



Dynamic Processes on Gold-Based Catalysts Followed by Environmental Microscopies

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Abstract: Since the early discovery of the catalytic activity of gold at low temperature, there has been a growing interest in Au and Au-based catalysis for a new class of applications. The complexity of the catalysts currently used ranges from single crystal to 3D structured materials. To improve the efficiency of such catalysts, a better understanding of the catalytic process is required, from both the kinetic and material viewpoints. The understanding of such processes can be achieved using environmental imaging techniques allowing the observation of catalytic processes under reaction conditions, so as to study the systems in conditions as close as possible to industrial conditions. This review focuses on the description of catalytic processes occurring on Au-based catalysts with selected in situ imaging techniques, i.e., PEEM/LEEM, FIM/FEM and E-TEM, allowing a wide range of pressure and material complexity to be covered. These techniques, among others, are applied to unravel the presence of spatiotemporal behaviours, study mass transport and phase separation, determine activation energies of elementary steps, observe the morphological changes of supported nanoparticles, and finally correlate the surface composition with the catalytic reactivity.

Keywords: gold catalysis; Au-based catalysts; environmental microscopies; in situ microscopies; surface science; FEM; FIM; PEEM; LEEM; E-TEM

1. Introduction

In the current technology era, an increasing amount of restrictions is imposed on industrial applications, e.g., fine chemical industry and automotive industry, to safeguard the environment. Moreover, the development of novel, economically cheap and green processes in the chemical industry requires the use of catalytic materials. According to the current estimations, the major part of chemical processes uses a catalytic substance in at least one of the production steps (90% of the processes). Among these, the dominant part uses heterogeneous catalytic materials (solid catalysts principally), while the homogeneous catalysts and biocatalysts are used in only 20% of the catalytic processes. One important and challenging role of any catalyst is to increase the selectivity towards the desired products. This is achieved by providing an alternative reaction pathway, of lower activation energy, for the reaction towards the desired products. In the specific case of heterogeneous catalysis, the materials used can be divided into two types: the noble-metal based catalysts (Rh, Pd, Pt, Ru, Au...) supported on oxides (Al₂O₃, ZrO₂, CeO₂, SiO₂, TiO₂...) or zeolites [1–8]; and the metal-oxide catalysts [9–18]. Regarding the first class of catalytic materials, the deposition method of the active species, principally in case of noble metals, allows highly dispersed nanoparticles (NPs) to be obtained [19,20] resulting in an important number of active sites available for catalytic reactions.

These catalysts include mainly platinum, palladium and rhodium supported on oxide supports and play key roles in many important chemical processes. For instance, platinum-based catalysts are useful in hydro-treatment and complete oxidation processes, whereas palladium and rhodium are crucial components in the abatement of automotive emissions [20-23]. Indeed, the coexistence of these three noble metals in the catalytic converters corresponds to the optimum of catalytic activity at low temperature for the NOx selective catalytic reduction and CO and hydrocarbon oxidation reactions. One of the main problems during this process concerns the emissions of pollutants when the system has not reached its working temperature, also called cold start period. The majority of the emissions are produced after the starting of the vehicle engine when the temperature in the exhaust system is lower than 425 K. For the catalytic oxidation of carbon monoxide, the Pd or Pt-based catalysts do not bring a total conversion at this temperature [24,25]. Understanding how the catalyst works in order to improve its catalytic activity at low temperature is thus of great importance. At the end of the 1980s, Haruta and co-workers showed an outstanding catalytic activity for the CO oxidation at low temperature (200 K) with the use of highly dispersed gold nanoparticles on reducible support. The deposition of nanoparticles with diameters of 2 to 5 nm on reducible support such as Fe₂O₃, Co₃O₄, and NiO exhibits the highest reactivity [26–29]. Together with the research performed by the Hutchings group on the hydro-chlorination of acetylene, Haruta's discovery triggered a great scientific interest for catalysis by gold [30–32]. This is shown in Figure 1 representing the growing trend in term of publications per year involving gold-based catalytic materials and the use of environmental microscopies.



Figure 1. Increasing number of published papers from 1980 to 2017 in the Scopus Scholar database using "gold-based catalysts" and "environmental microscopy" as search keywords (March 2017).

Regarding the evolution of the number of studies using gold-based catalysts, the major part of the scientific publications concern the methods used for the deposition of gold nanoparticles with very narrow and well-defined sizes [33–47]. Besides conflicting papers on the influence of the support materials, the effect of the size on the reactivity of gold catalysts for environmental reactions is well established. The small size of gold nanoparticles induces a modification of the electronic structure of these nanoparticles. In addition to this, a decreased size of the particles increases the dispersion, and accordingly the number of edges and kinks on the metallic particles: such highly uncoordinated surface sites could correspond to the preferential adsorption sites, and their increasing number could then be responsible for the increasing reactivity of the catalytic materials. During the last decade, the research on catalytic materials with gold mainly focuses on the development of bimetallic gold-based catalysts [48-52]. Indeed, the use of bimetallic catalysts allows, to some extent, the size sensitivity of gold catalysts to be diminished. Moreover, the metal particles composed of two metallic components can be used to obtain different and improved catalytic properties as compared to the monometallic catalysts. These new properties are principally due to the synergistic effects between the two metals. Moreover, the catalytic reactivity of the bimetallic particles can be influenced by their structure, the atoms of the nanoparticles can be organised as solid solutions or core-shell structures [51,53–55].

Furthermore, the studies are also focused on the development of synthesis methodologies for the size of the gold nanoparticles deposited on catalytic supports or stabilized in solution [56–59]. The functionalisation of the nanoparticles in solution are mainly used for solid-liquid heterogeneous catalysis [60–62]. Others studies are reported on the immobilisation of enzyme on the gold nanoparticles by the stabilisation of these NPs with π -conjugated dithiol ligands for biocatalytic applications [63–67].

In order to understand the influence of parameters such as the size, the chemical composition or the crystallographic structure of the nanoparticles on the reactivity, the use of environmental microscopy techniques brings outstanding new viewpoints on this catalysis. The development and use of environmental microscopies enables the influence of different atmospheres on the behaviour of the catalytic phase to be analysed. The variety of these techniques makes possible the observation of either the evolution of the active phase or the evolution of the catalytic material at different scales (from nm to mm) and under reaction pressures ranging from 10⁻⁶ to 10⁵ Pa. This review is divided in two parts. The first one concerns the description of the principal environmental microscopies commonly used to study catalytic systems. The second part of this review presents the selected case studies of gold or gold-based catalysts with these environmental microscopy techniques. The main techniques used in these studies and presented in this contribution are Photoemission Electron Microscopy (FEM), Low-Energy Electron Microscopy (LEEM), Field Ion Microscopy/Field Emission Microscopy (FIM/FEM) and Environmental Transmission Electron Microscopy (E-TEM).

2. Environmental Microscopies

Since the early developments of Ultra-High Vacuum (UHV) techniques, many research groups studied model surfaces following the so-called "surface science approach" [68–73]. In this way, the surface structure and their chemical composition can be routinely analysed with high-resolution, providing a better insight into fundamental surface chemistry phenomena. The control and the systematic study of the parameters related to the substrate (crystallographic orientation, density of surface defects, size effect...) is of great importance to get a better understanding of the elementary steps of surface reactions [74]. However, a fundamental problem arises considering the extrapolation of the obtained results towards studies with the applied formulation of catalysts used in industrial conditions. This issue of different working conditions i.e., low pressures and well characterised model surfaces as compared to high pressures and complex formulation of catalysts, is known as the "pressure and materials gap" [75]. Bridging the materials gap can be achieved by using samples with increasing complexity, as illustrated in Figure 2. It then becomes possible to work with samples that approach closely the complexity of industrial catalysts in terms of size, structure and presence of support.



Figure 2. Schematic representation of catalysts samples of increasing complexity studied by different environmental microscopy techniques allowing the materials gap to be bridged. (a) Reproduced from [76]. Copyright 2015, MDPI; (b) and (c) Adapted with permission from [77]. Copyright 2016, American Chemical Society; (d) Reprinted from [78]. Copyright 2015, Springer; (e) Adapted with permission from [79,80]. Copyright 2013, American Chemical Society; (f) Reproduced from [80]. Copyright 2016, Macmillan Publishers Ltd.

In the mid 50s, the importance of studying catalysts in their catalytically active state has already been underlined. Classical microscopy and spectroscopy techniques operate under UHV conditions because of the working principle of the techniques, but also to avoid misinterpretations of the collected data due to the presence of residual gases [81]. During the past decades, major breakthroughs were achieved in the development of surface sensitive analysis techniques working at higher pressures, thereby largely reducing the pressure gap [82], without excessive loss on the lateral resolution [81]. The combination of these techniques with in situ measurements makes them relevant and essential in the field of catalysis.

To study the catalytic systems in details, different samples with a large variety of morphological features, sizes and shapes can be used, as described in Figure 2. Therefore, the use of adapted analysis techniques becomes necessary. The techniques or combination of techniques must be able to resolve structural properties at different levels. As an example, for the analysis of single nanoparticles, atomic resolution is necessary, whereas for studies on foils, the catalytic behaviour can be studied at the mesoscale where the difference in reactivity between crystallographic planes of a few tens of µm may be sufficient. To further explain this idea, Table 1 gathers the basic principles, capabilities and main applications in catalysis of environmental microscopy techniques, e.g., FIM/FEM (Field Ion Microscopy/Field Emission Microscopy), PEEM/LEEM (Photoemission Electron Microscopy/Low-Energy Electron Microscopy), E-SEM (Environmental Scanning Electron Microscopy), E-TEM (Environmental Transmission Electron Microscopy) and STM (Scanning Tunnelling Microscopy). Figure 3 represents the approximate pressure domain of applicability and the size of the resolvable features in the presence of a reactive gas phase.



Figure 3. Application domains of different environmental microscopy techniques in terms of resolvable features and gas pressure permitted in the system under reaction conditions (see Table 1 for a detailed description of the techniques).

2.1. Environmental Microscopies in Catalysis

In the catalysis field, research mainly focuses on two different aspects: from the catalytic reaction point of view and from the catalyst point of view. The first one aims to resolve the mechanism of reaction by determining the presence of intermediate species, kinetic parameters, or the presence of specific dynamics in the system [83]. The second one focuses on the catalytic materials and their evolution during the reaction, for example the presence of surface reconstructions, segregations, sintering... [84,85]. These problems are difficult to tackle with classical microscopies working under UHV conditions, where the samples are typically characterised before and after reaction. Therefore, the presence of transient reconstruction or transient dynamics cannot be probed under UHV conditions. To some extent, environmental microscopies bring solutions as the sample imaging occurs in the presence of a reactive atmosphere. Either the chamber of the microscope is designed for in situ experiments, or special sample holders are adapted to allow in situ characterisations within a conventional microscope [86].

