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Reducible oxides as ultrathin epitaxial films

Paola Luches, Sergio D'Addato

Abstract

This chapter reviews and discusses recent work on two-dimensional films of reducible oxides supported on metal substrates. In general, peculiar chemical and structural phases, different from the bulk ones, can be stabilized depending on the oxygen chemical potential, on kinetic processes and on the specific substrate used. A peculiarity of reducible oxides is that the observed phases can often be reversibly transformed one into the other by applying reducing and oxidizing treatments.

1. Introduction

An oxide is defined reducible if it can be easily and reversibly reduced depending on the ambient conditions. Reducibility is linked to the existence of two or more oxidation states with comparable stability for the cations. Prototypical reducible oxides are represented by cerium and titanium oxides, although also other transition metal and rare earth oxides can be considered reducible; among these for example other rare earth oxides, like PrO_x , SmO_x , TbO_x , and other $3d$ metal oxides, like VO_x , MnO_x , FeO_x , CoO_x , but also HfO_x , TaO_x , NbO_x , WO_x and many more.

Indeed, reducibility is very relevant for catalysis, since materials based on reducible oxides can act as oxygen buffers, which can store and release oxygen and/or charge, promoting redox reactions with a unique regeneration ability [1]. Furthermore, reducibility is an important property also in view of the application of oxides in other fields, like for example energy conversion and storage,¹ biomedicine [2] and memories [3]. Reducibility is not only linked to the material capacity of easily forming oxygen vacancies but also to the reversibility of the process and to oxygen transport properties within the material.

The influence of **reduced dimensionality** on the **reducibility** of an oxide is certainly relevant in view of the optimization of the properties of the material through the understanding of the modifications which arise with confinement. Pronounced changes of the **electronic structure** of oxides at reduced dimensionality have been widely investigated, and interesting properties have been observed on reducible oxides in particular [4,5]. In cerium oxide particles of nanometric size the oxygen vacancy formation energy has been found to be greatly reduced and to reach a minimum at a specific size [4], thereby causing a surprisingly enhanced reactivity of catalysts made of metal nanoparticles supported on nanocrystalline cerium oxide [6], possibly also facilitated by easy oxygen transfer to the supported metal nanoparticles [7].

If only one of the dimensions of the considered material is confined to the nanoscale, i.e. if the system under study is a two-dimensional film, several interesting modifications are introduced and intriguing phenomena can take place. The stabilization of metastable structural phases by epitaxy or by the mere spatial confinement, the presence of lattice strain or rumpling, the interaction with the substrate, in terms of charge transfer but possibly also of interfacial atom exchange, the different

stability of structural defects, are all expected to determine relevant changes in the electronic structure, which give origin to systems with different properties compared to the corresponding bulk phases.

The studies of two-dimensional reducible oxides are typically carried out on **model systems**, i.e. on ultrathin films supported on single crystalline substrates, prepared and analyzed in high or ultrahigh vacuum. The investigations aim at identifying interesting aspects on simple systems, which can be analyzed by **surface science techniques** and understood with the help of theoretical descriptions. A proper **modeling** of the systems under investigation often represents a crucial step for a complete atomic level understanding of the material properties and potential performances. Only through the fundamental insight of the new properties which arise on simple systems, the much more complex real systems, based on the same materials, can be understood and optimized in their functionality. For example, catalysts made of metallic nanoparticles supported on reducible oxides typically involve some degree of encapsulation of the metal within the oxide and the formation of very thin oxide layers on the metal nanoparticle surface, through the so called strong metal-support interaction (SMSI) [8]. Ultrathin oxide layers exposing different facets, with different structures and morphologies can be prepared in the form of model systems to identify the most active species towards the considered reaction. The results of these studies can be used to design the catalysts with an optimized activity and selectivity by finding suitable synthetic routes to maximize the density of required active sites for the chosen specific functionality.

In this chapter recent studies on the properties of ultrathin films of reducible oxides are reviewed, focusing on the aspects which influence the reducibility of the materials at the two dimensional limit. New properties of cerium oxide two-dimensional films are discussed in section 2. Section 3 reports studies of titanium oxide two-dimensional films, which are interesting for comparison and for complementing the concepts outlined on cerium oxide based systems. Selected studies of ultrathin films of different reducible transition metal and rare earth oxides are finally reported in section 4.

2. Cerium oxide two-dimensional films

The most stable **cerium oxide** phase is CeO_2 , also known as *ceria*, which has a fluorite structure with cerium ions in the 4+ oxidation state. The Ce_2O_3 phase, with an orthorhombic structure (A-type phase), is relatively less stable at ambient conditions. Several additional metastable phases with different stoichiometry can be stabilized in specific conditions [9], among them the Ce_2O_3 bixbyite (c-type) phase deserves a particular relevance, since it can be obtained from the most stable fluorite-type CeO_2 phase by removing 25% of the oxygen lattice atoms in an ordered way and allowing for a slight structural rearrangement [10].

The (111) surface of the CeO_2 phase is the thermodynamically most stable one, followed by the (100) and (110) surfaces, the latter two being polar [11]. The repeating unit along the (111) direction is an O-Ce-O triple layer, which in the following will be referred to as a monolayer (ML). Cerium oxide ultrathin films exposing the (111) surface can be obtained by epitaxial growth on metallic single crystal substrates with six-fold surface symmetry such as (111) surfaces of cubic structures [12-18], or hcp (0001) surfaces [14,19,20]. The lattice mismatch between cerium oxide

and most of the metal substrates is quite large and it amounts to approximately 30-40%. Nevertheless, films with a (111) surface orientation and a very good epitaxial quality have been obtained [15,17,21,22]. The (100) and (110) surface orientation can be stabilized through epitaxy using substrates with a very small lattice mismatch with CeO₂. These are typically non-metallic and include Si [22,23], YSZ [24] and SrTiO₃ [25,26]. However, most of the studies performed using non-metallic substrates focus on thick films with several tens of nm thickness, possibly because of the difficulties in applying surface science techniques, necessary to obtain information at the ultrathin limit, on substrates with low conductivity.

The first and main part of this section will be dedicated to studies of **cerium oxide (111)** ultrathin films, and the final part to the investigations concerning ultrathin films exposing less stable surfaces. Important aspects connected to reducibility at the two-dimensional limit will be reviewed and discussed, with focus on: i) structure, morphology and defectivity, ii) charge transfer and intermixing at the interface, iii) modifications in structure and morphology induced by reduction, iv) stabilization of metastable structures induced by dimensionality and/or epitaxial constraints.

Pioneering works studying ultrathin cerium oxide films on the Pt(111) surface date back to the mid-nineties and they were motivated by the idea that a deep understanding of ceria-based materials obtained from studies of single crystalline surfaces and films of different thickness, structure and morphology could help improving the activity of cerium oxide supports in three way catalytic converters [12,13,27,28]. Cerium oxide films were grown either by deposition of metallic cerium followed by post-oxidation [12,27] or by oxidation of Ce-Pt surface alloys [13,28]. Already these early works pointed out that ultrathin films in the ML and sub-ML range contain a non negligible Ce³⁺ concentration, while thicker films can be fully oxidized [12,13]. As will be clearer in the following, this aspect is rather crucial in trying to identify a charge transfer from the metal substrate and/or a dimensionality-induced decrease of oxygen vacancy formation energy induced by the lower average O coordination of Ce ions at the early stages of the growth. The **structure** of the films was found to be disordered unless thermal treatments at temperatures higher than 700 K, which stabilize the fluorite (111) phase, are performed [12,13]. **Thermally-induced instabilities** were observed above 1000 K [12,13]. Interestingly, films which fully cover the substrate showed a significant activity towards CO oxidation, higher than the one of the bare Pt(111) surface [12]. An important role of the metal-oxide interfacial sites and of non-(111) oriented surfaces was also hypothesized, based on the high activity of films which do not fully cover the substrate [12].

The results of the early studies provided a strong motivation for subsequent works, using more controlled growth procedures, combined with a variety of techniques for surface characterization down to the atomic level. Cerium oxide ultrathin films with the (111) fluorite structure were grown on Pt(111) [17,21,28,29], Rh(111) [15,16], Ru(0001) [14,20], Ni(111) [14], Cu(111) [30,31], Au(111) [32], Pd(111) [33] and Re(0001) [34] substrates. On one side these studies confirmed the enhanced reactivity of bidimensional cerium oxide films compared to the clean substrates and the importance of metal-oxide interface sites [31,35], on the other they clarified important fundamental aspects linked to the reducibility of low-dimensional supported cerium oxide systems [15,29,30].

In general, the studies of ultrathin supported cerium oxide films and islands were motivated by the need to identify the active sites in real catalysts made of metallic nanoparticles supported on cerium oxide. Several works specifically addressed the **reactivity** of metal-supported cerium oxide systems

and towards selected reactions [31,35,36,37]. A catalytically active role of the interface between cerium oxide and Rh(111) in CO oxidation was suggested for example by Eck et al., who identified preferential CO adsorption sites near the phase boundaries of the oxide islands [36]. Suchorski et al. showed that the catalytic activity towards CO oxidation of cerium oxide nanoislands on Pt(111) is remarkably increased compared to the bare Pt(111) surface [35]. By comparing systems with different density of step edges they suggested the observed increase to be due to the different electronic properties at the islands perimeter [35]. A similar role for the oxide metal interface, combined with the low dimensionality of cerium oxide, was identified for the CO oxidation reaction and for the water gas shift reaction on the cerium oxide/Cu(111) inverse catalyst model system [31,37]. Studies of this kind stimulated more accurate work on cerium oxide bidimensional structures and systematic investigation of the influence of the preparation conditions and of the substrate used on the structure, morphology, defectivity and reducibility of the investigated systems.

On most substrates post-growth thermal treatments in O₂ have been found to improve the surface morphology, stoichiometry and structure of the cerium oxide terraces obtained [17,20]. The ultrathin films have relatively large coincidence cells with the underlying metal surface giving origin to **moiré patterns** in scanning tunneling microscopy (STM) images [20,38] or in low energy electron diffraction (LEED) patterns [15]. The coincidence cells with the lowest strain typically involve 5×5 ceria surface cells and 7×7 substrate surface cells (hereafter termed 5:7 coincidence), however at the ultrathin limit smaller coincidence cells, implying a larger **strain**, can be stabilized. This phenomenon has been observed for example on the Pt(111) substrate, where ultrathin films have a compressed surface structure and an interatomic distance comparable with the 3:4 coincidence cell, while thicker films have the relaxed bulk structure [17,21,39]. At the ultrathin limit a contraction of the in plane lattice parameter has been observed also using Rh(111) as a substrate [15], and even on Cu(111) substrates [38]. The evidence for a lattice contraction also on Cu(111), on which cerium oxide could adopt a 3:2 coincidence with the substrate with negligible strain, demonstrates that the tendency for lattice contraction is probably an intrinsic property linked to reduced dimensionality [38].

