lower-level simulations on the real systems in the initial-state approximation. Finally, electrochemical scanning tunneling microscopy stands as a particularly promising approach to monitor with atomic-scale precision the catalytic activity of subnanometer species, such as *e.g.* single iron atoms anchored to carbon vacancies in graphene.¹⁸⁹

Lastly, size-selected subnanometer silver and iridium clusters have been also found as efficient electrocatalysts in Li–O₂ batteries¹⁹⁰ and helped to identify the composition and structure of the active species in iridium-based Li–O₂ batteries,¹⁹¹ respectively.

Optical properties of clusters

Optical spectroscopy is potentially a powerful tool for characterization and manipulation (photocatalysis) of nanostructured catalysts. Indeed, the optical properties of larger metal nanoparticles, both pure and alloyed have been the subject of a wealth of studies in the past decades,^{87,192} with the interest on these properties stemming from their technological applications, which range from plasmonic materials, to sensors and single molecule spectroscopy (SERS). The delocalized electrons of metal nanoparticles are in fact strongly responsive to electromagnetic external fields, to which they react giving rise to coherent and collective excitations and correspondingly strong absorption peaks in the optical region of the spectrum, called surface plasmon resonances (SPR).¹⁹³ In the present review, it is worthwhile underlining that applications of optical properties include photo-catalytic studies.^{158,194} It can be added that, when the structure of the clusters is atomically precise as in the subtopic of monolayer-protected clusters, systematic investigations of optical absorption and ultra-fast response, including plasmonic effects, have been conducted in the last decade, providing a mass of often well rationalized data and interesting perspectives, as will also be recalled later.⁸⁵ This may lead one to expect a similar importance of the topic of chiro-optics also in the subnanometer supported cluster field. However, the technical difficulties associated with weak signals and the need of disentangling response of the catalyst, the reactants/products, and the support have so far hindered the full exploitation of the potentialities of this technique. In the following we will briefly review what has been achieved already together with some appealing perspectives, recalling that one of our goals in this review is to trigger interest on and promote developing new promising areas.

A systematic study of optical absorption spectra from the ultraviolet to the visible for size selected neutral Ag_{*n*} clusters (*n* = 5–120) embedded in solid Ne was published in ref. 195. With increasing size of the Ag_{*n*} clusters, several highly correlated electron excitations gradually develop into a single surface plasmon with energy between 3.9 and 4.1 eV. The plasmon energy is highest for clusters with atom numbers fully filling states with the lowest radial quantum number (*e.g.* 1s, 1p, 1d, ...). TDDFT calculations for clusters with several candidate geometrical structures embedded in Ne show excellent agreement with the experimental data, demonstrating that the absorption bands depend only weakly on the exact structure of the cluster (Fig. 11c).

Surface cavity ring-down (s-CRD) spectroscopy under ultra-high vacuum (UHV) conditions was used to measure extinction spectra of size-selected, supported Ag₂₀ and Ag₅₅ clusters.¹⁹⁶ The reaction with benzenethiol shifted the localized surface plasmon resonance observed around 3.2 eV to lower energies. This was attributed to an increased dielectric function of the surrounding medium as well as to a reduction of the free-electron density inside the silver clusters. Time-resolved study of that shift revealed that two thirds of the overall redshift can be attributed to an increased dielectric constant of the



surrounding medium, whereas a reduction of the free-electron density accounts for one third, in agreement with previous theoretical TDDFT predictions.¹⁹⁷

The oxidation reaction of supported, size-selected small silver clusters between Ag₉ and Ag₅₅ was monitored using surface second harmonic generation spectroscopy (s-SHG).¹⁹⁸ A rapid decline of the SH-intensity was attributed to the formation of silver–oxygen-bonds. A double-exponential character of the time evolution of the SH-intensity for all investigated cluster sizes was observed. The two independent pathways of SH-intensity loss were attributed to a surface- and an interface-oxidation of supported clusters, respectively.