Techniques	Description	Advantage	Inconvenient	Applications in Catalysis	References
Field Ion Microscopy (FIM)	 Field ionisation of imaging gas Resolution: 2 Å Pressure: up to 10⁻² Pa Samples: prepared as tips modelling a single nanoparticle (15–100 nm diameter) 	- Direct imaging - Atomic resolution - Possible coupling with mass spectrometry	 Limited to conductive samples Presence of an electric field Limited pressure 	 Surface structure and reconstruction Spatial and temporal reaction dynamics Structure/reactivity influence Diffusion/front propagation 	[87–93]
Field Emission Microscopy (FEM)	 Field emission of surface electrons Resolution: 15 Å Pressure: up to 10⁻² Pa Samples: prepared as tips modelling a single nanoparticle 	- Direct imaging - Possible coupling with mass spectrometry - Great variety of gases	 Limited to conductive samples Presence of an electric field Limited pressure 	 Spatial and temporal reaction dynamics Structure/reactivity influence Diffusion/front propagation Local activity Non-linear behaviour 	[94–98]
Photoemission electron microscopy (PEEM)	 Photoemission of surface electrons Resolution: 50 nm Pressure: up to 10⁻¹ Pa Samples: single crystals, foils, micro composites 	- Direct imaging - Non-destructive - Possible coupling with mass spectrometry	- Limited to flat samples	 Spatial and temporal reaction dynamics Diffusion and front propagation Structure and reactivity dependence Determination of active sites 	[89,99–104]
Low energy electron microscopy (LEEM)	 Reflection of backscattered low energy electrons Resolution: 2–10 nm Pressure: up to 10⁻² Pa Samples: single crystals, foils, micro composites 	 Tuneable probing depth Combination with LEED (Low Energy Electron Diffraction) 	- Limited to flat samples - Limited pressure	- Surface structure and reconstruction - Film growth - Phase transitions - Reaction dynamics - Front propagation	[102,105–111]
Environmental scanning electron microscopy (E-SEM)	 Detection of secondary and backscattered electrons Resolution: 5 nm Pressure: up to 1000 Pa Samples: porous materials, supported nanoparticles 	- Variable pressure (VP-SEM) - Operational at high pressures	 Conductive samples Time resolution (scanning method) Nature of gases 	- Surface structure - Surface composition - Local activity	[74,112]
Environmental transmission electron microscopy (E-TEM)	 Detection of transmitted electrons Resolution: 2 Å Pressure: up to 10⁵ Pa Samples: porous materials, supported nanoparticles, unsupported nanoparticles 	 Direct imaging Reaction cells Combination with diffraction and spectroscopy 	- Thickness of samples - Beam damage - Data treatment	 Surface structure and reconstruction Growth studies of nanostructured materials Chemical composition changes Atomic diffusion 	[113–120]

Table 1. General description of the main relevant environmental microscopy techniques used for catalysis applications, together with their advantages, inconvenient, and domains of applications in catalysis.

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Ellipsomicroscopy for surface imaging (EMSI)	 Variations of the surface ellipsometric parameters Resolution: 20 μm Pressure: up to 10⁻¹ Pa Samples: single crystals, foils 	- Wide range of pressures (UHV to ambient)	- Light source - Height sensitivity	 Thickness of adsorbate layers, Chemical composition Surface structure Diffusion Spatial and temporal reaction dynamics 	[69,71,103,121]
Reflection anisotropy microscopy (RAM)	- Local variations of anisotropy - Resolution: 10 μm - Pressure: up to 10 ⁻² Pa - Samples: single crystals, foils	- Wide range of pressures (UHV to ambient)	- Light source	 Surface reconstructions Surface adsorption and adsorption energies Diffusion Spatial and temporal reaction dynamics 	[69,103,121,122]
Scanning tunnelling microscopy (STM)	 Measurement of the tunnelling current Resolution: 2 Å Pressure: up to 10⁵ Pa Samples: single crystals, foils, nanoparticles 	- Atomic resolution - Direct real space imaging - Large surface analysed - No pressure limit	- Time resolution (scanning method) - Electric field	- Atomic diffusion - Surface structure - Spatial reaction dynamics - Determination of active sites	[70,73,123–128]

2.2. Basic Principles of Relevant Techniques

As stated in the introduction, this review will focus on the analysis of catalytic behaviours on three main family of techniques: Field Ion Microscopy/Field Emission Microscopy (FIM/FEM) to study the catalytic behaviour on a single nanoparticle, Photoemission Electron Microscopy/Low-Energy Electron Microscopy (PEEM/LEEM) to investigate catalytic processes on flat samples, and Environmental Transmission Electron Microscopy (E-TEM) allowing the analysis of unsupported, supported and self-supported catalysts. These techniques are described in more details in this section, together with striking examples on their possibilities.

2.2.1. Field Emission Microscopy FEM/Field Ion Microscopy FIM

Field emission techniques such as Field Emission Microscopy (FEM) and Field Ion Microscopy (FIM) are used to study samples prepared as sharp tips. The extremity of the tip-sample is a highly curved surface, which is a good model of a catalytic nanoparticle with a diameter of 20–100 nm. [98,129]. Pure gold samples are routinely prepared by electrochemical etching of a gold wire using KCN solutions, but also in CaCl₂, NaCl, KCl, and H₂SO₄ + HNO₃ solutions [130–134]. Alternatively, inert metallic tips such as tungsten can be prepared and then covered with a few atomic layers of Au by vacuum evaporation or laser deposition [135–137]. In the case of alloys, the samples are prepared either by electrochemical etching of the alloy wire [138], or by evaporation of the two metals followed by annealing to ensure the homogenisation of the alloy-layer [139].

Field emission techniques are not widely used these days as surface science techniques for catalysis applications, probably because of the scarcity of these instruments. However, the unique features of these techniques allow in situ catalytic studies in the presence of a reactive atmosphere as well as characterisation under high vacuum conditions, for pressures ranging from residual ones to 10^{-2} Pa and for cryogenic temperatures (≈ 50 K) to ≈ 900 K. In FIM mode, the application of an electric field between the sample and the detector screen induces the process of field ionisation of an imaging gas. The ions are accelerated by the field towards the detector screen where image formation occurs. The micrographs provide the crystallographic surface structure of the sample, with an atomic lateral resolution, as seen in Figure 4a [140]. Atoms of the imaging gas are ionised over the most protruding atoms of the surface (kink and step sites). With this, a magnification of 10^6 is typically obtained. The FIM images show an excellent agreement with a ball-model of a model hemisphere (Figure 4b). Morphological changes such as facetting or missing-row reconstructions can easily be imaged during their formation [98].



Figure 4. Analysis of surface reconstruction by FIM/FEM: (**a**) FIM micrograph of a Ni crystal tip and the corresponding Miller index annotations. The magnification corresponds to ~10⁶ on a detector screen of 3 inches diameter. Conditions of acquisition: T = 61 K, F = 3.5 V/Å; (**b**) corresponding ball model where withe balls represent kink and step sites; (**c**) FEM micrograph of the same crystal under

hydrogen atmosphere. Conditions of acquisition: T = 873 K, $P_{H2} = 1 \times 10^{-4}$ Pa illustrating the facetting process; (d) corresponding ball model of the facetted structure; (e) FEM micrograph of the crystal under C₂H₂ atmosphere. Conditions of acquisition: T = 873 K, $P_{tot} = 2.5 \times 10^{-4}$ Pa; (f) schematic representation of the growth of the open (113)-facets. Adapted with permission from [140]. Copyright 2009, American Chemical Society.

In FEM mode, a negative polarisation of the sample as compared to the screen-detector initiates the emission of surface electrons, as can be observed in Figure 4c,e. The field emission depends on the work function of the crystallographic orientation of the facets. In Figure 4c–f, the work function variations and changes in morphology of the sample after the formation of extended facets can be followed during the early stages of C-nanotube formation on a Ni tip-sample. Since the work function depends on the presence of adsorbates, variations of the work function due to adsorption, dissociation or desorption of different chemical species lead to variations of the brightness pattern, allowing the presence of adsorbates to be followed, and therefore, the presence of specific dynamics at the surface [141].

Although field emission techniques have barely changed since their invention in the 1950s, they represent a powerful tool in surface science and heterogeneous catalysis. The unique possibility to image reaction dynamics make them a fruitful technique towards the understanding of fundamental phenomena occurring at the surface of catalytic particles. For example, processes such as nonlinear behaviour, surface explosions [142] or oscillations [143–146], but also reconstructions, morphologyreactivity relationship and diffusion [147] can be studied with these techniques. Moreover, these techniques can be coupled with mass spectrometry and are known as Pulsed Field Desorption Mass Spectrometry (PFDMS), or sometimes referred as the 1-Dimensional Atom Probe (1DAP) [140,148]. The general idea is to apply electric field pulses to the tip, with a specific frequency, so as to initiate field desorption of surface species. Desorbed species are analysed by time-of-flight mass spectrometry. By varying the pulsed frequency, it is possible to follow the evolution of chemical species during the progress of the reaction and to determine kinetic parameters. A similar concept is the base of the 3-Dimensional Atom Probe/Atom Probe Tomography (3DAP/APT) principle [149,150]. The main idea here is not to analyse the single layer of adsorbates, but rather the distribution and the nature of atoms within the sample [151]. The system is equipped with a position sensitive detector, allowing a tomographic reconstruction of the sample, to be obtained with atomic resolution in the z-direction, together with the chemical identification of the ions. Analysis of the surface segregation of catalytic nanoparticles following exposure to reactive gases [85,88] is thus possible. It has to be noted that this technique requires UHV conditions. We also mention that the FEM/FIM instrument can be coupled with spectroscopy (Field Ion Appearance Energy Spectroscopy [152]).

2.2.2. Photoemission Electron Microscopy PEEM/Low Energy Electron Microscopy LEEM

Photoemission Electron Microscopy (PEEM) is based on the illumination of a sample with UVlight, and detection of the resulting photo-emitted electrons. The contrast arises from the local differences in the yield of electron-emission, which in turn depends on the local work function. As in the case of FEM, the local work function depends on the presence of adsorbed species. Qualitative measurements of adsorption and dissociation phenomena are thus possible [143,153]. Low Energy Electron Microscopy (LEEM) is based on the emission of electrons when the sample is exposed to a flux of low energy electrons. The impinging electrons give rise to the emission of backscattered electrons from the analysed surface allowing for real space imaging of the surface structure. By using the back focal plan as objective lens, it is easy to switch to diffraction mode and obtain diffraction patterns with the technique known as Low Energy Electron Diffraction (LEED). The contrast arises mostly from structural differences [105,154,155].

LEEM and PEEM are powerful tools to analyse numerous physical phenomena such as growth dynamics, the study of magnetic domains or adsorption processes, but they are not able to perform chemical characterisation. Because of the relatively low maximum pressures and resolution under in

situ conditions, the use of PEEM as a surface science technique for catalysis applications is limited to a few studies such as the determination of diffusion coefficients, and some reactions on composite surfaces to study the synergy between heterogeneous domains [156]. Another well-known example is the formation of spatiotemporal patterns during the CO oxidation on Pt surfaces, as can be observed in Figure 5. Propagation fronts corresponding to different surface compositions are clearly discernible during in situ imaging. Indeed, the contrast is obtained via changes in the work function, which depends on the nature of the adsorbates. The work function is different in the presence of O(ads) or CO(ads), and the transitions from one domain of stability to another can be observed via the propagation of waves [157].



Figure 5. PEEM study of surface reaction and spatiotemporal patterns formation: Temporal evolution of diffusion patterns during the oxidation of CO on Pt surface. Conditions of acquisition: T = 448 K, $Po_2 = 4 \times 10^{-4} \text{ Pa}$, $P_{CO} = 4.3 \times 10^{-3} \text{ Pa}$. Reprinted from [158]. Copyright 1993, AIP Publishing.