The deposition of cerium oxide amounts in the ML range typically results in the formation of large **flat islands** of ML or multi-layer height [15,17,20]. Procedures to obtain **continuous films** of monolayer thickness with almost complete coverage have been identified on the Cu(111) substrate, using low temperature (110 K) growth in O₂ followed by post-annealing in O₂ at 770 K [18]. Ultrathin cerium oxide films grown in strongly oxidizing conditions contain a non-negligible concentration of Ce³⁺ ions on the Pd(111) [33], Pt(111) [17] and on the Rh(111) substrate [15]. In the latter case a preferential localization of the Ce³⁺ sites at the interface was deduced by comparison of x-ray photoemission spectroscopy (XPS) and resonant valence band photoemission spectra. Valence band resonant photoemission, having a smaller probing depth than XPS, does not show any significant Ce³⁺ concentration on the surface. The non-negligible Ce³⁺ concentration detected by XPS is therefore ascribed to interface sites [15]. This represents a strong evidence of possible **charge transfer** from the metallic substrate towards the cerium oxide films. Effects such as charge transfer can indeed significantly alter the properties of two-dimensional cerium oxide films, and a definite assignment of the origin of reduced interfacial states requires a theoretical description of the system. On the Cu(111) substrate density functional theory (DFT) calculations showed that a very relevant charge transfer from the metal to the oxide takes place, yielding to the reduction of a

full layer of interfacial Ce ions [40]. DFT calculations on a 1 ML cerium oxide film supported on Pt(111), instead, evidenced electrostatic interactions between the topmost Pt layer and the interfacial oxygen atoms in the oxide, which cause a significant corrugation of both the film surface and of the interfacial Pt layer [41]. The charge transfer between Pt and Ce ions is significant only in the interfacial sites where the distance between the two atoms is short [41].

Not only the size, but also the shape and defects of the terraces obtained on the surface of ultrathin cerium oxide films seem to depend on the preparation conditions. On Ru(0001), using high temperature ($T > 800$ K) growth and post-growth annealing ($T \sim 1000$ K) in oxygen atmosphere ($P > 10^{-7}$ mbar), ultrathin films with point defects, linear defects ascribed to grain boundaries and round terrace step edges have been obtained [20]. The presence of **grain boundaries**, possibly induced by three-dimensional growth mode at the early stages of film formation [20], is not suppressed by room temperature growth followed by annealing of ceria films using a Pt(111) substrate [17]. On this substrate ultrathin films present **terraces** with straight **edges**, oriented along the substrate main symmetry directions [17]. Thicker films instead show more rounded terrace edges [17]. This is possibly caused by the stabilization of less stable step edges by the proximity of the metallic substrate in ultrathin films. A detailed characterization of the film step edges for various preparation conditions showed that depending on the heating temperature different step types can be obtained. The different steps are rationalized in terms of stability of the exposed facets and analyzed in terms of electronic properties [42]. One-dimensional electronic states have been found to develop on specific step types, and at domain boundaries as shown in figure 1, which reports STM images acquired at different sample biases.

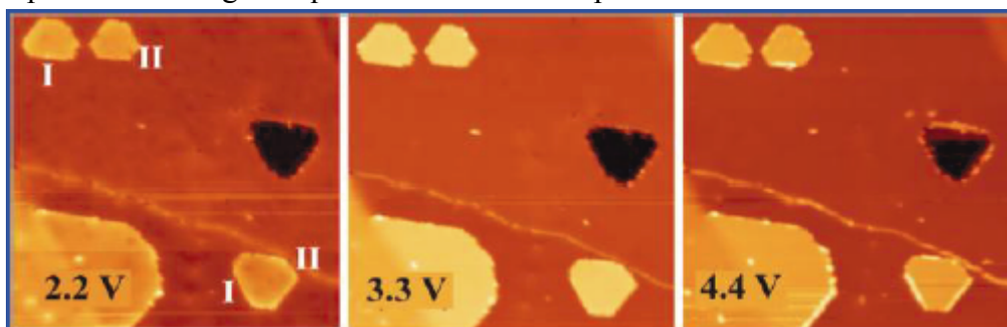


Fig.1: 65×65 nm² STM topographic images of a 6 ML cerium oxide film grown on Ru(0001) with islands and holes exposing different step edge orientation. Steps indicated as I and domain boundaries show an increased apparent height at 4.4 V sample bias, while steps indicated as II do not change apparent contrast with bias. Reprinted with permission from Nilus et al. [42] Copyright 2012 American Chemical Society.

The step edges exposed by cerium oxide two-dimensional films show similarities to those exposed by bulk ceria surfaces, although in this case depressions rather protrusions are most frequently observed on the surface [43]. A non-negligible density of **point defects** has been often observed on ultrathin film terraces [21]. The defects are very similar to the ones formed on bulk surfaces, the most common being surface oxygen vacancies, also imaged as trimers and as linear arrays, and subsurface oxygen vacancies [21]. Defects can also be intentionally induced by electron bombardment of the film surface [44]. A work by Jerratsch et al. identified some degree of delocalization of the charge left after O vacancy formation [44]. The delocalization of charge after O release has been highly debated also on bulk surfaces [45,46], and at reduced dimensionality some non negligible degree of delocalization has been clearly identified [5,47].

Cerium oxide films can be **reduced** by different procedures, including thermal annealing in vacuum [15,17,29,35], growth of metallic Ce in low oxygen background pressure [14,16,48], ion bombardment [49] or exposure to reducing gases [49-51]. **Vacuum thermal treatments** often induce also a structural and morphological modification, pointed out to be strongly dependent on the initial film thickness [15,29]. In particular, the temperature at which reduction starts is lower for thinner films, which also show an apparently higher final degree of reduction [15,29]. Indeed it is difficult to determine the absolute amount of Ce^{3+} ions formed in the different samples by the different treatments, given the unknown shape of the depth profile of O vacancy concentration, and the different depth sensitivity of the techniques used to measure the $\text{Ce}^{3+}/\text{Ce}^{4+}$ ratio. However, the idea of having a higher density of Ce^{3+} sites in the surface layer after reduction is generally accepted [29]. A dependence of the onset of cerium oxide surface reduction on the growth temperature of the film, and hence on the density of reduced coordination sites, has been pointed for films grown on the Cu(111) substrate [38]. A very interesting effect, shown to take place on the surface of a thin ceria film on Rh(111) after reduction, is the formation of an ordered array of **surface defects**, clearly visible in the STM images (figure 2), and ascribed to triple oxygen vacancies [52]. The smaller energy for the formation of vacancies at specific sites of the 5:7 coincidence lattice is ascribed to the local surface stress [52].

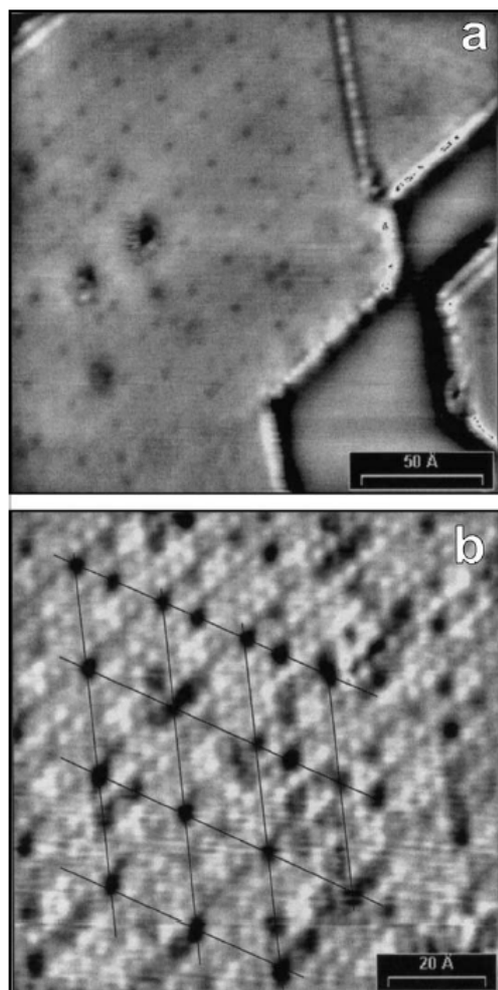


Fig.2: STM images of a 0.5 ML CeO_{2-x} film grown on the Rh(111) surface after annealing at ~ 900 K. (a) $200 \times 200 \text{ \AA}^2$; 0.93 V; 0.86 nA. (b) $100 \times 100 \text{ \AA}^2$; 0.80 V; 1.05 nA. The grid of black lines evidences the superlattice of defects,

ascribed to triple oxygen vacancies. Reprinted with permission from Casterllarin-Cudia et al. [52] Copyright 2004 Elsevier.

The surface of reduced ceria films on Pt(111) instead shows a corrugated **morphology** with bias-dependent features tentatively ascribed to electronic modifications induced by reduction [29]. Interestingly, ultrathin reduced cerium oxide phases show peculiar **surface reconstructions** (figure 3), whose origin is still under investigation and appears to be linked to the presence of the underlying substrate [29]. The effect of the heating time and heating rate on the final degree of reduction of the films has been also pointed out to be very important [29]. Some extra-periodicities have been observed also on the Cu(111) substrate after deposition of variable amounts of metallic cerium on a CeO₂ buffer layer, followed by annealing in UHV [53]. Some of them are commonly observed also on the surface of thick films after reduction [54] and correspond to metastable bulk phases such as the Ce₇O₁₂ phase and the c-type bixbyite phase.