In addition to simple absorption, more sophisticated phenomena such as photoluminescence have also been studied. Intense fluorescence of Au₂₀ in He matrix held at temperature of 6 K was reported in ref. 199, related with the HOMO–LUMO diabatic bandgap of the cluster. Since Au₂₀ has only one stable isomer (tetrahedral) and due to low temperature, Au₂₀ shows a very rich absorption fine structure reminiscent of well defined molecule-like quantum levels, which are fairly well reproduced both in absorption and intrinsic fluorescence by DFT and TDDFT calculations (Fig. 11d).

Ag clusters confined within Na-exchanged Linde type A zeolites were studied by X-ray absorption and steady-state and time-resolved photoluminescence spectroscopies in a coordinated effort to reveal the photophysical properties and link them to the precise cluster structure.²⁰⁰ At room temperature, the emission of the hydrated sample decays with 3.4 ns from a state with the same multiplicity as the ground state. Upon dehydration, the entire excited-state dynamics speeds up to 1.2 ps. The microsecond-scale lifetimes observed at 77 K suggest the occurrence of two main decay processes for the initially populated singlet state: intersystem crossing and charge transfer. The intersystem crossing yields the formation of a long-lived (409 μs) triplet state ³P from the ¹P state located at lower energy from which luminescence occurs. There is evidence that electron transfer is followed by electron–hole recombination or back electron transfer, yielding a relaxed singlet excited state. The dependence of luminescence lifetime on the water content indicates a possible way to speed or delay the electron–hole recombination in a controlled way (Fig. 11a).

An ultrafast preparation of subnanometer silver particles with high abundance, uniformity (7 Å), and stability into the cages of nanosized zeolite crystals was presented in ref. 41. The particles were prepared using UV excitation of a water suspension of nanozeolite containing photoactive vanadate clusters in the presence of ethanol (as an electron donor) and silver precursor. The subnanometer Ag clusters exhibit exceptional photocatalytic activity and selectivity in the reforming of formic acid to H₂ and CO₂ under visible light (Fig. 10b).

A charge engineering strategy to improve the photo-oxidation activity and stability of Au₂₅(PPh₃)₁₀(SC₃H₆Si(OC₂H₅)₃)₅Cl₂ nanocluster (AuNC) taking advantage of Z-scheme AuNC/ultra-thin BiOCl nanosheets (2D-BiOCl) heterojunction for visible light oxidative self-coupling of amines was reported. The enhanced photocatalytic activity of AuNC/2D-BiOCl was predominantly

attributed to the formation of Z-scheme heterojunction, which effectively accelerated the transfer and separation of photo-generated carriers. Especially, the consumption of holes in AuNC by injected electrons from 2D-BiOCl can inhibit the self-oxidation of AuNC, thus improve its stability.²⁰¹

High-energy-resolution X-ray absorption spectroscopy detection of ethylene and CO ligands adsorbed on catalytically active iridium centers isolated on zeolite HY and on MgO supports, supported by density functional theory and FEFF X-ray absorption near-edge modelling, together with IR spectra, was reported in ref. 45. The high-energy-resolution X-ray absorption spectra near the iridium L_{III} (2p_{3/2}) edge provided clearly ascribable, distinctive signatures of the ethylene and CO ligands and illustrated effects of supports and other ligands. The X-ray absorption technique was more sensitive than conventional IR spectroscopy for characterizing surface intermediates, and it should be applicable to samples having low metal loadings and in reactive atmospheres.