One of the limitations of the PEEM is the lack of chemical information during imaging. However, by exploiting the synchrotron radiation, XPEEM (X-ray Photoemission Electron Microscopy) has been developed to image surfaces with chemical sensitivity. It is then possible to study the evolution of the surface composition during the ongoing processes [106,159]. PEEM can be combined with mass spectrometry to follow the global gas-phase composition during the catalytic reaction [160]. This permits catalytic activities of different catalytic materials to be compared with different crystallographic orientations [161]. Further improvements of the technique include the combination of PEEM with LEEM which add the structure sensitivity and diffraction to the capabilities of the PEEM instrument, and is known as SPELEEM (Spectroscopic Photoemission and Low-Energy Electron Microscope) [155,162–164]. The availability of XPEEM in synchrotron facilities increased the interest of this technique and derivate for catalysis applications.

Also, the samples used by PEEM techniques are single crystals, foils or composite surfaces. The deposition of homogeneous layers can be performed by evaporation, and the preparation of composite surfaces is done via lithography techniques.

2.2.3. Environmental Transmission Electron Microscopy E-TEM

Transmission Electron Microscopy (TEM) is widely used in the catalysis community to determine the structure, morphology and, with complimentary techniques, the chemical composition of materials used in catalysis [82,165]. Traditional TEM microscopes can be modified to respond to the latest demands in materials science and catalysis [82,165]. A critical step towards the development of microscopes operational at high pressures is the need to have high pressures of reactive gases around the sample and high vacuum conditions in the rest of the apparatus. For this, a specific design of the microscope can be used, or alternatively, specific sample holders that can be easily adapted on the non-environmental TEM are available [166,167]. This comes from the fact that it is strictly necessary to confine the gas atmosphere in the vicinity of the sample in order to avoid a reduction of the mean free path of the electrons altering of the results. The use of E-TEM allows studying materials with high resolution and in temperature and pressure conditions close to those used in applied formulations of catalysts present the advantage of being active at relatively low temperature, enabling the study of catalysts under reactive environment without significant loss of spatial

resolution. This allows transient phenomena to be imaged that may be of great importance to understand the catalytic processes.

In E-TEM mode, high energy transmitted electrons are detected after interacting with very thin samples to produce images in direct space [84]. E-TEM allows to follow changes of the catalyst, e.g., structure, size, sintering... [168], and complementary analyses can be performed by diffraction—or spectroscopic techniques-Energy Dispersive X-ray (EDX) and Electron Energy Loss Spectroscopy (EELS). EDX can give information about changes of the chemical composition of the sample due to phenomena like surface segregation [169], while EELS is used to determine the chemical composition of the system, as well as surface bonding properties [84,170,171]. Both EDX and EELS can achieve atomic resolution analysis with the new generations of TEM and under specific conditions. A relatively wide variety of samples can be analysed by TEM: supported or unsupported particles, powders, structured materials... [28,76] which makes this tool useful for catalysis studies. An example of the capabilities of the E-TEM are the morphological changes occurring on a nanoparticle of Cu (from Cu/ZnO catalyst) under H_2 and $H_2 + O_2$ environments, as is presented in Figure 6. Other processes such as particle growth due to sintering [86], as well as nanotube growth can also be followed. The parallel characterisation of the catalyst surface as well as the reactive gas phase under working conditions, allows conclusions to be drawn on the structure reactivity relationship of catalytic systems [84,172].



Figure 6. E-TEM analysis of catalyst reconstruction under reactive conditions: (**A**,**C**,**E**) Micrographs of Cu nanoparticles deposited on ZnO captured with in situ TEM analysis, (**B**,**D**,**F**) together with their corresponding Wulff constructions illustrating the reactive gas phase induced shape transformations of the Cu nanoparticles. Conditions of acquisition: T = 493 K, (**A**) $P_{H_2} = 150$ Pa; (**C**) $P_{tot} = 150$ Pa with $P_{H_2}/P_{O_2} = 3.1$; (**E**), $P_{tot} = 500$ Pa with $P_{H_2}/P_{O_2} = 19.1$. Reprinted from [173]. Copyright 2002, AAAS.

3. Case Studies

This section is dedicated to an in-depth analysis of Au and Au-based catalytic systems imaged by PEEM/LEEM, FIM/FEM and E-TEM instruments. The combination of those three techniques covers a wide range of catalytic materials of increasing complexity, from single crystals to structured materials. Most of these studies are also combined with spectrometry/spectroscopy techniques to get a better insight of the systems, and are thus also discussed as a complement of the microscopic part of the studies. The remaining of this section is divided as follows: The first part is dedicated to the study of processes on single crystals by PEEM/LEEM. CO oxidation on Au-covered Pt is analysed by PEEM to highlight the presence of spatiotemporal behaviours; meanwhile the O₂ + H₂ reaction is probed on Au- and Au + Pd-covered Rh by LEEM and XPEEM to study the mass transport occurring under reactive environment. The second part corresponds to the study of processes on a single grain of catalyst by FEM/FIM. This includes the FIM imaging of CO/O₂ interaction and water gas shift reaction on Au tips; the experimental determination of activation energies on the H₂O/Au system by FEM; and the study of NO adsorption and reduction on Au-Pd alloys by FEM, correlated with the

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Pd segregation studied by atom probe. The third part focuses on the study of processes on supported and self-supported catalysts by E-TEM. This comprises the study of dynamic behaviours occurring during CO oxidation on supported Au nanoparticles, i.e., the coalescence of nanoparticles, the morphological changes of both the catalyst and the support, as well as their correlation with the catalytic activity. The study of catalytic oxidations on nanoporous gold samples will also be addressed. Finally, the main conclusions are drawn.

3.1. Study of Processes on Single Crystals by PEEM/LEEM

The selected examples presented in this section will describe the presence of non-linear spatiotemporal behaviours on Au-covered Pt surfaces, as well as the observation of mass transport and phase separation on Au-covered Rh surfaces. Finally, the spectromicroscopy of a model water-gas shift catalyst is presented.

3.1.1. Processes on Au-Covered Pt Single Crystals: Influence of Au on the Presence of Kinetic Instabilities on Pt(110)

The PEEM technique is particularly suited to study, at the mesoscale, how the reactivity of surfaces may be affected by the presence of adjacent areas with different reaction properties. CO oxidation over platinum surfaces has been extensively studied in in the past [121,158,174–177] and consists of a solid base to study how the dynamics can be affected. For example, during CO oxidation on Pt(110), at a fixed temperature and oxygen (O₂) pressure, if the CO pressure is continuously increased, different dynamics are observed: a stable state covered by O(ads) species, an excitable state, and a stable state covered by CO(ads) species. The main feature of the excitable regime consists of the propagation of reaction-diffusion fronts, the presence of spiral waves and solitary waves [158]. The properties of these patterns are governed by parameters such as the sticking coefficient of both O₂ and CO, or the diffusion coefficients. The examples presented below will show how the spatiotemporal patterns occurring during CO oxidation are affected by the presence of Au at the surface at different coverages. The presence of gold is supposed to continuously affect the kinetic parameters; meanwhile the mesoscale homogeneity of the medium remains unaltered.

On bare Pt(110), CO is oxidised to CO₂, which is directly released in the gas phase. A reversible surface transformation $1 \times 1 \rightleftharpoons 1 \times 2$ is driven by the CO coverage. The O₂ sticking coefficient is higher on the 1×1 surface than on the 1×2 reconstructed surface, which is in turn responsible for the occurrence of kinetic instabilities for given values of the control parameters, i.e., temperature, O₂ and CO pressures. These instabilities can be observed via the changes of the work function: ϕ is increased in the presence of CO(ads), and increased even more in the presence of O(ads): O-covered regions thus appear dark and CO-covered regions appear grey.

The presence of deposited Au on Pt(110) surface affects the behaviour of CO oxidation [178–180]. Indeed, gold atoms cause a uniform modification of the kinetic parameters. The presence of Au locally inhibits the adsorption of CO by acting as a site blocker (but does not change its sticking coefficient); and inhibits the dissociative adsorption of O₂, since it requires several neighbour Pt atoms for the dissociation to occur. According to the authors, the presence of Au does not affect the properties of neighbouring Pt atoms. For an Au-coverage of 0.05 ML (monolayer), at the mesoscale, the gold atoms are uniformly distributed at the surface of Pt(110), and the properties of the spatiotemporal patterns are modified under reaction conditions. The main modifications are: (1) the excitable regime is reached at lower CO pressure; (2) the velocity of the front propagation is reduced, and (3) the density of nucleation centres for the formation of waves increases. This can be observed in Figure 7 corresponding to a sample with an Au-covered domain (left part) and a bare Pt domain (right part) at a fixed temperature and O₂ pressure. In Figure 7a, at low CO pressure, both domains are O-covered, represented by dark regions in the PEEM image. When the CO pressure is gradually increased, one can observe—in Figure 7b,c—the formation of CO domains of higher brightness on the Au-covered surface, but not on the bare Pt. In Figure 7d, spirals are formed on the Au-covered domains, and the emergence of CO domains on the Pt side is observed. The Figure 7e corresponds to a parametric region where both Au-covered and bare Pt are in a reactive state, but presenting

different wavelengths. Finally, in Figure 7f, the Au-covered region is fully CO-covered, but O-waves are still present on the Pt region.





Figure 7. Study of different reactive behaviours in the presence and absence of Au: PEEM snapshots during the CO oxidation on 0.05 ML Au-covered Pt(110) adjacent to a bare Pt(110) area. Conditions of acquisition: T = 460 K, $Po_2 = 4 \times 10^{-2}$ Pa. The pressure is increased from (**a**) $Pco = 1.0 \times 10^{-3}$ Pa: two regions in the O-covered stable states; (**b**) $Pco = 1.7 \times 10^{-3}$ Pa: formation of CO(ads) domains on the Au-covered side; (**c**) $Pco = 1.9 \times 10^{-3}$ Pa: formation of CO fronts nucleating by diffusion across the boundary; (**d**) $Pco = 3.0 \times 10^{-3}$ Pa: formation of spirals on the Au-covered domain and formation of fronts on the Pt domain; (**e**) $Pco = 4.65 \times 10^{-3}$ Pa: spiral patterns on both domains; (**f**) $Pco = 5.0 \times 10^{-3}$ Pa: Au-covered domain fully covered by CO(ads) and presence of O(ads) waves on the Pt domain. Reprinted from [179], Copyright 1995, AIP Publishing.

As discussed, the presence of Au decreases the sticking probability of O_2 at the surface and blocks, to a lesser extent, CO adsorption. Since the transition from the monostable O-covered region to an excitable regime is balanced by the coverage of O(ads) and CO(ads), the critical balance will be reached for a lower CO pressure, explaining the presence of different dynamics on Au-covered and bare Pt in Figure 7b,c. The same explanation stands for the transition from excitable to monostable CO-covered presented in Figure 7f at higher CO pressure. Au atoms on Pt represent an obstacle for CO migration, with the direct effect of decreasing the CO diffusion coefficient. Therefore, the velocity of the front propagation is lower: 2.2 μ m/s in the [110] direction on the Au-covered Pt as compared to 2.8 μ m/s on bare Pt. This effect has been studied at different temperatures and shows the same trend of reduced velocity, as can be seen in Figure 8.