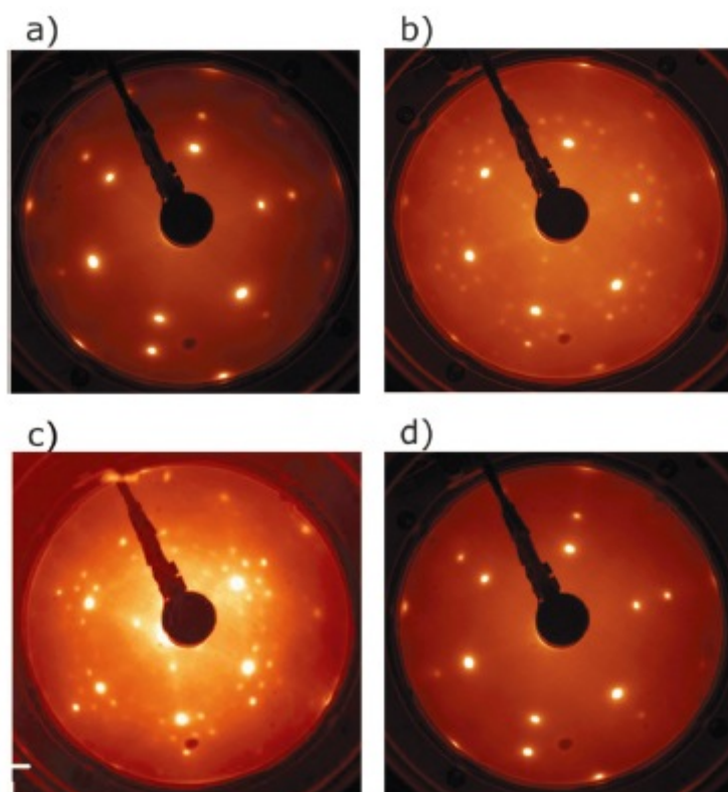


Fig.3 LEED patterns ($E_p = 80$ eV) of a 2 ML cerium oxide film (a) as prepared, (b) after intermediate reduction by heating in UHV at 770 K for 30 min ($c_{Ce^{3+}} \sim 40\%$, as measured by XPS), showing the (3×3) and the $9/4(\sqrt{3} \times \sqrt{3})R30^\circ$ phase (c) after strong reduction by heating in UHV at 1040 K for 15 min ($c_{Ce^{3+}} \sim 60-80\%$) showing the $9/4(\sqrt{3} \times \sqrt{3})R30^\circ$ phase (d) after re-oxidation by heating in O₂ at 1040 K. Luches et al. [29] - Reproduced by permission of the PCCP Owner Societies.

The full **reversibility** of the reduction process was demonstrated for ceria films of different thickness on Pt(111) [29]. We note here that this aspect is not trivial, and it is linked to the reducibility of the films. Cerium oxide films on a Si(111) substrate reduced by vacuum thermal treatments under conditions similar to those used in ref. 29, showed a non reversible reduction of Ce ions from the 4+ state to the 3+ state [23]. This process was ascribed to the formation of interface cerium silicate phases which are not reducible [23]. Interfacial atom exchange during growth was also observed using Ni(111) and Cu(111) substrates, and they were shown to have a non-negligible

influence on the stoichiometry of the films at the ultrathin limit [14,31]. On the Au(111) substrate the formation of Au/Ce alloys hinders the good ordering of extended two-dimensional films [32].

Substoichiometric films were also obtained by evaporation of Ce on Ru(0001) and on Ni(111) in low oxygen pressure ($P_{O_2} \sim 10^{-8}$ mbar) and they were found to have the fluorite structure with a high concentration of oxygen vacancies in at least the topmost oxygen atom layers compatible with a bixbyite structure rather than to a hexagonal Ce_2O_3 structure [14].

As reported above, several works have shown some degree of reducibility of cerium oxide ultrathin films, which is typically higher than the bulk, but they also evidenced that fully reduced states are very difficult to be achieved [15,29]. This may be linked to the fact that it is easy to form oxygen vacancies in the fluorite structure up to a certain density, or to the fast (partial) reversibility of the reduction process when high Ce^{3+} concentrations are obtained. The **cubic c-type bixbyite phase**, with 25% oxygen vacancies is in fact unstable in the bulk form and a full cerium oxide reduction may require a transition to the most stable hexagonal A-type phase. A metastable epitaxial cubic c-type Ce_2O_3 phase has been stabilized on the Cu(111) surface by deposition of metallic Ce on a CeO_2 buffer layer followed by 900 K annealing [55]. The film shows a well ordered surface with a (4x4) reconstruction with respect to the $CeO_2(111)$ surface (figure 4). The reconstruction is ascribed to ordered arrays of quadruple oxygen vacancy clusters and corresponds to the bulk termination of c- Ce_2O_3 [55]. Although the film thickness is as high as a few nm, the influence of the substrate is considered to be determinant for the stabilization of the observed phase, ascribed to the effect of the tetragonal strain induced by the Cu substrate [55]. A phase with a similar c-type bixbyite structure in the form of an ultrathin film was also stabilized on a Cl-passivated Si(111) surface [22].

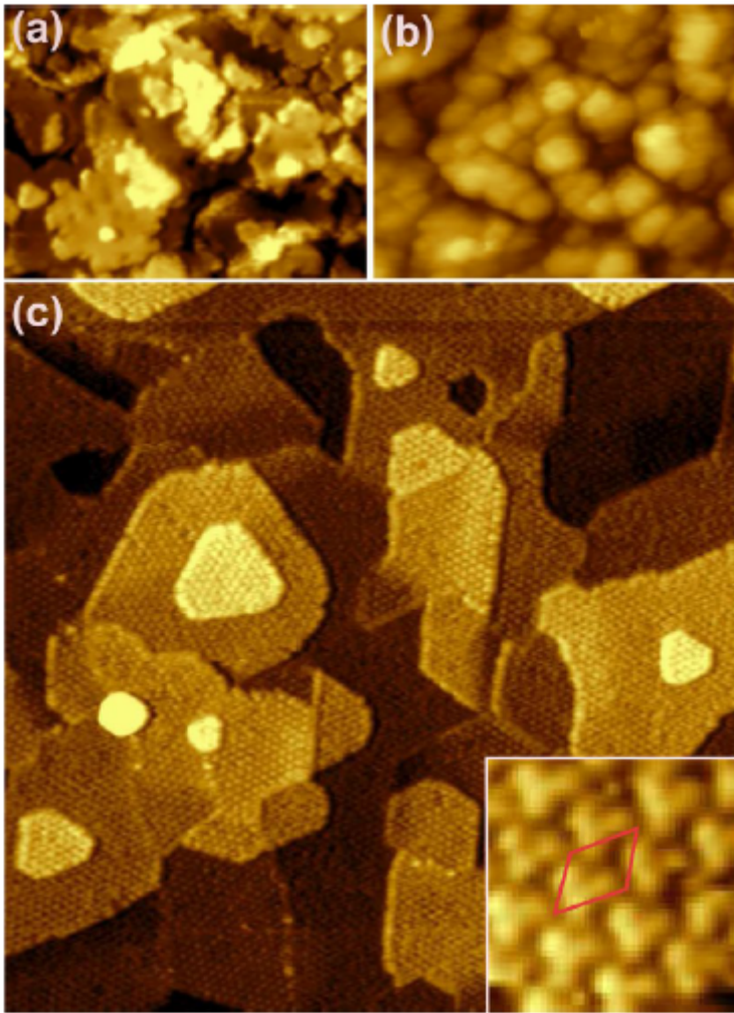


Fig.4: STM images showing the different stages of formation of $c\text{-Ce}_2\text{O}_3$ films on Cu(111). (a) CeO_2 buffer, (b) CeO_2 buffer with subsequent metallic Ce deposition, (c) ordered $c\text{-Ce}_2\text{O}_3$ layer obtained by annealing (b) in vacuum at 900 K. Inset: high-resolution image and surface unit cell (red rhombus) of the $c\text{-Ce}_2\text{O}_3$ layer. Images a–c are to scale. Image width (a,b) 60 nm, (c) 120 nm, (inset) $6 \times 6 \text{ nm}^2$. Reprinted with permission from Stetsovych et al. [55] Copyright 2013, American Chemical Society.

The exposed works show that ultrathin cerium oxide films offer new opportunities in view of obtaining modifications of the material properties, and in particular for the stabilization of **metastable structural phases**. This last issue has been investigated also in a rather recent **theoretical investigation** aimed at identifying the most stable Ce_2O_3 structures at the ultrathin limit [56]. The work, using simulated mechanical annealing searches and DFT calculations, shows that different ultrathin film structures are stable at different values of the in plane lattice parameter and it identifies in particular a new structure, which does not correspond to any known bulk crystalline **polymorph**, which is more stable than the A-type and than the c-type Ce_2O_3 structures at specific, relatively large, in-plane lattice parameters (figure 5) [56].

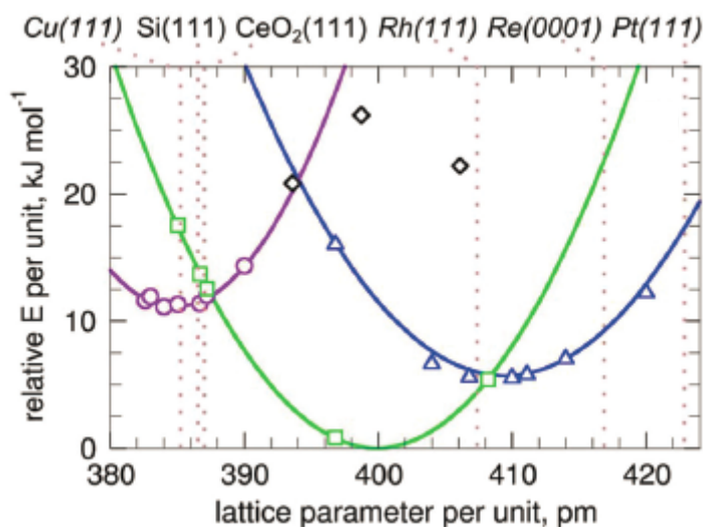


Fig.5: Results of DFT calculations for 4 ML Ce_2O_3 films with different structure: A-type (circles), bixbyite (squares) NF1 (triangles), NF2–4 (diamonds). Energies (relative to that of the optimized bixbyite film) and lattice parameters are given per Ce_2O_3 unit. Vertical dotted lines indicate lattice parameters of possible supports for film growth (multiplied by 3/2 for transition metals – highlighted by italics). Kozlov et al. [56] - Published by The Royal Society of Chemistry.

As mentioned at the beginning of this section, a few studies have also been focused on the stabilization of ultrathin cerium oxide films exposing surfaces different from the (111) on metal substrates. For example, the stabilization of cerium oxide nanoislands exposing (100) facets, with thickness down to 1 ML, has been shown to be possible using a Cu(111) surface and highly oxidizing growth conditions [57]. The stabilization mechanism invoked is the formation of a copper oxide with a rectangular unit cell at the interface, which imposes an epitaxial constraint and shares an O layer with cerium oxide to compensate polarity (figure 6) [57]. On the same substrate the presence of an interfacial $\text{CeO}_2(100)$ layer supporting three dimensional CeO_2 nanoislands with (100) surface orientation has also been observed to coexist with the (111) CeO_2 orientation [58].