A recent first-principles DFT and TDDFT study investigated the optical response of small (M/M')₆ and (M)₃(M')₃ six-atom clusters (unary and equimolar binary hexamers of group IB metals: Cu, Ag, and Au), both in the gas-phase and supported on the MgO(100) surface, chosen as a model of a simple oxide substrate.²⁰² UV-vis spectra were predicted and analyzed in detail in terms of ground- and excited-state wave functions and the involved Kohn–Sham orbitals, to rationalize origin and features of optical absorption in these systems. The interaction with the surface, in particular with the electric field generated by the charge-separated substrate, was found to induce a fragmentation and a shift toward lower energies of the absorption peaks, which was in tune with available experimental information and can be beneficial in terms of applications of these systems as photoenhancers or promoters. The orientation of the transition dipole moment with respect to the support (whether parallel or perpendicular) was also analyzed as a key parameter to translate the present result to non-inert supports to tune and optimize oscillator strengths and interaction with substrate excitations. These indications have been later confirmed in simulations on larger Ag₂₀ and Ag₂₀ clusters deposited on MgO(100), in which modeling was also extended to include defects on the oxide substrate such as oxygen vacancies and/or substitutional cation replacements. Interestingly, it was found that defect excitations can couple with cluster one and that the simultaneous presence of vacancies and cation doping may give rise to fine effects on transition dipole moment orientation.²⁰³ Finally, an exploratory study of optical absorption and photo-decay processes occurring in Ag₃(XCO₃)(C₂H₄)₂(O) ligand/cluster/support catalytic complexes (with X = H or F) after visible light absorption was recently conducted.¹³⁶ Although a “frailty” of bicarbonate and fluorocarbonate groups to optical adsorption was derived from this study, leading to C₂H₄ or CO₂ detachment or Ag₃ disaggregation, from the methodological point of view a palette of computational tools was tuned, enabling one to explore the promising field of subnanometer photo-catalysis.

It should be stressed that decisive progress has been recently made in this topic. Methods have in fact been recently