There is a gradual decrease of the available area for gas adsorption while the Au coverage is increased, which further affects the local dynamics. The pressure range to observe patterns in excitable state becomes narrower with increasing Au coverage. For example, at a coverage of 0.3 ML, both O₂ and CO adsorption is still possible, but the direct transition from O-covered to CO-covered is observed without any pattern formation. The effect of the presence of gold and the inhibition of reconstruction have to be accounted for in the full description of the catalytic system [121,179,181]. It can be concluded that surface inhomogeneities at the atomic scale affect the kinetic parameters averaged over mesoscopic length scale on which the surface still appears homogeneous [180].



Figure 8. How the presence of Au atoms affects the kinetic parameters during CO oxidation on Pt: Propagation speeds for CO fronts on bare and 0.05 ML Au-covered Pt(110) as a function of CO pressure with a base oxygen pressure of $Po_2 = 4 \times 10^{-2}$ Pa, at temperatures for conditions of dynamic bistability. This graph highlights the decrease of the CO diffusion coefficient due to the blocking effect of Au. Reprinted from [179], with the permission of AIP Publishing.

Another interesting effect observed in Figure 7c is the behaviour at the boundary. As a function of the pressure ratio, Au-covered and bare Pt domains present different regimes of stability and CO-waves produced in the Au-covered domain (1) cannot cross the border (the "forbidden area"); (2) can cross the border but are rapidly extinguished because of the O(ads) effect; or (3) can cross the border. The probability for the CO wave to cross the boundary obviously increases with higher CO pressures. It was noticed that there is a preferred creation of spirals at this boundary, which might be due to the higher density of defect sites acting as centres for nucleation.

3.1.2. Processes on Au-Covered Rh Single Crystals

The above described example mainly focused on the study from the viewpoint of the catalytic reaction. However, the catalytic system is in constant evolution and the catalyst may also undergo structural and compositional changes during the reaction. This can be probed by LEEM and XPEEM techniques as is it described below in the case of Au-covered Rh surfaces.

Reaction-Induced Patterning of Au/Rh(110) Surfaces

In this work, Au was deposited on O-precovered Rh(110) surfaces [153,182], and then the $H_2 + O_2$ reaction was monitored by LEEM. Characterisation of the sample proved that the Au, with a coverage of 0.5 ML, is initially evenly distributed onto the Rh surface. The reaction between H_2 and O_2 on Rh forms water which desorbs immediately under the reaction conditions (T = 550 K). Gold is chemically inert for the reaction and is thus used for its site-blocking effect. Starting from a homogeneously oxidised Au-covered surface, hydrogen is introduced in the chamber so as to ignite the formation of a reduction front. At this point, both O_2 and H_2 pressures are stabilised. Micro-XPS was used to obtain a quantitative chemical information of Au and O distribution, and the contrast variations observed in LEEM were in accordance with the Au 4f chemical maps. Therefore, LEEM was used to observe both the initiation of front propagation and the evolution of the surface under reaction conditions, as can be seen in Figure 9a where bright regions correspond to the reduced surface, dark regions to the oxidised surface, and grey regions to the initial surface.



Figure 9. LEEM and micro-XPS study of front propagation and mass transport: (**a**) LEEM images under O₂ + H₂ reaction illustrating the ignition and propagation of reduction/oxidation fronts. Field of view = 20 µm. Conditions of acquisition: T = 550 K, $Po_2 = 2.8 \times 10^{-5}$ Pa, $PH_2 = 3.3 \times 10^{-5}$ Pa; (**b**) Au 4f_{7/2} map acquired during front propagation illustrating the lateral inhomogeneity induced by the reduction/oxidation front; (**c**) Au 4f spectra measured in the different regions presented in (**b**). Reprinted from [153]. Copyright 2004, Elsevier.

When hydrogen is introduced in the chamber, the nucleation and growth of elliptical fronts can be observed (Figure 9a). This anisotropic behaviour is supposed to find its origin in the anisotropic diffusion of adsorbate species on the substrate [183]. These bright regions correspond to the surface reduced by the first reaction front. The dark regions correspond to the propagation of oxidation fronts, generated at the centre of the reduction fronts. During the $O_2 + H_2$ reaction on bare Rh(110), only reduction fronts are observed, and reaction fronts initiate a transition from an inactive 'Osaturated' phase, to an active almost 'O-free' surface [183,184]. In the case of Au-covered Rh, micro-XPS analysis revealed that the reduced area is Au-enriched and O-free, and the oxidised area is Audepleted (Figure 9b,c). In the reduced area, the accumulation of Au grows continuously as the front progresses, up to a value of ≈ 0.9 ML corresponding to the maximum coverage for that system. After some time, the reaction front slows down and a quasi-stationary state can be observed (Figure 9a at 5500 s). This patterned surface, consisting of Au-depleted and Au-enriched regions is preserved in an oxidising environment, proving that the mobility of Au on oxidised surfaces is relatively low. However, the patterned surface is rapidly destroyed in a reducing environment, a sign of the high mobility of Au on oxygen-free surfaces. The Au mass transfer process observed in this system can be explained by the presence of an energetically favourable configuration: the reorganisation of the catalyst surface into "Au-poor/O-rich" and "Au-rich/O-poor" phases. We can assume that optimisation of the energy of the system is obtained if Au-adatoms bond directly to the Rh surface, inducing a rearrangement of the oxygen layer. The presence of an active gas may promote surface diffusion of metal adatoms, and this work proves that the propagation of reduction/oxidation fronts destroys the lateral homogeneity of the Au adlayer, which results in the development of a patterned surface composed of Au-depleted and Au-enriched regions.

The initial surface composition is important in the development of such patterned structures. For example, if Au is deposited on a clean Rh(110) surface [105] as compared to a O-precovered Rh(110) surface, the Au mass transport starts with the ignition of oxidation fronts, and bright domains merge to larger islands, eventually forming segregated phases (Figure 10). The aggregation of Au into the bright islands is confirmed by XPEEM.



Figure 10. LEEM imaging of the formation of a segregated phase due to the gold mass transport observed under reaction conditions. Reprinted with permission from [182]. Copyright 2005, American Chemical Society.

The presence of a segregated phase is accompanied with a different reactivity. Indeed, reduction fronts are solely ignited in the Au-poor regions. The fronts then expand up to the Au-rich regions where they propagate further, but with a slower velocity. The two phases can thus be seen as independent microreactors. This proves that complex interfaces can develop, with different reactivity, even though the species responsible are chemically inert.

Chemically Frozen Phase Separation

One step further into the complexity of phase separation on catalytically active surfaces is the use of a mixed-layer of adatoms. For this, Au and Pd were deposited onto Rh(110) surfaces [162,164,185,186] and studied under H₂ + O₂ reaction conditions at different temperatures. It has been found that the water formation reaction triggered the presence of stationary 2D metallic and oxygen structures. In this case, because of the formation enthalpy of binary alloys (Au-Rh, Au-Pd and Au-Pd), which is only negative for Au-Pd [187], and oxygen adatoms are more strongly bonded on bare Rh, the patterned structure is composed of mixed Au-Pd and O islands. This can be seen on the XPEEM images in Figure 11a. It should be mentioned that O-covered regions may still contain very small amounts of metal adatoms depending on the reaction conditions. This lamellar structure results from chemically frozen spinodal decomposition, and is energetically favourable, but kinetically hindered in the absence of reaction. Once formed, such structures can be preserved if the temperature is quenched down to 300 K and the reaction is interrupted, proving the structure to be stationary rather than transient.



Figure 11. Chemically-frozen phase separation studied by LEEM and XPEEM: (**a**) LEEM and elemental XPEEM of the lamellar structures corresponding to separated Au-Pd and O islands. Conditions of acquisition: T = 820 K, $P = 5.3 \times 10^{-5}$ Pa, $P_{H_2}/P_{O_2} = 0.5$, $\Theta_{Au+Pd} = 0.5$ ML; (**b**) LEEM images of stationary structures produced under different gas pressures. Conditions of acquisition: T = 820 K, P = 100 K, $P_{H_2}/P_{O_2} = 0.8$, $\Theta_{Au+Pd} = 0.75$ ML; (**c**) Plot of the structure wavenumber (in the [110] direction) as a function of the pressure of the reactants. Adapted with permission from [185]. Copyright 2006, American Chemical Society.

The more important feature of such lamellar structures is that they are adaptive, which means that they can be created, preserved and reversibly modified by tuning the reaction conditions (either the gas pressure or the temperature). An example is presented in Figure 11b, where the wavelength of the structure is modified by increasing the pressure of the reactants while the pressure ratio remains the same. This wavelength, calculated along a specific direction, obeys a power law over two orders of magnitude in the pressure of the reactants (Figure 11c). Since the wavelength depends on the reaction conditions, the reaction rate could be finely adjusted by using variations of the pressure or temperature as a tool to control the wavelength, and therefore design specific microstructures at the surface of the catalyst. This also corresponds to an experimental proof of the growth of spatial modes of modifiers microsctructures in the presence of a surface chemical reaction [188,189].

3.1.3. Spectromicroscopy of Au-Supported Nanoparticles

One noticeable study performed on Au-based systems is the spectromicroscopy of Ausupported nanoparticles [190]. As described above, the use of SPELEEM instrument adds the structure sensitivity to the capabilities of the PEEM. Recent work [190] has been performed on gold nanoparticles supported on CeO_{2-x}(111) crystallites, themselves deposited on a Rh(111) substrate. This material can be used as a model catalyst for the water-gas shift reaction. Au/CeO₂ systems have been, and still are, widely used for catalytic applications such as the water gas shift reaction, CO and volatile organic compounds (VOCs) oxidation, and NO₂ reduction [33,191–194]. Therefore, the understanding of the redox behaviour of the gold/ceria system remains a crucial step in the fundamental understanding of its catalytic activity.

Because the resolution of XPEEM and LEEM techniques is limited to a few tens of nanometres, Au particles cannot be resolved, but spectroscopic information can still be obtained. For example, it was shown that as the coverage of gold increased, the ceria was reduced, which is consistent with charge transfer from the gold particles to the oxide support. The effect of the presence of an oxygen atmosphere and the beam-induced reduction of the sample was also successfully investigated. To examine this effect, Au 4f region is probed and a shift to higher binding energy is observed for Au particles located on ceria as compared to those on Rh. The use of LEEM imaging can be correlated to XPEEM imaging of gold to discern the ceria structures, as can be seen in Figure 12. By using energy-filtered XPEEM, it is possible to distinguish the Au signal originating from the ceria crystallites, or from the Rh support. This study shows promising results for the future of PEEM and derivate techniques for catalytic applications.



Figure 12. Spectromicroscopy of Au nanoparticles: XPEEM, LEEM and XPS analysis of Au nanoparticles deposited on CeO_{1.67}/Rh(111) showing the effect of beam-induced reduction of the ceria. Ceria islands can be discerned in XPEEM and LEEM images. The resolution of the technique allows observing the presence of Au on both the ceria and the Rh support. (**a**–**c**) analysis under $Po_2 = 5 \times 10^{-5}$ Pa; (**d**–**f**) analysis under UHV conditions. Reprinted with permission from [190]. Copyright 2014, American Chemical Society.

3.1.4. Perspectives

In the past and current research, Au-based samples were not widely studied by PEEM/LEEM techniques. Only a few systems were imaged: pure Au [105], Au-Si [105,156,195], as well as Au-Mo [105]. However, with the increasing interest in Au-based alloys for catalytic applications, model samples were produced to study the presence of specific dynamics, interfacial phenomena, the possible presence of subsurface species, as well as phase separation processes.