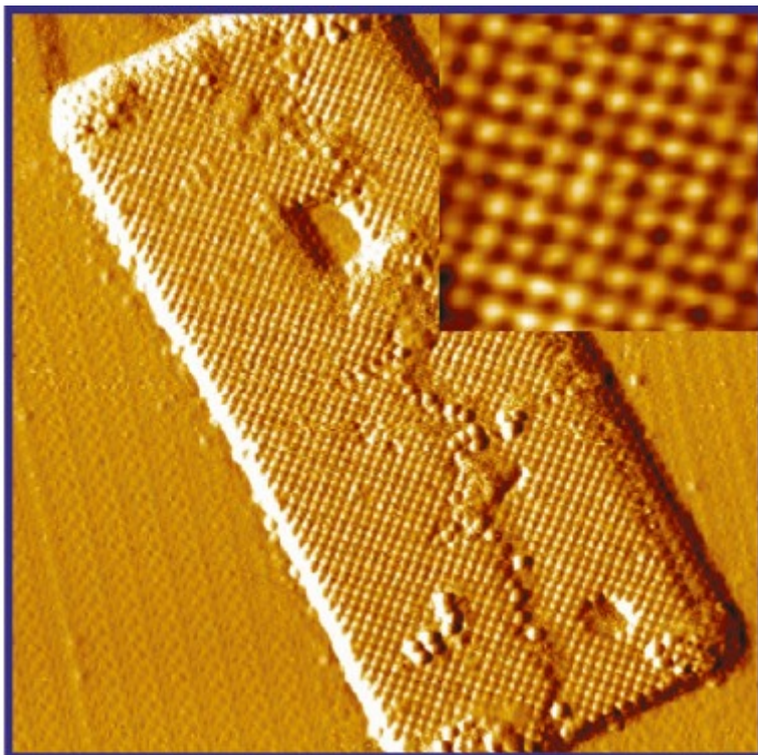


Fig. 6: High resolution STM images of CeO₂(100) phases prepared by Ce deposition in 5×10^{-7} mbar of O₂ at 650 K on a copper oxide surface, formed by oxidizing Cu(111) with NO₂ at 600 K. The large-scale images with size 40×40 nm² is shown in the derivative mode. Reprinted with permission from Yang et al. [57] Copyright 2011 American Chemical Society.

A recent study pointed out the formation of **ceria nanocrystallites** exposing **(100) facets** after thermal treatments in vacuum of continuous (111) cerium oxide films grown on Ru(0001) [59]. The nanostructures, of nanometric height and several tens of nm lateral size, compensate their polarity through surface reconstructions similar to those hypothesized for the bulk (100) orientation [11]. Open aspects which may contribute to a better understanding of less stable surfaces, also in view of preparing two-dimensional films with less stable orientation, are the understanding of the mechanisms which preferentially stabilize this phase under highly reducing conditions compared to other phases, and the possible reasons for the existence of the (001) orientation only at relatively large heights of a few nm. Although studies of cerium oxide films exposing less stable surfaces at the two-dimensional limit are in general rather scarce, they have a great potential interest in view of the expected smaller surface oxygen vacancy formation energy, due to the lower coordination of surface O atoms. Furthermore, the possible formation of new structural phases driven by the compensation of polarity at reduced dimensionality may also open up new perspectives, in analogy with the case of non-reducible oxides [60].

3. Titanium oxide two-dimensional films

Titanium oxide is another case study for reducible oxides. Unlike cerium ions which are stable only in the 3+ and 4+ oxidation states, titanium ions can have different oxidation states, the most common ones being Ti²⁺, Ti³⁺, Ti⁴⁺, giving rise to various oxides. The most stable oxide is TiO₂,

also known as *titania*, which can present different structural phases, the most important ones being rutile, anatase and brookite.

Ultrathin titanium oxide films grown on metal substrates have been the subject of extensive research over the last twenty years, as they can give rise to novel nanostructures and to a rich variety of phases which have been studied using surface science techniques. Stoichiometric and non-stoichiometric phases with different atomic structures can be obtained through the growth on single crystal metal surfaces, using simple **experimental procedures**: deposition of Ti in a vacuum chamber in controlled pressure of residual oxygen, i.e. reactive deposition, post-oxidation of Ti metal films, reduction in vacuum by heating the films in UHV or in controlled residual hydrogen pressure. A selection of results reported in the literature, with focus on peculiar properties arising at two-dimensions and in particular on reducibility-related issues, are hereby reported and discussed.

Experiments on titanium oxide ultrathin films grown on Cu(100) were carried out by different groups [61-63]. Maeda et al. [61] deposited metallic Ti on the Cu surface previously exposed to O₂ (P= 1 × 10⁻⁶ mbar at T=330 K), and oxidized the resulting film by post-growth O₂ dosage (P= 1 × 10⁻⁷ mbar at T=623 K). A two-dimensional growth of an O²⁻/Ti⁴⁺/O²⁻ trilayer with TiO₂ stoichiometry was deduced from XPS data. LEED patterns showed a **hexagonal geometry** with two domains rotated by 90° with respect to each other, and an in-plane lattice constant of 0.29 nm. The study also indicates that this phase is unstable for coverages beyond a single O²⁻/Ti⁴⁺/O²⁻ trilayer. The model proposed by Maeda et al. [61] for the atomic geometry of the trilayer is analogous to the one deduced for the quasi-hexagonal structure formed by titanium oxide on the oxidized (110) surface of the NiTi alloy [64]. A quasi-hexagonal phase was also observed for titanium oxide films obtained by reactive deposition of Ti (P= 1 × 10⁻⁶ mbar, T=573 K) on Cu(100) previously saturated with chemisorbed oxygen [63,65]. The chemisorption resulted in a (√2 × 2√2) R45° LEED pattern with Cu missing-row reconstruction. At low Ti coverage ($\theta_{\text{Ti}} < 0.5$ ML) the same group observed a LEED pattern with a centred rectangular unit cell indicated as c-(√2 × √2) R45°. STM data showed that the rectangular phase is associated with flat islands of uniform thickness, embedded within the outermost layer of the substrate. The stabilization of this phase is probably favoured by the missing-row reconstruction of the O-Cu(100) surface used as substrate [63,65]. The quasi-hexagonal phase is observed at $\theta_{\text{Ti}} > 0.5$ ML (figure 7), however when the whole substrate surface is covered by the film ($\theta_{\text{Ti}}=2.0$ ML) a regular hexagonal pattern is visible in the LEED, very similar to the one observed by Maeda et al. [61]. An in-deep X-ray photoelectron diffraction (XPD) and LEED intensity analysis of the quasi-hexagonal phase, compared with DFT calculations, confirmed the O-Ti-O trilayer model, allowing to identify also the specific registry of the titanium oxide film with respect to the Cu(100) surface (figure 7) [65]. Interestingly, DFT provided also information about the **electronic properties** of the film, showing that the O 2p and Cu 4sp states overlap and that the film does not show an insulating behaviour because of an upshift of the O 2p bands and a downshift of the Ti 4s states, compared to the case of the ideal unsupported films.

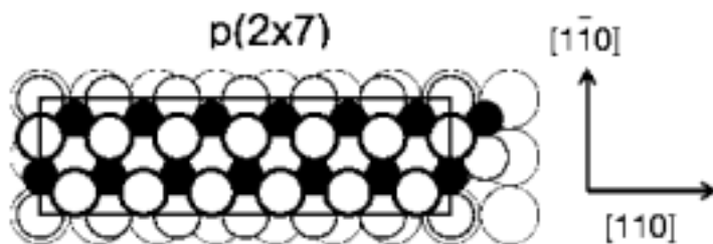
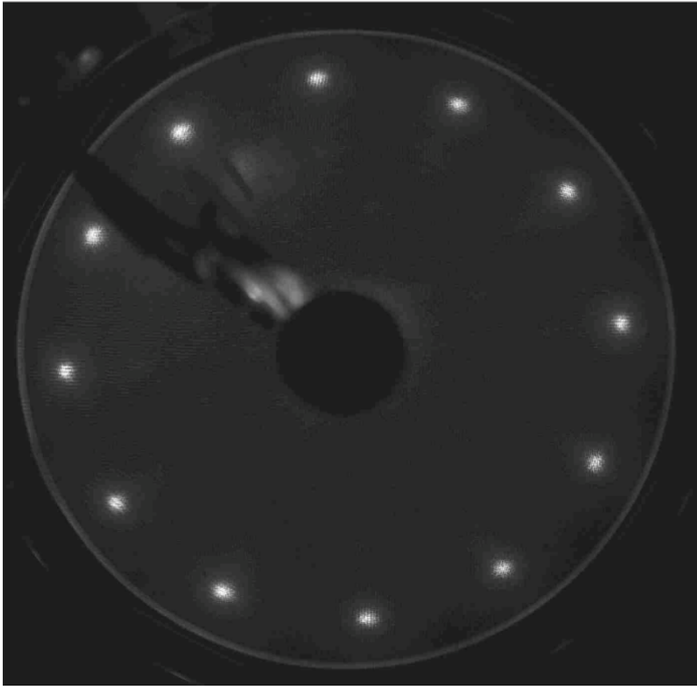


Fig. 7. a) LEED pattern observed a TiO_2 film with the quasi-hexagonal structure on the $\text{Cu}(001)$ surface. b) Atomistic model of the quasi-hexagonal phase of the O-Ti-O trilayer, with a rectangular $p(2 \times 7)$ unit cell, showing the registry of the film with the substrate. Reprinted with permission from Atrei et al. [65] Copyright 2009 American Chemical Society.

A TiO_2 **lepidocrocite-like structure** was instead obtained at submonolayer coverage when the substrate used for the growth was $\text{Ag}(100)$ [66]. Again, LEED, XPD and STM experiments results were compared with the results of DFT calculations. At coverages higher than 1 ML, islands with the rutile (110) surface begin to form [67]. Single domain titania nano-sheets with lepidocrocite structure were also observed when TiO_2 films were grown on the (1×2) $\text{Pt}(110)$ surface [68].