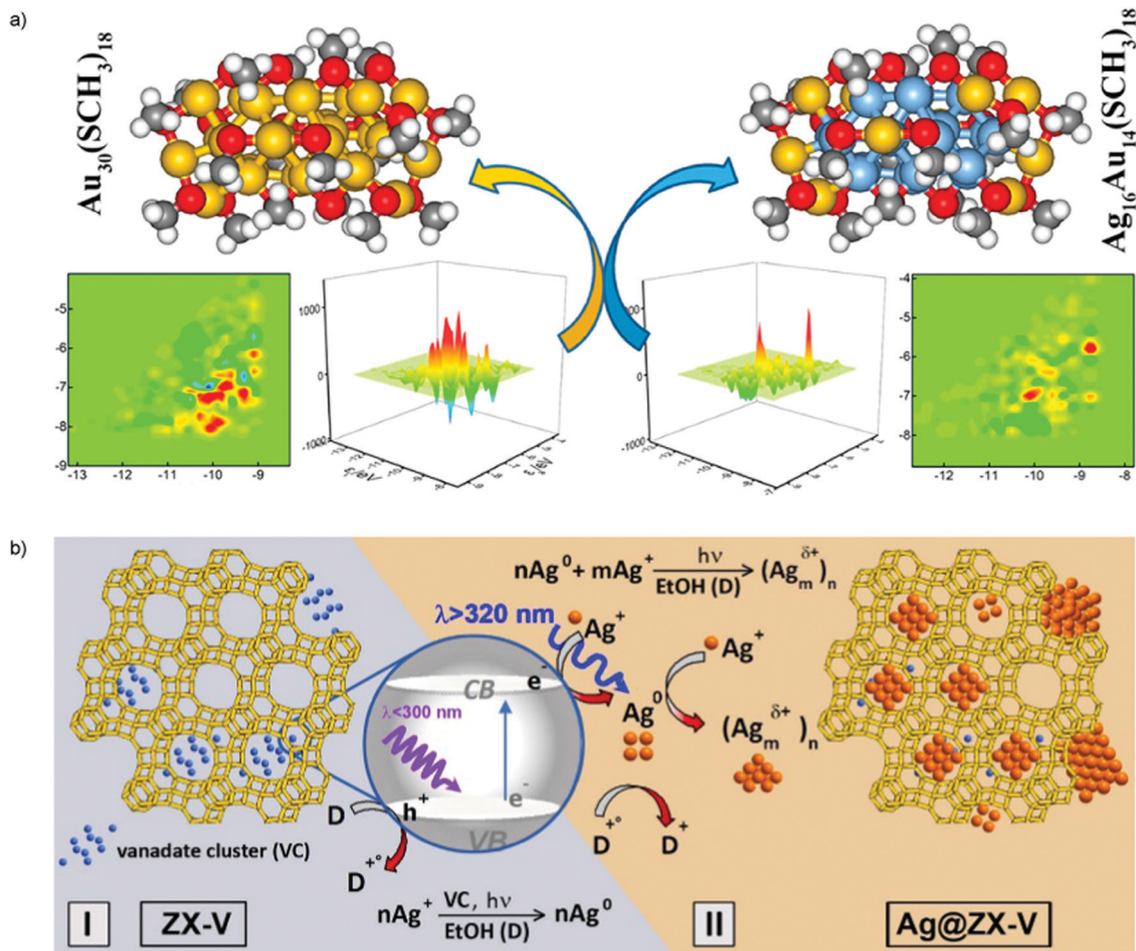


Fig. 10 (a) Individual component maps of rotatory strength (ICM-RS) and rotatory strength density (RSD) plots as analysis tools of chiro-optical linear response spectra deriving from time-dependent density functional theory (TDDFT) simulations to analyze circular dichroism spectra of selected $(\text{Ag}-\text{Au})_{30}(\text{SR})_{18}$ monolayer-protected metal nanoclusters. (b) Illustration of the Photocatalytic Process Leading to the Formation of Silver Clusters in the Supercage of FAU-Type zeolite (I) silver photoreduction by excitation of the vanadate clusters (VCs) with $\lambda < 300 \text{ nm}$ (D); (II) photoreduction Ag^0 formed in the stage (I) by excitation of silver clusters with $\lambda > 320 \text{ nm}$ (D). (a) is adapted with permission from ref. 209, Copyright 2018 American Chemical Society, (b) is adapted with permission from ref. 41, Copyright 2018 American Chemical Society.

developed to predict photo-induced processes in complex systems such as ligand-protected⁸⁴ and oxide-supported subnanometer clusters,^{204,205} with an accuracy quantitatively validated against experiment, thus opening a pathway for predictive modeling photo-induced and photo-chemical reactivity in these materials. Indeed, in ref. 204, a rutile $\text{TiO}_2(110)$ surface decorated with Ag_5 subnanometer clusters and its photon-absorption spectrum were investigated, predicting the formation of surface polarons on TiO_2 that has then been confirmed experimentally. Similar phenomena were predicted in ref. 205 on a rutile $\text{TiO}_2(110)$ surface either bare or decorated with Cu_5 subnanometer clusters. It was also predicted that electron donation from the Ag_5 or Cu_5 cluster enhances overall absorption and extends it somewhat into the visible region, with perspective applications in CO_2 photoactivation.²⁰⁶

Indirect nanoplasmonic sensing (INPS) coupled with mass spectrometry was used to study CO and oxygen adsorption as well as CO oxidation, on Pt nanoparticles, in the Torr pressure

range. A highly sensitive probe that can detect gas adsorption of a few hundredths of a monolayer, even with a very low number density of Pt particles was obtained. A similarity was observed between the sign and the evolution of the localized surface plasmon resonance (LSPR) peak shift and the work function measurements for CO and oxygen chemisorption.²⁰⁷

We believe that one important step in achieving a deeper understanding of optical phenomena is undoubtedly the development of analysis tools which allow one to decompose the various physical chemical effects at play into a few basic ingredients and single out descriptors able to orient research and design of optimal systems. For the sake of brevity, we here mention only two recent developments that, due to their general character, can be applied to a wide range of phenomena, also after needed methodological extensions. An Individual Component Maps of Oscillatory Strength (ICM-OS) analysis tool of TDDFT simulated spectra have first been introduced and applied in the field of monolayer-protected clusters (MPC) to investigate