3.2. Study of Processes on a Single Grain of Catalyst by FEM/FIM

The different examples presented in this section will show (1) how the coupling between FIM and mass spectrometry allows the mechanism of reactions to be resolved, (2) how FEM can be used to determine activation energy for adsorption/desorption/diffusion processes, and (3) how the coupling of FIM with a position-sensitive detector allows the surface composition to be correlated with the reactivity.

3.2.1. Processes on Au Tips

Propagation Waves during CO/O2 Interaction

Gold catalysts are known to be active for the low temperature CO oxidation [27,47], and it was thus straightforward to study this system by field emission techniques. One of the advantages of this technique is to allow the study of catalytic reactions in the absence of any influence of the oxidic support, which is of great importance for the fundamental understanding of the catalytic system. In addition to this, the behaviours observed are influenced by the presence of a steady electric field which could mimic conditions encountered in studies over Au^{b+}/metal-oxide catalysts [196]. In Figure 13a, we can observe a typical Au sample prepared and imaged by field ion microscopy. The study of CO/O₂ system was performed on such samples [197]. The authors mainly observed the presence of a propagating wave during the imaging of the gold nanoparticle in the presence of CO + O₂ gas mixtures at 300 K. It has to be noted that FIM work during CO oxidation on Pt field emitter tips proved that oxygen is used as an imaging gas [198]. Since FIM imaging lacks chemical analysis, further studies coupling the FIM with mass spectrometry [148] were performed for different systems of increasing complexity: CO/Au [199], O₂/Au [200] and finally CO + O₂/Au [201].

The study of CO adsorption on Au tips [199] suggests the formation of gold mono- and dicarbonyls, as shown in Figure 13b representing the mass spectra obtained during CO adsorption at room temperature. Gold carbonyls are indeed observed, in the presence of high electrostatic field, as AuCO⁺, AuCO²⁺ and Au(CO)²⁺ species. The mass spectra have been acquired over a region of approximately 400 atomic Au sites with a relatively high density of kinks, as shown in Figure 13a (black circle i.e., the probed area). The formation of carbonyls appears to be field-dependent, in agreement with the field-dependence of CO bonding energy to Au(111) [202], and DFT calculations concluded that the local field-induced surface charge accumulating on gold atoms stabilises the carbonyls on highly facetted gold nanostructures. It is suggested that in the absence of an external field, the local charges can be present at kinks and steps sites of small clusters, explaining the sizedependent reactivity of Au clusters [29,203].



Figure 13. Analysis of a single Au nanoparticle by FIM and 1DAP: (**a**) Field Ion Microscopy (FIM) image of a (111)-oriented Au sample where the atomic resolution can be observed on steps and kinks and inside flat planes of small sizes. The dark circle corresponds to a typical region of analysis for studies by mass spectrometry. The magnification in FIM corresponds to ~10⁶ on a detector screen of 3 inches diameter. Conditions of acquisition: T = 50 K, $P_{Ne} = 10^{-3}$ Pa. Reprinted from [204], Copyright 2006, Elsevier; (**b**) Mass spectra of an Au sample during interaction with CO in the presence of an external electric field. Mono- and di-carbonyls are formed on the sample. The region of analysis corresponds to ~400 atomic Au sites. Conditions of acquisition: T = 300 K, $P_{CO} = 10^{-3}$ Pa, steady field $F_R = 2.1$ V/Å, pulsed field $F_P = 1.1$ V/Å, frequency of the field pulses f = 1.67 kHz. Reprinted from [199]. Copyright 2003, AIP Publishing.

The next step towards the understanding of the observed propagation wave is the study of the O₂/Au system [200]. Previous work showed that the dissociative oxygen chemisorption on pure gold model nanoparticle is caused by the presence of surface impurities [205–207]. On Au tips, surface impurities are supposed to be absent due to the cleaning process following the sample preparation. Experiments by PFDMS showed that the adsorption of molecular and atomic oxygen on Au tips does not occur at pressures lower than 10^{-2} Pa in the temperature range 300–350 K. The dissociation of oxygen cannot be forced by applying an electric field up to 60 V/A. AuO₂⁺ species are observed in the mass spectra when the applied field is close to the onset of Au field-evaporation, which means that once formed, field evaporated Au⁺ ions undergo complexation by O₂ in the very vicinity of the "takeoff" location. The scenario is reinforced by the enhanced impingement rate of oxygen molecules in the presence of a strong inhomogeneous electric field. O²⁺ species are also present in the PFDMS analysis, but their presence is independent of the temperature, reaction time and probed region. The presence of O_{2⁺} thus reflects gas phase ionisation of oxygen close to the tip rather than molecular adsorption. These results are in agreement with DFT models proving that no O₂ dissociation will occur in the presence of an external electric field [208]. The size of the Au tip samples is of several tens of nanometres diameter and notoriously not active in catalysis for CO oxidation.

As stated, the interaction of CO and O₂ on an Au sample at room temperature induces the presence of a propagation wave, as it can be seen in Figure 14. Oxygen is first introduced in the reaction chamber, followed by CO, and the surface is imaged with a field of 1.2–1.5 V/Å. The initiation of the wave occurs close to the central pole, where the field is the highest, and then propagates away from the nucleation region down to the shank of the tip [201]. It has been shown that the application of an external field induces the presence of gold-carbonyls [199] but does not induce the adsorption/dissociation of oxygen species [200]. It is thus rather unlikely that the propagation wave corresponds to CO oxidation.



Figure 14. Imaging a chemical wave under reaction conditions by FIM: (**a–e**) Series of snapshots during the interaction CO/O₂ on an Au tip imaged by FIM during the ongoing process. A chemical wave is observed with its initiation at the apex part of the tip and propagation to the shank of the tip. The magnification in FIM corresponds to ~10⁶ on a detector screen of 3 inches diameter. Conditions of acquisition: T = 300 K, $Pco = 4 \times 10^{-5}$ Pa, $Po_2 = 10^{-2}$ Pa, imaging field = 1.4 V/Å; (**f**) Guide to the eye of the wave superimposed with a ball model of the sample. The black region corresponds to the regions probed by mass spectrometry (see Figure 15). Reproduced from [196]. Copyright 2007, John Wiley & Sons, Ltd.

To support this hypothesis, PFDMS was used to probe the chemical nature of the adsorbates during the occurrence of the wave propagation, as depicted in Figure 14f. The resulting mass spectrometry analysis is presented in Figure 15. The absence of CO₂ species in the mass spectra suggests that CO oxidation does not occur. As discussed above, the CO is chemisorbed on gold and Au-carbonyls are formed in the presence of an external field: AuCO⁺, AuCO²⁺, Au(CO)²⁺ and Au(CO)_{2²⁺}. The propagation wave is thus mainly composed of gold-carbonyls undergoing field evaporation in a wave. The mechanism for this wave propagation is as follows: CO is chemisorbed at the edge site of a terrace because of the local field enhancement weakening the Au-Au bond. The formed Au-carbonyl, denoted Au-CO, is now free to migrate in an inhomogeneous field. Due to the Au-CO dipole, the carbonyl species will be pushed towards the edge of the second layer terrace. The process of detachment/migration is then repeated, and the region where Au-CO desorption occurs, migrates as a wave, from the apex where the field is higher down to the shank of the tip where the electric field intensity drops. This decreasing electric field slows down the propagation, as observed experimentally and theoretically [209]. Regarding the oxygen species, since their quantities adsorbed on the surface are insignificant or even completely absent, the O⁺ and O₂⁺ species observed come from field ionisation in the gas phase, and oxygen species only have a spectator role and act as the imaging gas.

As a conclusion, oxygen does not lead to field-induced oxidation of CO, but the propagation corresponds to the Au-carbonyl wave made visible with oxygen as imaging gas. Simulations [209] were in good agreement with the experiments and showed the absence of oxygen dissociation, the absence of CO oxidation, and the wave propagation of carbonyls to a distance close to the radius of the sample. The study also notes that CO oxidation would be possible with a negative electric field (thus in FEM mode) for pressure in the range of 10⁴ Pa, not accessible in a field emission microscope.



Figure 15. Chemical probing of the carbonyls-wave by PFDMS: Mass spectrum acquired on an Au tip during the wave propagation (see Figure 14) and highlighting the presence of an Au-carbonyls wave and oxygen as imaging gas The region of analysis corresponds to \approx 300 atomic Au sites. Conditions of acquisition: T = 300 K, $P_{CO} = 2 \times 10^{-5}$ Pa, $P_{O_2} = 10^{-2}$ Pa, steady field $F_R = 1.2$ V/Å, pulsed field $F_P = 2.5$ V/Å, frequency of the field pulses f = 10 Hz. Reprinted from [201], Copyright 2006, AIP Publishing.

Water Gas Shift Reaction (WGSR)

The water gas-shift reaction is of importance for the both the conversion of CO gas and the production of hydrogen:

$$CO(g) + H_2O(g) \rightarrow CO_2(g) + H_2(g)$$
(1)

Previous works suggested synergistic effects between Au nanoparticles and their support: the dissociative adsorption of H₂O on Au is followed by spill over of OH species on the support [210,211]. The presence of OH species was also considered important in the promotion of the WGSR [212,213]. The CO/H₂O/Au system was also studied by FIM [204] to try to shed some light on this system. For this, water was adsorbed at the surface of a gold catalyst (as presented in Figure 13a) leading to the formation of bright spots, and then CO was introduced in the chamber at 300 K. When CO is present in the reaction chamber, bright waves ignite simultaneously on different regions of the tips and propagate. Water in the form of H_2O^+ is supposed to be the imaging species. If the CO supply is stopped during the experiment, the wave stops and the brightness fades. If CO is removed after the experiment, new bright spots appear, associated with water adsorption: these facts suggest a reaction between CO and water. The wave front never starts from the same position and there is no preference for certain crystallographic orientations. Accordingly, the reaction is not "demanding", i.e., is not restricted to specific crystalline orientations. Analyses by PFDMS highlight the presence of surface hydroxyls, signs of water dissociation. When CO and H₂O are present in the chamber, the intensity of the hydroxyl peak decreases, which is a strong argument in favour of a reaction between CO and OH species:

$$CO(ads) + OH(ads) \rightarrow CO_2(g) + H(ads) + *$$
(2)

where * stands for an empty site.

The absence of CO_2 in the mass spectra is commonly explained by its formation as a neutral species, which escapes ionisation by field pulses. It has to be noted that a field of 1.1 V/Å is necessary to activate the water dissociation on Au but not sufficient to ionise CO_2 .

3.2.2. Processes on Au-Covered Tips: Determining H₂O Adsorption/Desorption/Diffusion Energies by FEM

The previous section was dedicated to the investigation of reactive processes on Au tips by field ion microscopy. Here, we focus on the study of simple processes by field emission microscopy and the determination of parameters such as the activation energy of desorption and diffusion. Thanks to the FEM technique, these parameters can be obtained for a single grain of catalyst, rather than from a single crystal or from the collection of several particles. For this, Au-covered W tips were used. Au is deposited by evaporation and the Au coverage is estimated via measurements of the work function changes during the evaporation.