Rutile TiO_2 (110) films were also observed to grow on $\text{W}(100)$ [69,70] and $\text{O}(2 \times 1)$ - $\text{W}(100)$ [71] in two orthogonal domains along the $\text{W}[010]$ and $[001]$ directions. In the first case, the TiO_2 films at coverage values between 5 and 30 ML reverted to the bulk structure, and a systematic splitting of the spots in the LEED pattern was attributed to the formation of a stepped surface, probably caused by a strain relaxation mechanism due to the compression of the long axis of the TiO_2 unit cell. In the case of oxygen-reconstructed $\text{W}(100)$ surface, STM images showed the formation of titanium oxide islands at low coverage following the orientation of the original missing rows induced by the $\text{O}(2 \times 1)$ reconstruction (figure 8). At increasing coverage, the islands extend and become higher, but TiO_2 does not form a continuous film. The presence of point defects on the rutile islands, characteristic of the native rutile $\text{TiO}_2(110)$ - (1×1) surface was observed in high resolution STM

images. TiO_2 was also grown by reactive deposition and annealing on $\text{Mo}(100)$ [72,73]. At coverage values below 10 nm, LEED showed a $(2\sqrt{2} \times \sqrt{2}) R45^\circ$ pattern, while STM images showed ordered atomic rows along the [010] and [001] substrate direction. Annealing procedures allowed **reduction** of the film, with the Ti ions oxidation state changing from Ti^{4+} only, to a mixture of Ti^{4+} , Ti^{3+} and Ti^{2+} valence states, as evidenced by XPS. Similar studies were carried out also on $\text{Mo}(110)$ [74,75], on which films of TiO_2 with (100) orientation or $\text{Ti}_2\text{O}_3(0001)$ were obtained, depending on the preparation method. In particular the $\text{Ti}_2\text{O}_3(0001)$ phase was stabilized by initial deposition of a metallic Ti layer on the $\text{Mo}(110)$ surface, followed by reactive Ti growth in O_2 in the same conditions which lead to the formation of the $\text{TiO}_2(100)$ phase [74]. STM images evidenced smooth surfaces with distinct flat **terraces** and well-defined **step edges** after annealing in O_2 at $T = 900\text{-}1100$ K, while XPS data showed that the oxide films remained partially reduced after the treatment. In contrast, thin films annealed in oxygen at $T=1200$ K were fully oxidized but they exhibited a three-dimensional rough surface morphology [75].

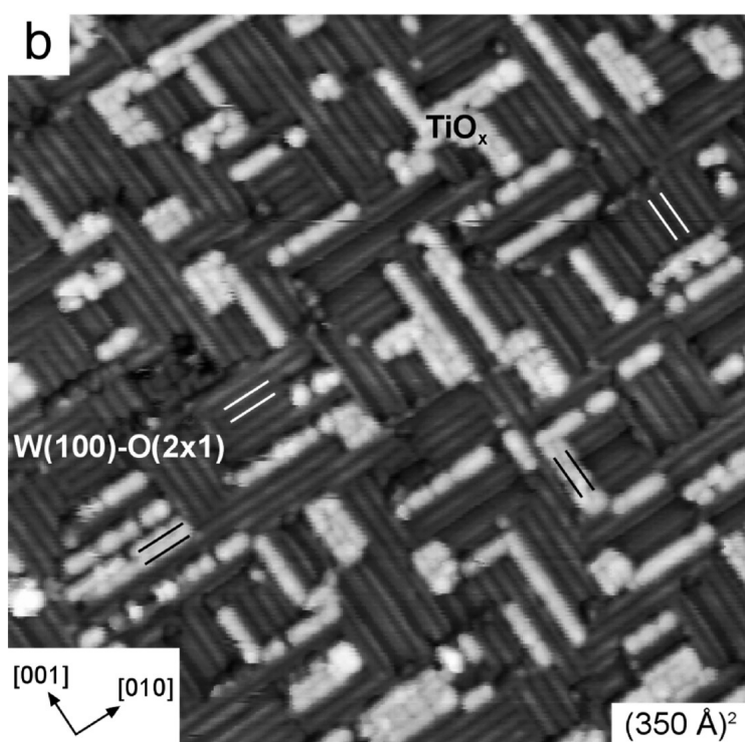


Fig. 8. STM image of a 0.2 ML film of $\text{TiO}_x(110)$ on $\text{W}(100)\text{-O}(2\times 1)$. An area of $\text{TiO}_x(110)$ and an area of the $\text{W}(100)\text{-O}(2\times 1)$ substrate are indicated. White lines are drawn over some of the $\text{W}(100)\text{-O}(2\times 1)$ rows then duplicated, in black, over the TiO_x rows to highlight the similar periodicity. The crystal directions correspond to those of the $\text{W}(100)$ substrate. Reprinted with permission from Pang et al. [71] Copyright 2013 American Chemical Society.

The same group also reported formation of a well-ordered (8×2) TiO_x film on $\text{Mo}(112)$, obtained by depositing Ti on $\text{SiO}_2(\text{ML})/\text{Mo}(112)$ followed by oxidation/annealing and a final anneal at 1400 K to completely remove residual Si [76]. This phase shows peculiar properties, including a strong interaction with Au, which allowed a complete wetting of its surface, at variance with other oxide supports [71,76].

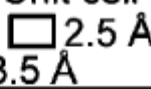
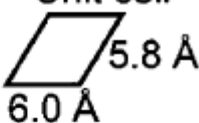
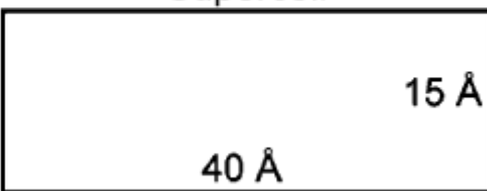
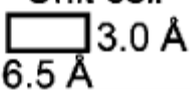
		Substrate: Ni(110) Unit cell 
Ti coverage (MLE) Oxidized to saturation by 10 ⁻⁷ mbar O ₂ Annealed to 900 K	< 1 MLE	TiO ₂ quasi hexagonal overlayer Unit cell 
	~ 1 MLE	TiO ₂ wetting overlayer Unit cell  Supercell 
	> 1 MLE	TiO ₂ (110) overlayer on wetting layer Unit cell 

Fig. 9. Summary of the TiO₂ phases grown on Ni(110). Reprinted with permission from Papageorgiou et al. [78] Copyright 2007 American Chemical Society.

Using a Ni(110) substrate for titanium oxide ultrathin films growth either a quasi-hexagonal phase or (110) rutile rods on a TiO₂ wetting layer could be obtained, depending on the initial titanium coverage [78,79]. In both cases, XPS and x-ray absorption near edge spectroscopy (XANES) results demonstrated that Ti was fully oxidized. Figure 9 gives a summary of the TiO₂ phases reported in ref. 76. The rutile and wetting layer phases showed a similar behaviour with respect to exposure to water at P=10⁻⁸ mbar, which resulted in coadsorption of both molecular water and hydroxyl groups. By dosing water at P=10⁻⁶ mbar hydroxylation of the thin films was observed. The process was found to be reversible by annealing at T=490 K. STM images on rutile rods revealed a (1×2) reconstruction in some areas, ascribed to a reduced phase, very close to the one observed on the surface of bulk rutile TiO₂(110) single crystals [79]. The (1×2) islands disappear completely after annealing at T=773 K in O₂ [79]. Further reduction of the surface could be induced by annealing in UHV at T=1110 K, with observation of {132} and {121} families of crystallographic shear planes. In this respect the behaviour of ultrathin films is analogous to the one of the TiO₂ (110) single crystal surface [80].

The study of titanium oxide films on Pt surfaces is particular relevant, in view of a deeper understanding of the phases formed in oxide-supported metal catalysts as a consequence of SMSI [8]. Experiments of Pt nanoparticles grown on TiO₂(110) surface revealed that the (111) oriented Pt islands were encapsulated within TiO₂ regular films after annealing at T=1100 K with a subsequent change of the catalytic activity [81]. The encapsulation was observed and studied in detail also in

the case of Pd on (1x2) TiO₂(110) [82]. Some of the structures obtained in these systems were also observed on the reverse catalyst model TiO_x/Pt systems.

Matsumoto et al. used a number of experimental techniques for a complete study of titanium oxide grown on Pt(100) [77,83]. In particular, two different routes for **film synthesis** were used: in the first one a Pt₃Ti surface alloy was oxidized with O₃ and annealed at T=1000K. The result was a smooth film composed of one layer of Ti₂O₃ with (3×5) superstructure, with flat terraces without islands, and with a structure similar to the (1×2) strands formed on the reduced TiO₂(110) surface. The second route was reactive deposition of Ti in O₂ (p=6.7 × 10⁻⁷ mbar) followed by annealing above 750 K in vacuum. In this case, a (3×5) structure could be observed for coverages lower than 1 ML, while a (4 × 3√5)R60° structure was obtained after deposition of 2 ML. The proposed model for the observed reconstruction consists of TiO₂ tetragonal nets with some O atoms in the second layer. The (3×5) structure was re-obtained after annealing above 950 K in vacuum. At increasing coverage and after annealing at T=1000 K, TiO₂ clusters were observed, coexisting with Ti₂O₃ (3×5) and clean Pt(100) domains. Finally, after further annealing at T=1300 K the TiO₂ clusters decomposed to form a (2√2 × 2√2)R45° structure, proposed to be Ti₅O₈, and (3 × 5) domains.

Boffa et al. investigated titanium oxide films grown on Pt(111) for coverages ranging from 1 to 5 ML [84]. Two **different structures** were obtained depending on the sample pretreatment. In particular, annealing at temperatures between 770 and 970 K in O₂ gave a (√(3 × √43))R7.6° three fold symmetric structure, with TiO₂ stoichiometry and primitive cell 18.2 Å × 18.2 Å size. Annealing in vacuum (820 -1120 K) gave a second ordered overlayer with primitive cell of 18.2 Å × 13.9 Å size and Ti₄O₇ stoichiometry. Partial dissolution of Ti in Pt was observed after annealing at high temperatures, reducing the thickness of the films to approximately 1 ML [84].

A systematic work on titanium oxide films grown on Pt(111) was carried out during the last years [85-90]. The study first concentrated in ultrathin layers, with an amount of Ti lower than 1.2 MLE, where 1 MLE corresponds to the surface density of Pt atoms per unit area on the Pt(111) surface. By varying the Ti dose and the annealing conditions (temperature and oxygen pressure) six **metastable phases** with different long-range order were originally found [85]. The conditions leading to the different phases are summarized in figure 10.