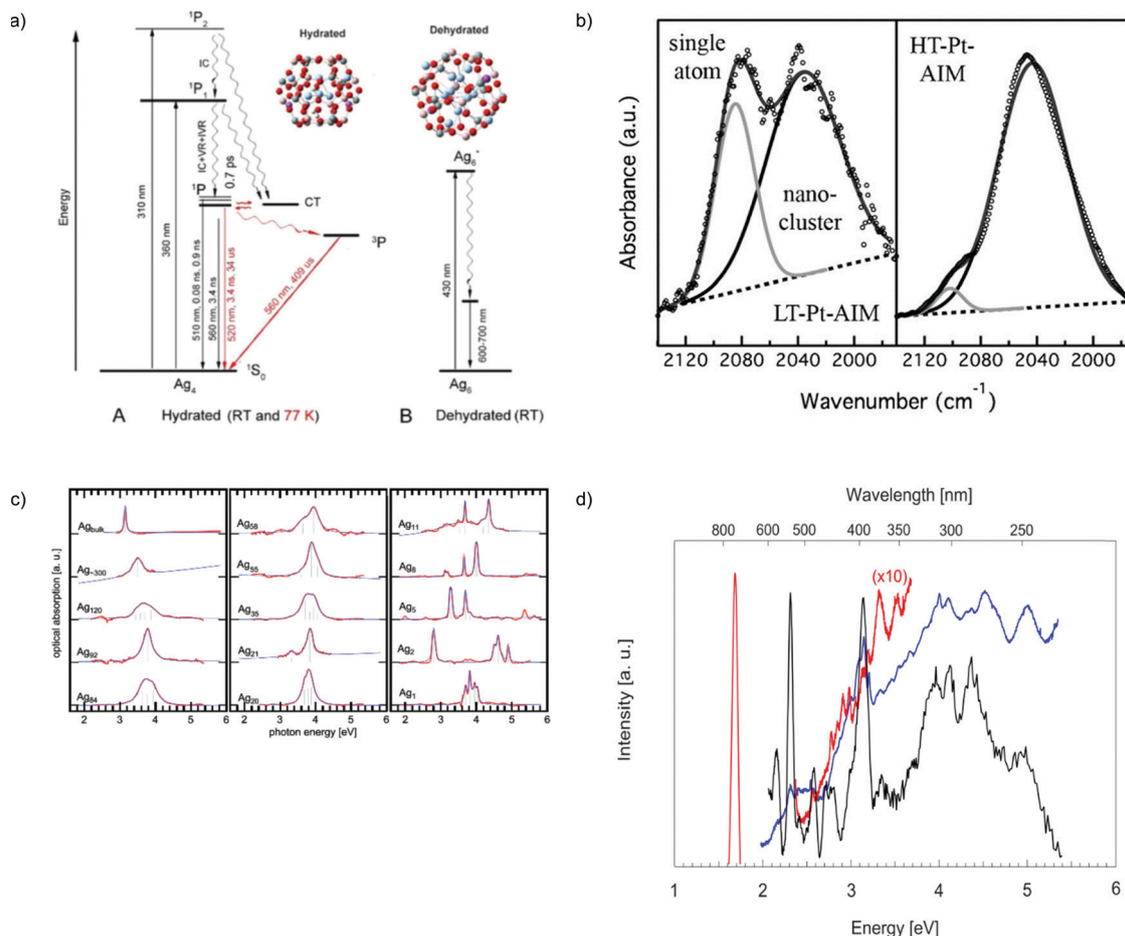


Fig. 11 (a) Ag clusters (AgCLs) confined within Na-exchanged Linde type A zeolites show dependence of luminescence lifetime on the water content indicating a possible way to speed or delay the electron–hole recombination in a controlled way. Proposed simplified kinetic scheme of the processes observed upon excitation of hydrated (A) and dehydrated (B) samples. The processes depicted in black and red are observed at RT and 77 K, respectively. (b) Single atoms and few-atom clusters of platinum exhibit high catalytic activity for ethylene hydrogenation while exhibiting resistance to sintering up to 200 °C. *In situ* IR spectroscopy reveals the presence of both single atoms and few-atom clusters that depend upon synthesis conditions. IR spectra of CO adsorbed on LT-Pt-AIM (left) and HT-Pt-AIM (right). (c) Measured optical absorption spectra (red) of monodisperse Ag_n ($n = 1-120$) in Ne at 6 K. The uppermost spectrum was calculated for bulk Ag and redshifted by 0.17 eV in order to account for the dielectric shift of the Ne matrix. (d) Au_{20}/Ne absorption spectrum (blue) (broad band illumination, spectral detection). Excitation spectrum (black) (monochromatic excitation, monochromatic detection at the main fluorescence line of 739.2 nm). Fluorescence spectrum (red) (excitation at 266 nm, spectral detection). (a) is adapted with permission from ref. 200, Copyright 2019 American Chemical Society, (b) is adapted with permission from ref. 169, Copyright 2018 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, (c) is adapted from ref. 195 with permission from the Royal Society of Chemistry, (d) is adapted from ref. 199 with the permission of AIP Publishing.

general phenomena such as coupling of aromatic conjugation and Ag doping, constructive/destructive interference due to long-range off-diagonal single-particle contributions, *etc.*²⁰⁸ This allowed one to clarify the basic principles of optical spectroscopy of aromatic monolayer-protected alloyed (Ag–Au) clusters and demonstrated how these are uniquely different from those of aliphatic systems. Analogously, Individual Component Maps of Rotatory Strength (ICM-RS) and Rotatory Strength Density (RSD) plots of TDDFT simulated spectra were also introduced as analysis tools of chiral linear response spectra.