In a first set of experiments [214], water adsorption and desorption was studied on Au-covered W tips. Water is condensed on the sample at liquid nitrogen temperature, in field-free conditions to avoid any field-effects on the adsorption energy. To calculate the desorption energy, thermal desorption is applied to the system and corresponds to TPD measurements on a single nanoparticle. In the absence of field and from 78 K, the sample is heated to a chosen temperature, then cooled down. At constant voltage, the presence of water induces a decrease in the emission current. Therefore, the authors monitored the voltage required to maintain a constant emission current (Fowler-Nordheim method). The voltage is measured at 78 K. The first observation from these experiments is the presence of two adsorption states since desorption happens at two different temperature ranges: 150–160 K and 220–240 K. All the water is completely desorbed at 240 K, signs that there is no water dissociation in these conditions (in this case, $\Theta \ge 2$, where Θ corresponds to the surface coverage). For the first adsorption state, thermal desorption isotherms can be plotted, as presented in Figure 16a. An Arrhenius plot can be extracted from this graph and is represented in Figure 16b. The analysis indicates an activation energy for water desorption of $E_{des} = 51.5 \pm 1.5 \text{ kJ/mol}$. The behaviour of water on gold-covered W tip can be compared to a pure W tip, or to a gold-covered tip with a monolayer or a sub-monolayer coverage. In the last case, water dissociation occurs because of the W influence.



Figure 16. Extracting kinetic parameters from FEM analysis: (**a**) Thermal desorption isotherms of water from the Au-covered W tip in the temperature range 145–155 K to probe the desorption energy of the first water-adsorbed configuration; (**b**) Arrhenius plot obtained from (**a**). Reprinted from [137]. Copyright 1997, Elsevier.

Similar experiments were performed for H₂O diffusion by plotting the isotherms in the temperature range 113–123 K and the corresponding Arrhenius plot, leading to an activation energy for diffusion $E_{diff} = 27 \pm 2$ kJ/mol [214]. A more detailed analysis was performed by the same group [215] to study the influence of the Au-coverage and the crystallographic orientation of diffusion on the activation energy. The procedure is rather similar to the adsorption of water at low temperature, followed by sample heating to the temperature range of 118–127 K to let the water diffuse, all this in the absence of an external field. The sample is then imaged at a constant potential and the field emission pattern is processed (by background subtraction and stacking of images so as to increase the contrast and to highlight the low-brightness processes). Figure 17 represents a clean W tip ($\Theta_{Au} = 2.5$) (Figure 17b), and a clean Au-covered W tip ($\Theta_{Au} = 4$) (Figure 17c). The remainder of the Figure 17d–i corresponds to snapshots of the water diffusion, presenting a sharp diffusion-boundary. The diffusing H₂O is represented by the brighter areas of the field emission pattern, and the white lines in Figure 17g–i correspond to the crystallographic direction

where the diffusion was measured. By heating the sample at the targeted temperature during specific time-lapses, it was shown that the square diffusion distance is linear to the diffusion time. Dependence on the Au coverage and crystallographic orientation with temperature shows an activation energy for surface diffusion from 28 kJ/mol to 38 kJ/mol. The differences in energy suggest different roughnesses of the sample.



Figure 17. Imaging water diffusion by FEM: (**a**) Field emission pattern of a clean W tip. The magnification corresponds to ~10⁶ on a detector screen of 3 inches diameter; (**b**) Field emission pattern of a clean Au/W tip with a coverage of $\Theta_{Au} = 2.5$; (**c**) Field emission pattern of a clean Au/W tip with a coverage of $\Theta_{Au} = 4$; Diffusion of H₂O (**d**) FE pattern after water condensation; (**e**) Processed FE pattern from (**d**) where the background has been removed and stacking of 2 identical images. The main crystallographic facets are represented by their Miller indexes: the bright parts correspond to regions where water is adsorbed; (**f**) FE pattern after 300 s of warming at *T* = 120 K; (**g**) Processed FE pattern from (**f**) with background subtraction and stacking; (**h**) Processed FE pattern after 650 s of warming and; (**i**) after 1130 s of warming. Reprinted from [215]. Copyright 1999, Elsevier.

3.2.3. Processes on Au-Based Alloys

Oxygen Interaction with Au-Ag Alloys

Similarly to the previous system, the adsorption of oxygen was studied on Au-Ag alloys [139]. In this study, Au and Ag atomic layers (five to 10 atomic layers are sufficient since they have similar properties to the bulk [216]) were deposited by evaporation and further annealed to ensure a complete mixing of the alloy. Due to the inertness of Au, a decrease of the Ag/Au ratio within the alloy is accompanied by a drastic decrease of its ability to adsorb oxygen, subsequently affecting its catalytic properties. FEM was used to monitor the oxygen adsorption as a function of the alloy composition, and further data processing was applied to calculate the heat of adsorption. The conclusions of this work are: (1) an increase in Au content decreases the sticking probability of oxygen, (2) there is no oxygen detected on pure Au at 300 K, and (3) there is a decrease of the heat of adsorption with added gold. It is interesting to note that 40 years after this study, Au-Ag systems are

still studied to understand the behaviour of Ag within an Au backbone and its influence on the catalytic behaviour [217,218], and that relatively simple studies by FEM enable the nanoscale effect of Au-Ag alloying on the energies of adsorption to be measured.

Towards deNOx Reactions on Au-Based Catalysts

Gold-based catalysts are frequently studied for deNOx reactions [219,220]. First, the adsorption of NO on Au nanoparticles at 300 K was studied [221] and induced, in addition to the expected NO⁺, the presence of dimer (NO)₂⁺, and of nitrous oxide N₂O⁺ species in the mass spectra. Previous works by kinetic analysis [222,223] suggested the formation of nitrous oxide via the combination of two NO(ads) molecules rather than decomposition of NO(ads) and recombination. The absence of N⁺ in the mass spectra suggests that indeed, this mechanism is plausible.

$$(NO)_2(ads) \rightleftharpoons N_2O(ads) + O(ads)$$
(3)

The next level of complexity consists in investigating the adsorption of NO on an Au-based alloy, e.g., Au-Pd. Changes in structure and morphology of the catalyst may be present, but the sample can also undergo changes in the surface composition, i.e., surface segregation of one of the components. The presence of surface segregation in binary alloys plays an important role in catalysis because of the synergistic effects between the two metals that directly depends on their concentration ratio. Furthermore, the control of surface segregation is a crucial step in the design of catalysts with fine-tuned properties. Segregation behaviour in catalytic systems can be suitably studied by atom probe tomography, an extension of the FEM/FIM techniques [88].

The study of the adsorption of NO on Au-62 at% Pd catalysts was performed by FIM, PFDMS and CAP (Catalytic Atom Probe [151]). Even though the presence of two different metals may induce an imperfect FIM image (as compared to pure metals), it is possible to image alloys and to discern different crystallographic facets with very good accuracy, as can be seen in Figure 18a. The adsorption of NO was studied by catalytic atom probe [138] and the main results obtained are as follows: there is no segregation when the sample is exposed to NO at 300 K, and there is no segregation in absence of NO at 573 K. However, exposing the sample to NO at 573 K induces the presence of Pd segregation at the surface, as can be observed in Figure 18b representing the cumulative plots of total atoms (Au + Pd) as a function of the Au atoms. The slope of such plots corresponds to the composition of the analysed sample, and any change in this slope indicates segregation or depletion behaviour. In the case of NO adsorption, we can clearly observe a Pd-segregation: the initial sample composition of Au-62 at% Pd becomes Au-80 at% Pd after NO exposure. This behaviour is explained by the capacity of Pd to dissociate nitrogen monoxide, and the subsequent formation of a strong Pd-O bond. The segregation is thus thermodynamically driven. To prove the dissociation of NO, PFDMS measurements were performed and the mass spectra indicates the presence of N⁺, O⁺, NO⁺ and NO₂⁺ species, proof of the NO dissociation and recombination on this alloy. This is in contrast to NO adsorption on pure Au where the formation of N2O was due to the dimerisation of NO rather than its dissociation [221].

Finally, FIM was used to image a catalytic reaction during the ongoing process [138]: a NOprecovered Au-62 at% Pd sample (NO exposure: t = 30 s, $P_{NO} = 3 \times 10^{-3}$ Pa, T = 450 K) was imaged in the presence of hydrogen. While H₂ pressure is gradually increased, a 'titration-wave' can be observed and is reproduced in Figure 18c for different times of reaction, t = 0 s corresponding to the introduction of H₂ in the reaction chamber. The changes in brightness pattern and the presence of this titration-wave prove that hydrogen species reacted with the adsorbate layer. The imaging field is such that H₂ ionisation is impossible, but sufficient to ionise H₂O which has the role of imaging gas. This observation combined with PFDMS studies suggests that NO dissociates on Pd, subsequent oxygen is more easily adsorbed on Pd than on Au, and O(ads) is reacted off with the H(ads) species. The combined approach of FIM, PFDMS and CAP experiments allows a better fundamental understanding of the system, down to the nanometer-scale to be obtained.



Figure 18. Study of Au-Pd alloy by field emission techniques: (**a**) FIM image of Au-Pd alloy (Au-62 at% Pd). The main discernable crystallographic orientations are represented by their Miller indexes. Conditions of acquisition: T = 45 K, $P_{Ne} = 10^{-3}$ Pa, F = 3.5 V/Å; (**b**) Cumulative plots of ions collected during atom probe experiments before any exposure and corresponding to a composition of Au-80 at% Pd, sign of the Pd surface segregation. Conditions of exposure: T = 573 K, $P_{NO} = 6 \times 10^3$ Pa, t = 15 min. Reprinted from [138]. Copyright 2009, Elsevier; Titration of a NO-precovered Au-62 at% Pd sample by hydrogen, proving the formation of water, and therefore, the dissociation of NO on this catalyst. Panel (**c**) Courtesy of M. Moors.

3.2.4. Perspectives

In the past, Au-based systems were studied by FIM for various applications such as the study of long-range migration of self-interstitial atoms [224], phase transformation [225], precipitation in neutron-irradiated alloys [226], conductivity [227], or in the application of nano-tips to electron microscopy [228]. Even though these studies were not focused on catalysis, they proved the feasibility of imaging various Au-based alloys by FIM; alloys which are now used for catalysis applications: Au-Mo [228] for the reverse water-gas shift reaction [229,230] or as N₂ dissociation catalyst for the Haber-Bosch process [231], Au-Pt [224] for CO oxidation [49,232] and selective toluene oxidation [52], Au-Fe [226] for CO oxidation [233] and N₂ dissociation catalyst for the Haber-Bosch process [231], and Au-Cu [225,234] for the water gas-shift reaction [235,236] and CO oxidation [50].

Finally, three more applications are relevant for catalysis studies: the interaction of cyclohexane on Au tips [237], which finds its interest in the abatement of Volatile Organic Compounds (VOC); the behaviour of potassium-gold films [227], which could be used to study the influence of promoter in a catalytic reaction; and eventually, the imaging of Au nanoparticles by FIM after deposition on a field emitter tip [238,239], opening a way to study the catalytic process on a single Au nanoparticle rather than on a tip-model.