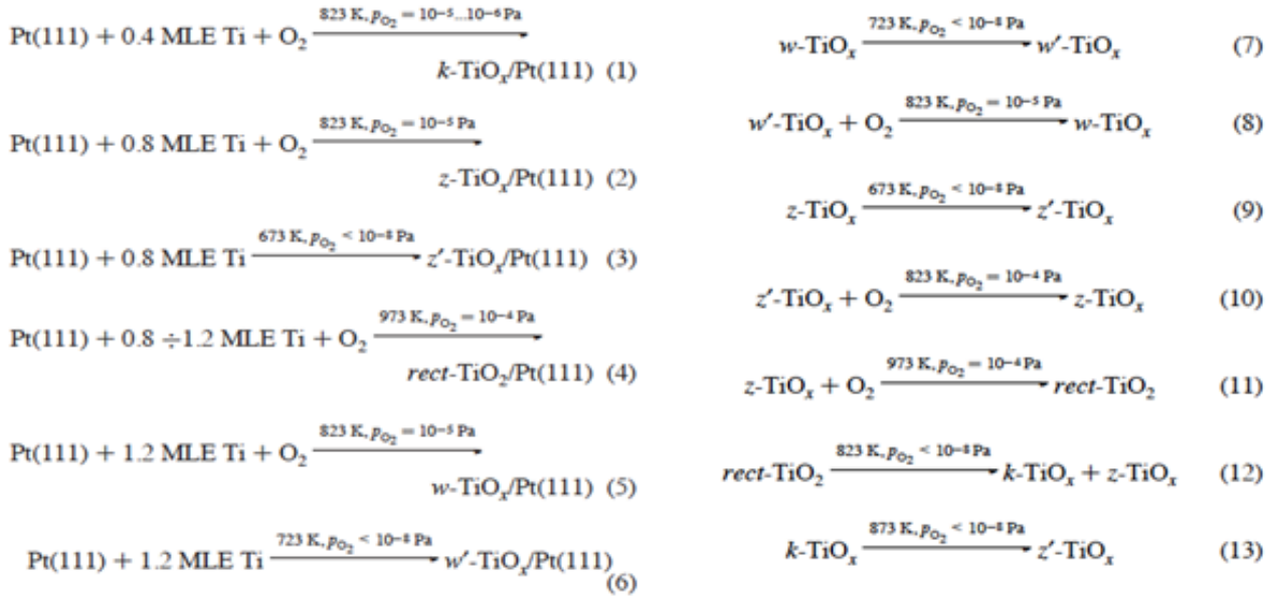


Fig.10: a) Summary of the different ultrathin titanium oxide phases obtained on Pt(111) in different conditions. *k* stands for *kagomè*, *z* for *zig-zag*, *rect* for *rectangular* and *w* for *wagon-wheels*. b) Summary of the structural and chemical transitions observed to occur under different thermal treatment conditions. The prime symbol indicates a reduced phase. Reprinted with permission from Sedona et al. [85] Copyright 2005 American Chemical Society.

The different phases are identified by the geometry of the surface structures observed in atomically resolved STM images. Some of the observed structures include reduced titanium ions. Upon thermal treatments the films could undergo some **structural transformations** while in O₂-rich and O₂-poor conditions reversible **film oxidation and reduction** were observed (figure 10). All the identified phases, with the exception of the stoichiometric TiO₂ phase with rectangular surface symmetry, correspond to a film composed of a Ti-O bilayer, with interfacial Ti. The TiO_x phases wet the Pt substrate, and present long-range order with complex superstructures, observed in LEED patterns. The combination of LEED, XPD and STM data with DFT calculations allowed to identify the atomic geometry of all the studied phases [88]. For instance, the *z*-phase has an incommensurate unit cell with 6.8 Å × 8.6 Å size, and a Ti₆O₈ geometry, while the *z'*-phase has a commensurate unit cell of 16.6 Å × 14.4 Å size (with Ti₂₄O₃₀), geometry (figure 11). Ti 2*p* and O 1*s* core level XPS and valence band photoemission studies [86] allowed also to separate the observed phases in two main groups: a group of three stoichiometric films (*k'*, *rect* and *rect'*) and a group of sub-stoichiometric films (*z*, *z'* and *w*). The valence band photoemission data also revealed some peculiar aspects of the **electronic properties**, like for instance the mixing of the Ti-Pt states near the Fermi level in the sub-stoichiometric films, a clear indication of the presence of the Pt-Ti interface in this group.

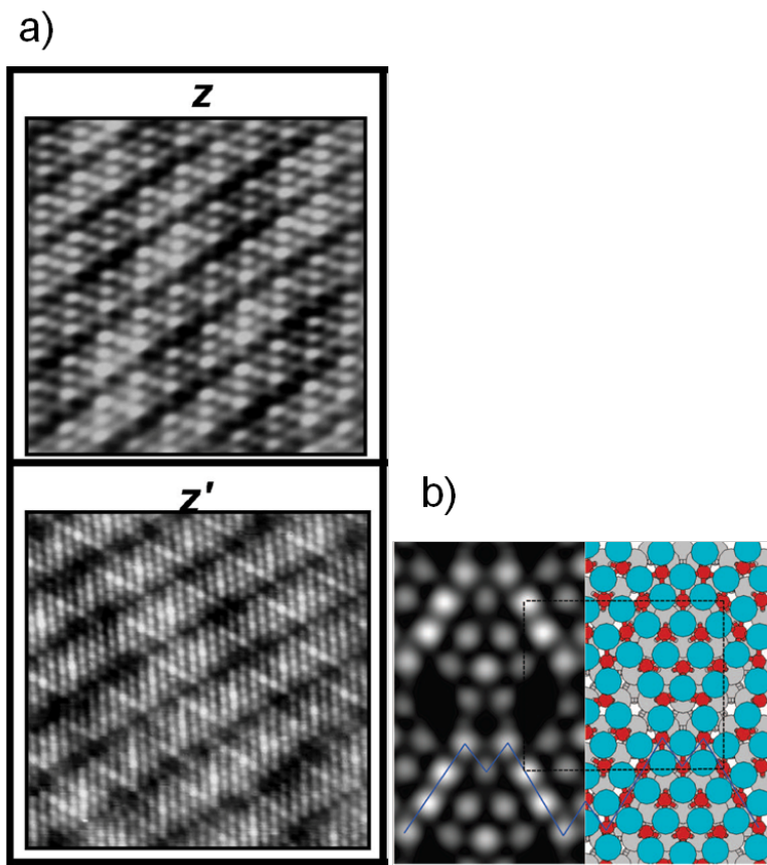


Fig. 11 a) atomically resolved image of the z - and z' - TiO_x phase on Pt(111). b) simulated STM image and structure of the z' phase, as obtained by DFT calculations. Reprinted with permission from Barcaro et al. [88] Copyright 2009 American Chemical Society.

Another interesting phase is the stoichiometric TiO_2 *rect* phase, showing an incommensurate rectangular unit cell of size $3.8 \text{ \AA} \times 3.0 \text{ \AA}$. STM revealed also that the apparent height of the *rect*- TiO_2 islands can be only explained by a multilayer sequence, with oxygen atoms at the interface with Pt occupying only top and bridge sites, an arrangement similar to the one proposed for *rect*- VO_2 [89]. Recently, other reduced hexagonal phases $h(6 \times 6)$ and $h(9 \times 9)$ were discovered by annealing z' phases at high temperature [90].

4. Two-dimensional films of other reducible oxides

4.1 Transition metal oxide two-dimensional films

Indeed, most transition metals, due to their specific electronic structure, can have different oxidation states and can form oxides which are reducible. Within the wide number of studies done in the field of two-dimensional reducible transition metal oxide films, we will here focus on those reporting aspects strictly related to reducibility and we will limit to the strictly two-dimensional case, i.e. to films with thickness of the order or below 1 nm.

A very interesting example of a study of this kind is represented by the work by Li et al. on [manganese oxide two-dimensional films](#) [91]. By STM, LEED and XPS analyses different low dimensionality phases were identified in films of approximately 1 ML thickness as a function of the

oxygen chemical potential during preparation on a Pd(100) substrate. Figure 12 summarizes the obtained results. Among the observed structures, some represent the two-dimensional limit of bulk truncated structures, like the hexagonal structure observed under mildly oxidizing condition, which recalls the MnO(111) structure. Most of them on the contrary do not resemble any known manganese oxide phase and they are often found in coexistence on the Pd surface [92].

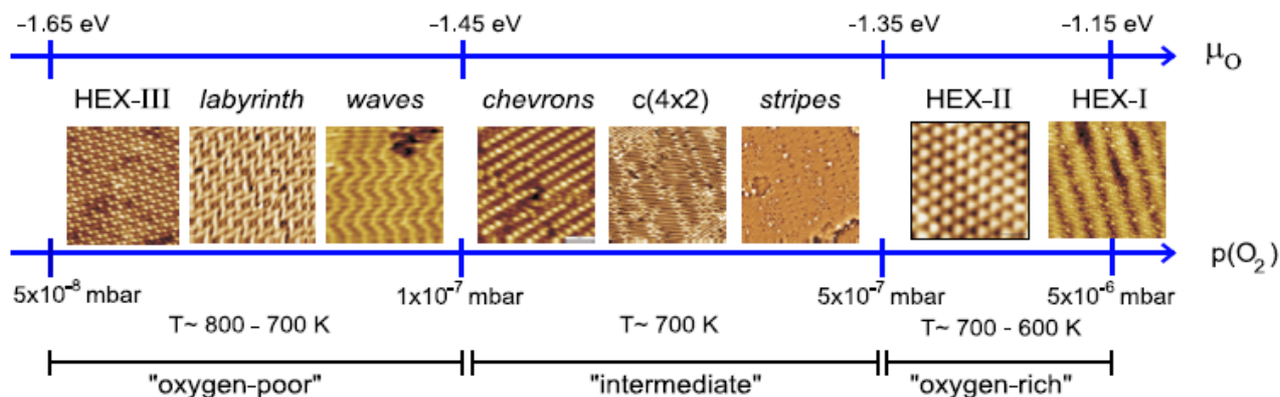


Fig. 12: Schematic phase diagram of the two-dimensional Mn oxides, presented as a function of the oxygen pressure $p(O_2)$ and of the oxygen chemical potential μ_O . The nominal coverage of Mn on Pd(100) is 0.75 ML. © IOP Publishing. Reproduced with permission from Li et al. [91]. All rights reserved.

Also vanadium cations in **vanadium oxide** can have different oxidation states (from 2+ to 5+), giving a wide variety of phases. The most important ones are V_2O_5 , VO_2 , V_2O_3 and VO , each one having a different crystal structure and showing a variety of physical and chemical properties [92]. Furthermore, mixed valence oxides can also be formed, with the cations in two possible oxide states, like for example V^{5+} and V^{4+} in V_6O_{13} . Of peculiar interest is the crystal structure of V_2O_5 (layered orthorhombic), which is essentially composed of zigzag double chains of square VO_5 pyramidal units sharing edges and running along the b direction. The VO_5 pyramid has the vanadium atom in the centre, four oxygen atoms in the basal plane and a vanadyl-type oxygen atom at the apex (see also inset of figure 14d). The layers of chains are stacked along the $[0\ 0\ 1]$ (c) direction. The resulting solid is composed of distorted tetrahedrally coordinated VO_6 unit. The VO_6 unit is also present in the tetrahedral rutile and monoclinic phases of VO_2 which are stable below and above $T=340$ K respectively, while the V_2O_3 phase has a corundum structure above 160 K. All these structures can be found in ultrathin films grown either on oxides or on metals, but other phases can also be stabilised by the interaction with the substrate [89,92].