²⁰⁹ ICM-OS, ICM-RS and RSD allow one to visualize the origin of chiro-optical response in momentum or real space, including signed contributions and therefore highlighting cancellation terms that are ubiquitous in both absorption and chirality phenomena, and should be especially useful in

analyzing the spectra of complex systems, to derive insight and understanding, and eventually rational design, in chiro-optical studies of complex systems²⁰⁹ (Fig. 10a), as explored more recently in ref. 210 where the rotatory strength of chiral gold nanowires was designed and brought from negligible to very large values following the leads of the ICM-RS analysis.

We don't have room to report all the detailed studies conducted on the optical response of MPC. Here, the unique stoichiometry and reasonably well-defined structure have allowed in-depth (and unprecedented in the metal cluster field) investigations of structure/optical-properties relationships, at the same time producing very interesting systems and tracing a path for future studies in parallel fields of heterogeneous and electrochemical catalysis. We refer the reader to recent excellent reviews such as ref. 85, 211 and 212.



The power of *operando* spectroscopies not only in the electrochemical but also in heterogeneous catalysis context was highlighted in ref. 213 where, in addition to systematic high-resolution scanning transmission electron microscopy, a simultaneous monitoring of the Pt dispersion, oxidation state, and CO oxidation activity by *operando* fast XAS and DRIFTS, both combined with MS was reported. Poorly active single Pt^{m+} atoms ($m \geq 2$) were converted gradually and irreversibly into highly active ~1 nm sized Pt^{δ+} clusters ($\delta < 2$) throughout the heating-cooling reaction cycles. As a final note, we stress that it is expected that such *operando* modalities (including other spectroscopies such as IR, Raman, and synchrotron ones such as XANES, EXAFS, *etc.*) will develop quickly in a next future and will provide essential information to achieve the goal of a physical-chemistry-informed rational design of nanocatalytic systems (Fig. 6a and b).