3.3. Study of Processes on Supported and Self-Supported Catalysts by E-TEM

The selected examples presented in this section will describe the shrinking process of nanoparticles, the morphological changes of supported nanoparticles during exposure to different atmospheres and the support-dependence of these changes, and finally, the use of self-supported Aubased catalysts, i.e., nanoporous gold, for oxidation reactions. As a preliminary comment, we mention that imaging by TEM is based on the interaction of the sample with a high-energy electronbeam. It is known that the sample/e-beam interaction may induce undesired effects on the samples. In the specific case of gold, one example of this effect can be observed in Figure 19, which represents a maturation process of two Au nanoparticles [240]. The images are obtained by HR-TEM, and each image was acquired after a few minutes of illumination under the electron beam. We can observe that the small nanoparticle disappears layer by layer (Figure 19b–d), down to a single atomic layer.

This process is similar to the Ostwald ripening and the transport of atoms from the shrinking particle to the bigger particle may occur by surface diffusion along the CeO₂ surface. Further studies highlight the structural changes of nanoparticles under e-beam radiation on different support (TiO₂, Fe₂O₃...) [241,242]. Another study showed an increase in coordination of Au nanoparticles under e-beam irradiation [243]. Since the size, structure and the under-coordinated sites are known to affect the catalytic activity, one has to be careful in the interpretation of results obtained under e-beam.



Figure 19. Series of high-resolution TEM revealing the shrinking process of an Au nanoparticle supported on CeO₂ solely induced by e-beam illumination: (**a**) one large and one smaller Au nanoparticles can be observed; the smaller nanoparticle shrinks layer by layer (**b**) 3 remaining layers; (**c**) 2 remaining layers; (**d**) 1 remaining layer. Reprinted from [240]. Copyright 2005, Springer.

3.3.1. Processes on Supported Au Nanoparticles

Coalescence of Au Particles

One of the key parameters for the catalytic activity of gold is the size of the nanoparticles. It is then important to understand the factors affecting the sintering of nanoparticles, under reactive conditions, so as to control them. The mechanism of model Au nanoparticles sintering was studied on graphene support [244,245] under vacuum and hydrogen flow at different temperatures. As depicted in Figure 20, Au nanoparticles present a size ranging from 1 to 5 nm. The graphene support is stable at temperatures up to 673 K. At room temperature, the nanoparticles are rather stable, meaning that they make irregular movements around a fixed site, both in vacuum and in the presence of hydrogen. Once the temperature is increased, nanoparticles become dynamic and coalescence of nanoparticles can be observed. Figure 20 represents such coalescence through surface mobility under hydrogen gas at a temperature of 373 K.



Figure 20. Coalescence of Au nanoparticles supported on graphene via migration of nanoparticles. The snapshots correspond to time t = 0 s, 56 s and 94 s. Conditions of acquisition: T = 373 K, $P_{H_2} = 200$ Pa, 300 kV. Reprinted with permission from [245]. Copyright 2014, American Chemical Society.

We can observe that Au particles diffuse as crystalline entities on the graphene substrate until they make contact. At this point, the particles are linked together (Figure 20 middle) via the formation of a crystalline bridge, allowing the atoms to diffuse from one nanoparticle to eventually reach another. If the temperature is increased up to 673 K (and still under a pressure of $H_2 = 200$ Pa), a different mechanism takes place, similar to the Ostwald ripening where bigger particles grow to the detriment of smaller ones via mass transfer. This proves that coarsening can follow multiple routes, particle migration and Ostwald ripening, which depends on the reactive conditions [244–246]. From a structural point of view, the choice of the support is thus crucial to avoid sintering and ensure the activity of gold catalyst.

Importance of the Support for the Stability

Catalysis by gold is most likely affected by the electronic state of the Au nanoparticles, which strongly depends on their interface with the support [247–250]. Metal-oxide supports such as CeO₂, TiO₂ and Fe₂O₃ are commonly used, their key function being the stabilisation of gold nanoparticles through the surface oxygen vacancies, which in turn depend on the size and shape of the support. On ceria support [251], it has been proved that the Au nanoparticles move reversibly, back and forth between equivalent sites by approximately 0.09 nm at room temperature and in presence of a reactive environment. This movement happens to a lesser extent in vacuum, but more frequently under O₂ gas: this is explained by the compensation of oxygen vacancies by the O₂ gas, decreasing the anchorage sites of the Au nanoparticles on the support. It was concluded that gold-based catalysts are relatively loosely bound to the support and may then diffuse on the surface with low activation energy. The authors also note that in the case of experiments by E-TEM, the low energy can be supplied by e-beam radiation. Using CeO₂ nanorods rich in surface oxygen vacancies, Au nanoparticles are strongly anchored at the surface, active for CO oxidation and WGS reaction and stable under realistic reaction conditions. Indeed, no sintering was observed up to 573 K [252].

Another striking example of nanoparticle/support interaction is the structural reorganisation on the Au/TiO₂ system under beam irradiation [253]. The study was performed under vacuum, inert atmosphere (N₂-100 Pa), oxygen atmosphere (O₂-100 Pa) and reactive atmosphere (1 vol% CO in air-100 Pa).

The facetting of the Au nanoparticles, which will be described in the next section, can be summarised as follows: facetted morphology is observed under vacuum, inert and reactive environments; and round-shaped morphology is observed in only O₂ atmosphere. By switching from one gas to another, a systematic change of the morphology can be observed in the majority of the nanoparticles [253–256]. Regarding the active phase/support interaction, it was observed that by increasing the electron dose *D*, the nanoparticles were increasingly decorated with material originating from the support. This can produce decorated, pillared or encapsulated particles, as can be observed in Figure 21A. The structural evolution diagram is presented in Figure 21b, where the regions of observation of different structures are plotted as a function of the electron dose *D* and electron emission current ϕ . For low *D* and ϕ , corresponding to the green region, no structural reorganisation can be observed. However, by increasing *D* and ϕ , the structural changes become an irreversible process. Finally, by extrapolating for *D* = 0 and ϕ = 0, the intrinsic structure of the catalyst may be deduced and is located in the green region.

This study proves that the e-beam radiation must be accounted during E-TEM experiments, and that a careful control of the environment is required to avoid any artefacts. Indeed, earlier works explained the difference in catalytic reactivity of Au/TiO₂ as compared to Au/CeO₂ due to the presence of decorated/pillared/encapsulated nanoparticles. These structures are however present only in the presence of e-beam irradiation, and not in the applied formulation of catalysts.





Figure 21. Au/TiO₂ nanoparticles changes under e-beam exposition: (**A**) Structural reorganisation of Au/TiO₂ nanoparticles under e-beam irradiation in vacuum, O₂ environment (100 Pa) and CO environment (1 vol% CO in air 100 Pa). *D* corresponds to the electron dose and ϕ to the electron current density; (**B**) Structure evolution diagram of Au/TiO₂ nanoparticles under e-beam showing the regions for the formation of decorated, pillared and encapsulated nanoparticles as a function of *D* and ϕ . The intrinsic structure can be extrapolated for *D* and ϕ = 0. Reproduced from [253]. Copyright 2012, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

Morphology of the Active Nanoparticles

Previous sections focused on the behaviour of gold nanoparticles during coalescence, as well as the imaging of the behaviour of the substrate materials with nanoparticles. E-TEM can be applied to study the morphological changes of the gold catalyst during the reaction and to correlate the morphology to the catalytic activity. The behaviour of Au/CeO₂ and Au/TiO₂ under vacuum, inert, oxidising and reducing conditions has been extensively studied.

On CeO₂ support, the Au nanoparticles present different morphologies as a function of the composition of the surrounding gas phase [257]. Under vacuum, inert gas atmosphere (N₂) and during CO oxidation (in a mixture 1 vol% CO/air at 1 mbar, corresponding to the reactive environment), the nanoparticles appear in the form of a stable polyhedron enclosed by the major {111} and {100} facets. This morphology is represented in Figure 22A(a,b). The facetted morphology becomes "rounded" under O₂ atmosphere, as shown on further TEM images in Figure 22 where the CO/O₂ pressure ratio is decreased, inducing a more oxidising environment. This change in morphology is however reversible. By applying a systematic change in the partial pressures of CO and O₂, a morphology diagram can be plotted and is represented in Figure 22B. The statistically mixed region corresponds to a region where less than 70% of the particles present a well-defined shape. With this, it can be concluded that CO molecules stabilise the {111} and {100} facets. CO molecules are indeed adsorbed on the on-top sites of an Au{100}-hex reconstructed surface under catalytic conditions (1 vol% CO in air at 100 Pa and room temperature), and this structure is energetically favourable [258].

The origin of rounded morphology observed in O₂ gas comes from fluctuating multi-facetted surfaces. Indeed, a careful examination of each frame of the E-TEM experiments showed that the minority {110} facets emerged frequently in addition to the major {111} and {100}. It is concluded that atomic oxygen from the dissociation of O₂ causes the rounded shape of the nanoparticles. The authors also mentioned this because of the residual gases in the E-TEM, oxygen-related species may be formed and induce the rounded morphology. However, the studied nanoparticles present the same morphology as TEM experiments in vacuum after exposure to an environment at different gas

pressures. The studies at low gas pressure are thus useful to understand the morphology of active nanoparticles under realistic catalytic conditions. Indeed, the morphology changes are in good correlation with the catalytic activity [257,259]. It has to be noted that even in the presence of strong interfacial bonding of Au with CeO₂ nanorods [252], the reshaping of the nanoparticles and the nanoparticle/support interface under reducing environment (42 vol% CO/6 vol% O₂/N₂) can be observed and prove the dynamic nature of the Au/support interface.



Figure 22. Study of the morphology of active Au nanoparticles: (**A**) Morphological change of an Au nanoparticle when the pressure ratio CO/O₂ is decreased; (**B**) Morphology diagram representing the dependence of the morphology of Au nanoparticles on the partial pressure of CO and O₂, with mostly round-shaped particles in oxidising environment and facetted particles in reducing environment. Reproduced from [257]. Copyright 2011, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

Importance of the Support for the Activity

As discussed above, CeO₂ and TiO₂ supports allow morphological changes of the Au nanoparticles to be observed. A comparative study was performed between two different Au/CeO₂ catalysts and less-active catalysts: Au/SiO₂ and Au/TiC [260]. The nature of the support is different: reducible oxide for CeO₂, non-reducible oxide for SiO₂ and carbide for TiC. By exposing the samples to 1 vol% CO/air at 100 Pa and to O₂ at 100 Pa, both Au/CeO₂ catalysts switched from facetted to rounded morphology. In the case of Au/SiO₂ and Au/TiC, the nanoparticles mostly remained facetted in both reducing and oxidising environment, as depicted Figure 23.

In active catalysts, the rounded morphology is due to O₂ gas in the reaction cell, and it is more likely that oxygen-species are generated at the periphery interface between the nanoparticles and the support. One of the hypotheses concerning the activation of oxygen on gold catalysts is the presence of surface oxygen vacancies on the support, contributing to a strong bond between the support and the Au nanoparticles. Therefore, electron transfer becomes possible and leads to negatively charged Au nanoparticles, enabling O-species adsorption on Au. When the Au nanoparticle and the support are structurally coherent, the nanoparticle becomes a sphere. The resulting periphery interface is important, and thus the resulting catalytic activity becomes high. Whether it is because of a lack of surface oxygen vacancies or an amorphous structure, the Au/support interaction might not be the key feature to explain catalytic activity. Therefore, the latter may be related to the "morphology-changeable" character of the nanoparticles, which directly depends on the nature of the support and its synergy with the nanoparticle [260]. As can be seen in Figure 23, this morphology-changeable character correlates with the catalytic reactivity.