A systematic work was carried out on vanadium oxide nanostructures and films deposited on Rh(111) by Netzer and co-workers. The experimental results were supported by *ab initio* DFT calculations in order to obtain complete information the structure and on the thermodynamics of the different phases obtained [89,93-96]. A **phase diagram** of the vanadium oxide nanostructures on Rh(111) is reported in figure 13. Concentrating on the two-dimensional phases, it was found that $(\sqrt{7} \times \sqrt{7})R$ 19.1° or $(\sqrt{13} \times \sqrt{13})R$ 13.8° structures can be formed under highly oxidative conditions, with $P_{O_2} = 2 \times 10^{-7}$ mbar and by keeping the substrate temperature at 670 K.

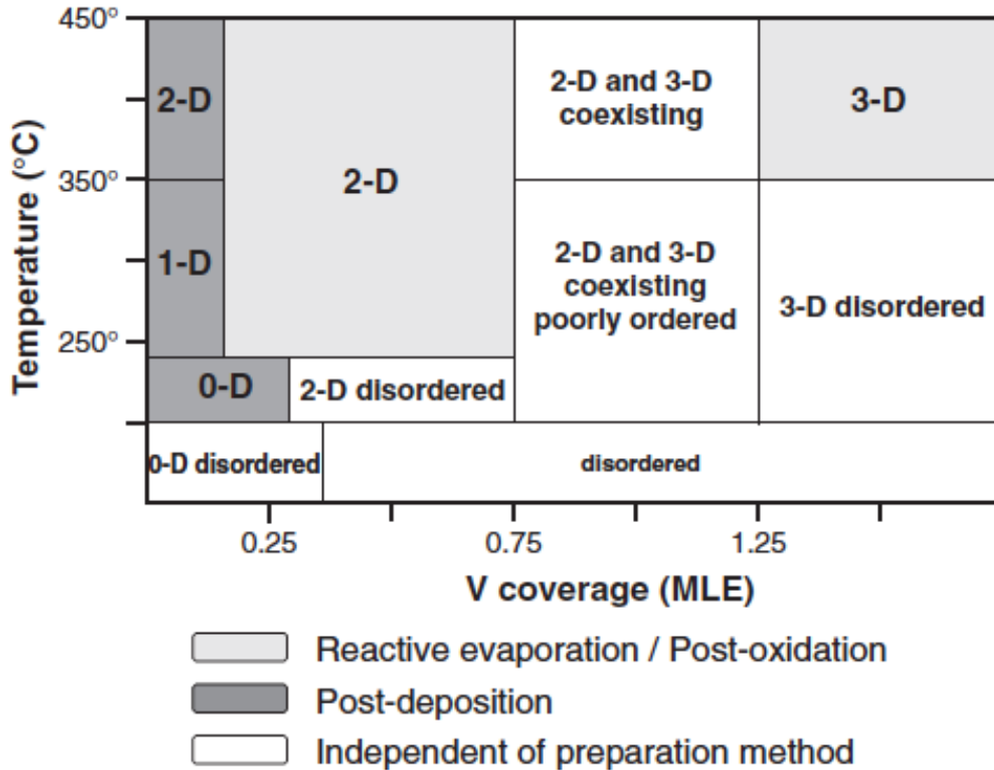


Fig. 13: Phase diagram of vanadium oxide nanostructures on Rh(111) as a function of vanadium coverage and of substrate temperature. © IOP Publishing. Reproduced with permission from J. Schoiswohl et al. [89]. All rights reserved.

As obtained by careful analysis of STM images and by DFT calculations, the $(\sqrt{7} \times \sqrt{7})R 19.1^\circ$ nanolayer consists on a V_3O_9 oxide phase with VO_5 square pyramids as building blocks. The same holds for the $(\sqrt{13} \times \sqrt{13})R 13.8^\circ$ phase, which has a V_6O_{18} stoichiometry.

By exposing the film to reducing conditions (i. e. to annealing in UHV or in hydrogen atmosphere) **reduced two-dimensional phases** can be obtained with $V_{11}O_{23}$, $V_{13}O_{21}$, V_2O_3 and VO stoichiometry. The reduction process occurs essentially by removing the vanadyl groups.

Zero-dimensional structures could also be obtained by evaporation of small quantities (less than 0.2 ML) of vanadium on $O(2 \times 1)$ -Rh(111) and by flashing at 250°C in UHV. The obtained structures are identical planar star-shaped V_6O_{12} molecules [96]. It was shown also that under high substrate temperature the clusters can diffuse and assemble in 2-D overlayers with well determined phases, like (5×5) or the $(5 \times 3\sqrt{3})$ -rect. The different phases are obtained either in oxidizing or in reducing conditions [89].

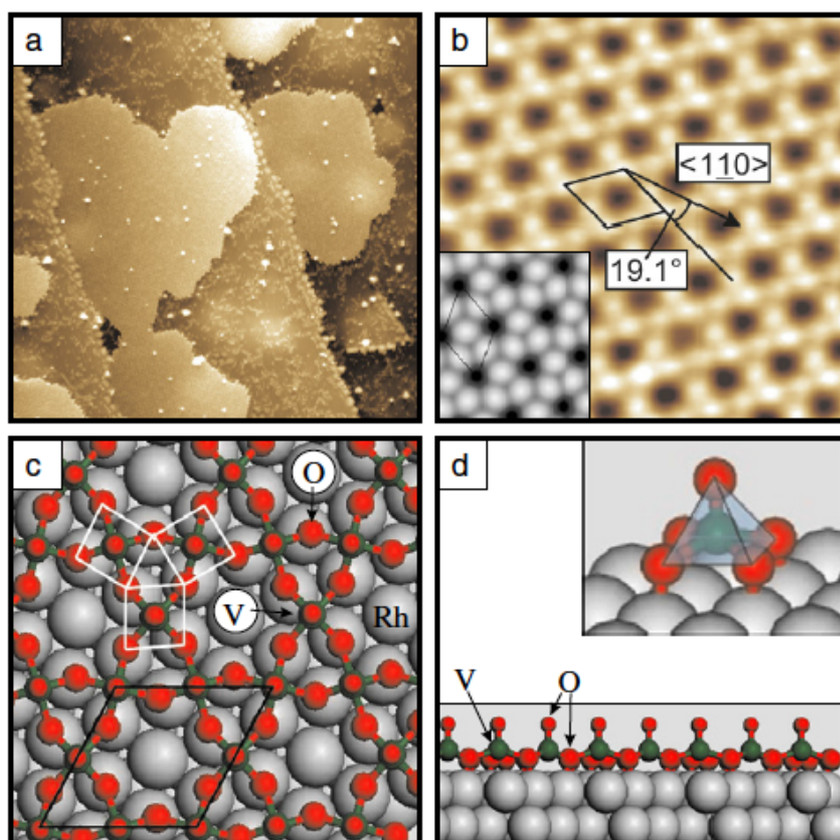


Fig. 14: a) Large scale ($100 \times 100 \text{ nm}^2$) and b) high resolution STM images of the $(\sqrt{7} \times \sqrt{7})R 19.1^\circ$ vanadium oxide nanolayer on Rh(111). Inset simulated STM image obtained by DFT calculations. c) Top view and d) side view of the structural model obtained by DFT. Inset: the VO_5 square pyramid constituting the building block of the film. © IOP Publishing. Reproduced with permission from J. Schoiswohl et al. [89]. All rights reserved.

Another interesting example is the growth of V_2O_y ($y \approx 5$) and V_2O_5 on Au(111) reported by the Freund group [97,98]. The films were obtained by physical vapour deposition of vanadium, subsequent oxidation under 50 mbar of oxygen at 670 K and annealing in UHV at 470 K. The high oxygen pressure used allowed to obtain a high oxidation state (V^{5+}) even for films thicker than 1 MLE (1 MLE corresponding to the same number of V atoms as one layer of Au(111)), at variance with previous studies, where the standard reactive deposition and post-oxidation procedures gave V^{2+} , V^{3+} or V^{4+} states (see for example ref. 89,93,99-101). At low coverage, from 0.26 to 1.04 MLE, two different coincidence monolayer structures could be observed by STM and LEED, which are determined by the **interaction with the Au(111) substrate**. The first structure has a rectangular unit cell, with size $3.6 \text{ \AA} \times 15 \text{ \AA}$, while the second one (observed at 0.52 MLE of V/Au(111) which correspond to one full layer of oxide film), has a $3.6 \text{ \AA} \times 10.8 \text{ \AA}$ ($\alpha = 60^\circ$) oblique unit cell. In analogy with the case of vanadium oxide monolayers on Rh(111) [89,93-96], it was supposed that the full monolayer film consisted of VO_5 pyramid building blocks sharing corners and edges at their bases, and on the basis of the XPS and XANES measurements it was also shown that the oxidation state in the V cations is close to V^{5+} , characteristics of V_2O_5 . Increasing the coverage again to 1.05 MLE, a different structure was observed, which was found to be similar to V_6O_{13} with (001) orientation, while for higher coverage V_2O_5 islands were formed, extending in size and giving rise eventually to V_2O_5 (001) films containing a low number of **point defects**. These films were composed of large (20 nm size) single crystal domains with some **azimuthal disorder** (figure 15) [97].

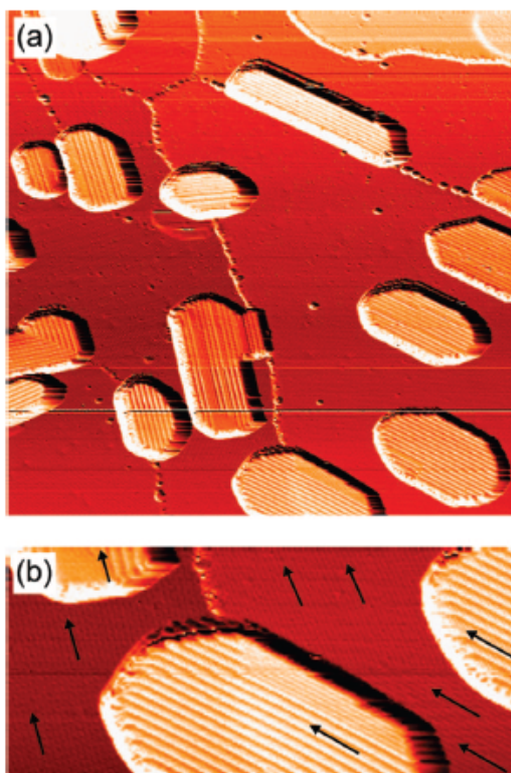


Fig. 15: STM images of a) 100 X 100 nm², b) 44 X 20 nm² of a film formed by the oxidation of 1.56 MLE V/Au(111). The images show the presence of V₂O₅(100) islands growing on V₆O₁₃(001) film. Reprinted with permission from S. Guimond et al. [97] Copyright 2008 American Chemical Society.