Summary and outlook

Modern society requires a transition to green chemistry, thus green catalysis. Presently available catalysts and catalytic processes, developed in a world in which environmental and energy issues were relatively minor, are for the vast majority not “green”. A societal sustainable growth requires developing atom-economical (optimal use of precious elements), stable (non-leaching undesired toxic compounds or endangered elements), efficient (low-energy spent in high-throughput), selective (low product separation costs) catalysts and catalytic processes, which in turn translates into finding novel catalysts and catalytic sites with all the desired features and performance. In this *scenario*, as promising new materials we focus our attention on metal and metal/oxide clusters at the subnanometer scale, in the size range from 1 to few tens of atoms, deposited on a proper stabilizing substrate. Several reasons justify our choice and make these materials especially appealing to find novel active sites: the unusual structural motifs encountered at this length scale, the strong quantum mechanical confinement effects, the fact that all metal atoms are at the surface and therefore potentially active for adsorbing and activating reactants, the fact that they can be prepared mass-selected and therefore with a precise definition of size and composition. Then, we review the physico-chemical studies that have been conducted on these systems aimed at clarifying and precisely characterizing their static and dynamic structure (Section 2), and the crucial topic of stability (Section 3). The reasoning behind our choice is our belief that a most promising path to the development of new systems is a physical-chemistry-informed rational design, in which the detailed knowledge achieved from monitoring, characterizing, and testing nanocatalyst materials at work enables researchers to understand the limits and problems but also the potentialities of these systems and to find the way to overcome issue and design drastically improved materials as desired. Separate sections have been dedicated to electrochemical systems (Section 4), due to their specific features and issues, and to optical properties (Section 5), in view of the increasingly important field of

photocatalysis. Throughout this review, we tried to emphasize the potential (and often realized) synergy between theory and experiment, and how they can complement each other to achieve at least in part the desired in-depth knowledge and understanding. We have occasionally mentioned topics where further progress is desirable, and is indeed actively pursued by several groups.

To conclude, let us summarize the three main messages and the perspective challenges that can be drawn from the above review. The main messages are that, first, we have shown that the field of subnanometer size- and composition-selected clusters indeed hold great promises in the catalysis field, and that subnanometer clusters can indeed be very active systems. Second, we have reported clear cases of the extreme dependence of such activity on support, size and composition, implying then that it requires a higher degree of knowledge to be able to guide researchers in the discovery of new, active and selective systems, so far often found *via* a trial-and-error approach. Third, the results achieved so far and here extensively reviewed have proved that a physico-chemical approach combining advanced characterization, modelling and controlled synthesis can in principle provide the required degree of knowledge. On the side of perspective challenges, it is clear that this agenda has not yet been fulfilled. Despite the amount of information and sophisticated results so far accumulated, there is yet basically no subnanometer systems for which an ultra-precise knowledge of static and dynamic structure, and of reaction intermediates and their energetics, has been fully achieved. In this respect, we believe that a promising strategy could be to focus the attention of several groups on the most promising systems singled out thus far and to concentrate on these systems a combined effort of the most advanced characterization and modelling tools, also pursuing the development of *e.g.* new microscopy characterization techniques that could overcome the issues of the perturbing effect of the electron beam,²¹⁴ together with new approaches, such as possibly the use of MPC internal standards for increasing the accuracy of the techniques, and many others. An additional challenge is to pursue in a more systematic way the new areas that have been scarcely explored so far, such as electrochemical applications and especially the chiro-optical properties of ultra-nanoclusters. We believe we have provided enough evidence that these actually represent exciting areas of research and possible applications, and that *e.g.* tools to properly model them are rapidly being developed.

We hope that the present excursus, by necessity only partial, will be useful to motivate a larger number of researchers and further studies along this most promising line of research.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

JJ and SV acknowledge support from the European Union's Horizon 2020 research and innovation programme under grant



agreement No. 810310, which corresponds to the J. Heyrovsky Chair project (“ERA Chair at J. Heyrovský Institute of Physical Chemistry AS CR – The institutional approach towards ERA”). The funders had no role in the preparation of the article. AF and SV are also grateful to CNR (National Research Council in Italy) and CAS (Czech Academy of Sciences) for support *via* the bilateral project “Subnanometer Cluster Catalysts for Green Chemistry (SNaGC)”. They also gratefully acknowledge the contribution of the International Research Network IRN on Nanoalloys (CNRS).

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