Figure 23. Correlation of the catalytic activity of supported Au catalysts with the "morphologychangeable" character of the nanoparticles, which is directly influenced by the nature of the support. Reproduced from [260]. Copyright 2017, Elsevier.

3.3.2. Processes on Self-Supported Au

The current research in gold catalysis consists of the study of catalytic reactions on nanoporous catalysts [261–265]. The nanoporous Au, npAu, consists of a 3D structure of interconnected pores and ligaments with a ligament size in the 20–100 nm range. The 3D network of ligaments and pores ensures the mechanical stability of the catalyst, as well as the mass transport of reactants and products through the catalyst [266]. These structures are prepared by free corrosion of a bulk Au-Ag alloy in an electrolyte [267], leading to the selective removal of the least noble metal, Ag in this case. As a function of the initial concentration, different ligament sizes can be obtained [268,269], allowing the structure of the final catalyst to be fine-tuned. Residual Ag is present in the final structure with a concentration of few percent. This Ag is known to influence the catalytic activity [270–273] and thus its presence has to be considered.

It is known that the catalytic activity on gold nanoparticles depends on the size of the Au [29,203] and that the support strongly influences the reactivity, as discussed in the previous section [260]. However, as is described below, significant activity can be observed in nanoporous systems, even though the characteristic length-scale is one order of magnitude higher (20–100 nm) and the catalyst is self-supported. In the same way that sintering decreases the activity of gold nanoparticles, coarsening decreases the activity of nanoporous gold. It has been proved that coarsening occurs during CO oxidation, and that planar defects can effectively prevent structure coarsening [274], and deformation mechanisms have to be accounted for [275]. Annealing and exposure to O₂ gas can lead to coarsening, material loss and material migration [276].

Nanoporous Au as Catalyst for CO Oxidation

Nanoporous Au catalysts with a ligament size of ≈ 30 nm and a residual Ag content of 1.2 at% were studied under environmental conditions to understand their reactivity towards CO oxidation [277]. The reactivity of npAu is similar to the reactivity of Au nanoparticles. High-resolution imaging allows observing the presence of a high density of atomic steps and kinks on the curved surfaces of npAu (on both convex and concave surfaces), which is one of the features also present on Au nanoparticles. Under CO oxidation conditions at room temperature, the surface is catalytically active, and significant surface reconstruction is observed at a few Pa, corresponding to {111} facetting dynamics along the {011} surface. The atoms in top-most surface are involved in CO oxidation. The facetting occurs while the e-beam is blanked, and is not observed under N₂, O₂ and CO atmosphere under the same imaging conditions: reconstructions are thus neither due to the e-beam irradiation, nor to the gas pressure. Residual silver may thus play a role. It is supposed that Ag segregates to step and kink positions on the surface so as to reduce the free energy for steps formation (as compared to pure Au). To prove this, an npAu sample of similar pore size and similar internal surface but comprising 20 at% of residual Ag has been used for CO oxidation. With this catalyst, {111} facetting dynamics are significantly suppressed during CO oxidation, leading to the conclusion that Ag

significantly stabilises the surface steps and kinks. Therefore, the presence of a higher Ag concentration accelerates the catalytic oxidation by suppressing the surface reconstruction dynamics and providing more Ag active sites for O₂ dissociation.

Nanoporous Au as Catalyst for Selective Alcohol Oxidation

The catalytic activity of nanoporous gold catalysts with a ligament size of \approx 50 nm (Figure 24a) and a residual Ag content of \approx 3 at% were studied for the selective oxidation of methanol [217]. Both the ligament size and the Ag content are expected to influence the catalytic activity as compared to the previous example. In this work, the authors focused on influence of ozone activation on the catalyst to improve the selectivity towards alcohol oxidation reactions. One of the key steps towards this goal is the dissociation of oxygen species at the surface of the catalyst, and Ag provides active sites for oxygen dissociation [261–265].

Experiments show that the ozone treatment creates a thin silver-rich layer $(1.1 \pm 0.1 \text{ nm})$ (Figure 24b), as well as crystalline islands (Figure 24c) and amorphous oxides (Figure 24d). EELS analysis (Figure 24e,f) also prove the silver enrichment at the surface of the ligament. From a structural point of view, the oxide layer can be reduced by CO, leading to the formation of small irregular nanoparticles on highly stepped regions of the catalyst. Further reduction in methanol results in the formation of 2–2.5 nm crystalline nanoparticles which may re-alloy into the bulk. From a catalysis point of view, the oxide layer is highly active for both CO and non-selective methanol oxidation. However, the reduction by CO leads to the formation of a distinct oxygen species stabilised by Au-Ag alloy sites. This distinct oxygen is inactive for CO oxidation, but active for selective methanol oxidation into formaldehyde. Ambient-pressure XPS and the temporal analysis of products (TAP) method corroborate these observations [217].



Figure 24. Nanoporous Au ad self-supported catalyst for selective alcohol oxidation: E-TEM analysis of nanoporous gold after ozone treatment showing (**a**) the porous and ligament structure of the sample; (**b**) the presence of an oxide layer due to ozone treatment; (**c**) the presence of crystalline islands; (**d**) TEM of amorphous oxide at the surface corresponding to silver-oxide species; (**e**) spots analysed by EELS; (**f**) EELS analysis showing the region of high Ag and O content. Reprinted by permission from [217]. Copyright 2016, Macmillan Publishers Ltd.

It can be concluded that the surface undergoes structural and compositional changes during activation by ozone and that the Ag segregation plays an important role in the formation of oxygen species for selective oxidation of methanol. The last two examples presented show that similar catalysts may behave completely differently as a function of the residual concentration of Ag, but also as a function of the pretreatment/activation steps applied to that catalyst. The presence of Ag, or any other metal, could be used to engineer catalysts for specific and selective reactions.

3.3.3. Perspectives

Currently, Au-based systems are studied by TEM and/or E-TEM for applications different to catalysis, but it may be of interest for heterogeneous catalysis applications to further study the role of the support and the synergy between the support and the catalyst. Such studies comprise the use of iron oxide and gold catalysts for the formation of carbon nanotubes [278], the analysis of unsupported Au particles deposited on Si [279], the Au assisted growth of MgO crystals [280], and Au assisted growth of Ge nanowires [281]. Different Au-based alloys were studied by TEM or E-TEM; these studies could be extended to their behaviour under catalytic conditions. These alloys are Au-Ni [282], Au-Cu alloys [283–285], Au-Pd, Au-Ag and Au-Pt nanorods [286].

As mentioned, e-beam irradiation may induce negative effects on the catalyst. However, in some cases, these effects are useful to produce Au-based alloys with specific structure. It has been proved that it was possible to grow Au layers on Pt particles by low-dose TEM [287,288], or to produce Au@Ag core-shell structures by exposing Au and Ag nanoparticles to a high dose of e-beam [242]. Finally, the e-beam irradiation has also been used in in situ liquid cell TEM to synthesise Au-Pd core-shell nanostructures [289]. The use of liquid cell for TEM [290] could thus be used to study the catalyst formation, as well as to follow liquid-phase reactions with Au as catalyst.

4. Conclusions and Perspectives

This review presents the main results that can be obtained from the analysis of gold and goldbased catalysts by environmental microscopies. By presenting case studies from three main families of techniques, it has been possible to cover the wide range of the catalytic complexity studied in fundamental research so as to get closer to the imaging of applied formulations of catalysts under industrial conditions. The PEEM/LEEM techniques address flat samples (single crystals, foils or composite surfaces); a model single nanoparticle of catalyst can be imaged by FIM/FEM techniques, whereas E-TEM can be used to study unsupported, supported and self-supported catalysts. Of course, these techniques present some limitations. PEEM experiments have a relatively low resolution, but allow global processes to be studied. FIM/FEM studies are, most of the time, performed with an imaging field, but allow Au catalysis in the complete absence of support to be studied; E-TEM imaging suffers from beam damages, but the technique can be used for a wide variety of catalysts.

PEEM was used to study how the presence of Au, even at low coverage, on a Pt surface affects the dynamics by modification of the kinetic parameters, whereas LEEM was used to probe the mass transport of the catalytic material, due to the reaction. In the latter case, the initial structure is important: starting from an O-covered Rh surface or a clean Rh surface before the deposition of gold, the behaviour under reaction conditions differs from stationary waves to segregated phases. By further increasing the complexity and depositing Au and Pd, the phase separation can be reversibly obtained by changing the control parameters. FIM experiments allowed wave propagation during the interaction of CO with O_2 on a single nanoparticle of Au to be observed. A detailed analysis of this system proved that oxygen eventually had the role of imaging gas and that the CO oxidation did not occur on an unsupported Au nanoparticle. However, a mixture of CO and H₂O seems to react via the water gas shift reaction. FEM was used to determine activation energies for water desorption and diffusion, to study the adsorption of oxygen on Au-Ag alloy, and also to investigate the NO hydrogenation on Au-Pd nanoparticles. Finally, E-TEM is of great interest to move closer to real conditions, as was shown by the study of Au nanoparticles sintering and the influence of the support. The morphological changes during gas exposure is again correlated with the nature of the support and the nature of gas studied; and finally the study of nanoporous structures made of Au-Ag where the catalytic activity depends on the size of the ligaments, the Ag content and its distribution.

Environmental microscopies are thus crucial to get a better understanding of catalytic systems, but complimentary techniques have to be applied to obtain chemical information during imaging.

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Spectroscopy, spectrometry and/or diffraction methods are thus necessary to get a full picture of the catalytic process. To cite but a few, surface propagation during CO oxidation on Pt induces a surface transformation that cannot be observed by PEEM, but is probed by LEED; the brightness in the LEEM patterns is associated with different surface compositions via their comparison with micro-XPS measurements; mass spectrometry was extensively used to prove the presence or absence of dissociative processes observed by FIM, as well as the segregation of Pd in Au-Pd alloys; and EELS is used to highlight the Ag segregation on nanoporous gold catalysts.

The comparison of three different techniques allows us to draw a few conclusions regarding gold catalysis:

- The addition of a minor amount of surface species may severely affect the global reactivity of the catalyst, as it was demonstrated on Au-covered Pt in PEEM and Au-Ag alloys in nanoporous gold by E-TEM;
- Alloying induces synergistic effects, as shown on Au + Pd-covered Rh by LEEM and Au-Pd catalysts for NO hydrogenation by FEM, a reaction that was not possible on pure Au;
- The size-effect in gold catalysis arises from the step and kinks density, as shown on nanoporous structures by E-TEM where the reactivity of the npAu depends on the size of the ligaments. These studies also prove the relevance of using a tip-sample of 20–50 nm by FIM to study fundamental processes;
- The presence of silver can be used to stabilise the structure, but also to guide the selectivity of the reaction, as it was studied by FEM and E-TEM;
- CO oxidation has been studied by PEEM, FIM and E-TEM; the water gas shift reaction by FIM and E-TEM, but current research focuses on more complex reactions over Au-based catalysts.

The emergence of environmental microscopies greatly improved the understanding of catalytic processes. The constant technological developments allow better imaging resolutions under reaction conditions to be obtained and more importantly, the developments towards correlative microscopy and spectroscopy/spectrometry under reaction conditions promise a golden future for the study of catalytic processes closer to conditions used in industrial applications.

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