4.2 Rare earth oxide two-dimensional films

In analogy with cerium oxide also in other rare earth oxides (REOs), like praseodymium, terbium and samarium oxide the cations can have different oxidation states, and the oxides can be defined as reducible. For samarium oxide the most stable oxidation state is Sm₂O₃, which can form different structural phases. Samarium oxide is the only REO which can also form a monoxide, SmO, with a rock-salt structure, stable under reducing conditions. The most stable valence state for terbium is 3+, leading to the Tb₂O₃ phase, however also the dioxide TbO₂, as well as intermediate phases like Tb₄O₇, can form. Praseodymium can have the 2+, 3+ and 4+ oxidation states and the most stable oxides are Pr₂O₃, PrO₂ and Pr₆O₁₁. The sesquioxides are typically stable in the c-type bixbyite structure, at variance with cerium oxide, for which the hexagonal A-type phase is favoured in the Ce₂O₃ stoichiometry.

Some studies report the growth of REO in the form of ultrathin films on metallic substrates and point out interesting aspects related to those observed in cerium oxide ultrathin films. Temperature programmed desorption (TPD) allows to monitor oxygen release in these REO films, while in the case of cerium oxide the release is considered to be too fast to be monitored [102,103].

Samarium oxide in the form of an ultrathin film has been shown to form by controlled oxidation of surface SmRh and SmRu surface alloys, in turn obtained by heating metallic Sm films deposited on Rh(100) and Ru(0001) surfaces respectively, leading to a SmO_x phase [102-104]. On both substrates specific sites for **CO absorption** have been shown to form at the perimeter of the samarium oxide ultrathin islands, due to the interaction with the underlying metal [102,103]. A

more recent study by Jhang et al. investigated samarium oxide films grown on a Pt(111) substrate by reactive deposition at 600 K followed by annealing in O₂ (P~10⁻⁷ mbar) at 1000 K [105]. In close analogy with cerium oxide films grown in similar conditions, the LEED pattern for submonolayer coverage shows a (1.37 × 1.37) structure. Interestingly, for coverages between 1 and 3 ML the LEED pattern shows additional faint spots in the so called quasi-(3×3) superstructure, very similar to the one observed in reduced ultrathin cerium oxide phases on Pt(111) (figure 16) [17,29]. The authors ascribe this features to a Sm₂O₃(111) phase with a **defective fluorite structure**, in which the oxygen vacancies are randomly distributed within the crystal, and to the formation of a 8:11 coincidence with the underlying Pt. A similar origin may be invoked also for the LEED pattern of reduced cerium oxide films, although in the latter case a similar coincidence may be expected also when the films are oxidized, and the reasons for its appearance only when the films are reduced are not clear. **Reduction** by thermal treatments in UHV at 1000 K for 30 min of the Sm₂O₃ films lead to the coexistence of Sm₂O₃(111) phase and of a SmO(100) phase, giving a superstructure rotated by 30° with respect to the (111) spots in the LEED pattern [105]. The **reversibility** of the structural and morphological modifications is observed also in this case [105].

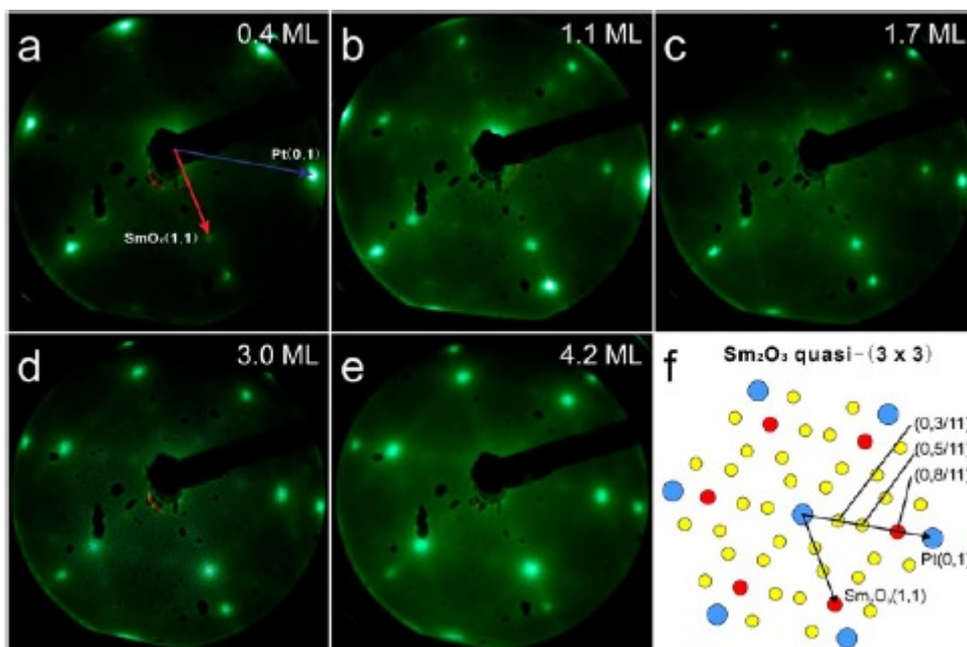


Fig.16: LEED patterns obtained after growing Sm₂O₃(111) films on Pt(111) to the coverages indicated followed by annealing in 7 × 10⁻⁷ mbar of O₂ at 1000 K for 10 min: (a) 0.4 ML of Sm₂O₃ shows a hexagonal (1.37 × 1.37) superstructure in registry with the Pt(111) (1 × 1) spots, E = 58 eV, (b) 1.1 ML, E = 48 eV, (c) 1.7 ML, E = 56 eV, (d) 3.0 ML, E = 52 eV exhibit a quasi-(3 × 3) superstructure, (e) 4.2 ML, E = 56 eV; the quasi-(3 × 3) becoming blurry, and (f) shows a schematic representation of the quasi-(3 × 3) pattern that is determined from FFT analysis of a structural model of superposed, hexagonal Sm and Pt lattices that form a hexagonal (8 × 8) coincidence lattice with respect to the Sm lattice, i.e. (11 × 11) with respect to the Pt lattice. Reprinted with permission from Jhang et al. [105] Copyright 2013 American Chemical Society.

Terbium oxide films were prepared with similar methods, i. e. using reactive deposition and post growth annealing in O₂ [106]. The LEED pattern up to 4 ML thickness shows the (1.32 × 1.32) structure, compatible with the terbium oxide lattice parameter of the Tb₂O₃ phase in a fluorite structure with disordered **oxygen vacancies** in close analogy with the case of samarium oxide [107]. In this case however diffraction from possible coincidence superlattices could not be observed. The

films could not be oxidized by using thermal treatments in O₂ at P ~ 7×10⁻⁷ mbar and temperatures from 95 to 1000 K, while plasma-generated atomic oxygen was shown to be effective in oxidizing a film at 300 K to TbO₂ and in generating weakly bound surface O species. The reduced Tb₂O₃ phase could be re-obtained by annealing TbO₂ in vacuum at 1000 K [106].

Praseodymium oxide has been studied only in the form of relatively thick films on Si(111) substrates [107-109]. As for terbium oxide thin films, also in this case, an **oxidizing treatment** in oxygen plasma was shown to be necessary to oxidize the Pr₂O₃ phase to the PrO₂ fluorite phase [107]. UHV thermal treatments to reduce the PrO₂ phase have been shown to give origin to a phase including a mixture of Pr₆O₁₁ and Pr₅O₉, the latter being unstable in the bulk phase [109].

5. Conclusions

The studies discussed in this chapter show that reducible oxides at two dimensionality may show important modifications of their properties, and indeed also of the reversible reduction and oxidation processes. Ultrathin films, typically supported on metal substrates, at different degree of reduction show new structural and chemical phases, often unstable in the bulk, which have been deeply characterized by surface science techniques, combined with the extremely important support of theoretical modeling. In some cases the different phases can be reversibly transformed one into the other by reducing and oxidizing treatments. The studies point out interesting challenging aspects, on which future studies might be focused, such as for example the stabilization of phases with less stable surface orientation, or more complex systems such as ternary two-dimensional compounds or mixed oxide phases. We believe that investigations following these lines may open up unexpectedly interesting aspects within the field of reducible oxide based materials.

Acknowledgements

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References

1. Trovarelli A, Fornasiero P (2013) Catalysis by Ceria and Related Materials. Catalytic Science, 2nd edn. Imperial College Pr., London
2. Celardo I, Pedersen JZ, Traversa E, Ghibelli L (2011) Pharmacological potential of cerium oxide nanoparticles. *Nanoscale* 3 (4):1411-1420
3. Waser R, Dittmann R, Staikov G, Szot K (2009) Redox-Based Resistive Switching Memories - Nanoionic Mechanisms, Prospects, and Challenges. *Adv Mater* 21 (25-26):2632

4. Migani A, Vayssilov GN, Bromley ST, Illas F, Neyman KM (2010) Dramatic reduction of the oxygen vacancy formation energy in ceria particles: a possible key to their remarkable reactivity at the nanoscale. *J Mater Chem* 20 (46):10535-10546
5. Cafun JD, Kvashnina KO, Casals E, Puentes VF, Glatzel P (2013) Absence of Ce³⁺ Sites in Chemically Active Colloidal Ceria Nanoparticles. *Acs Nano* 7 (12):10726-10732
6. Carretin S, Concepcion P, Corma A, Nieto JML, Puentes VF (2004) Nanocrystalline CeO₂ increases the activity of an for CO oxidation by two orders of magnitude. *Angew Chem Int Edit* 43 (19):2538-2540
7. Vayssilov GN, Lykhach Y, Migani A, Staudt T, Petrova GP, Tsud N, Skala T, Bruix A, Illas F, Prince KC, Matolin V, Neyman KM, Libuda J (2011) Support nanostructure boosts oxygen transfer to catalytically active platinum nanoparticles. *Nat Mater* 10 (4):310-315
8. Tauster SJ, Fung SC, Garten RL (1978) Strong Metal-Support Interactions - Group-8 Noble-Metals Supported on TiO₂. *J Am Chem Soc* 100 (1):170-175
9. Bevan DJM (1955) Ordered intermediate phases in the system CeO₂-Ce₂O₃. *J Inorg Nucl Chem* 1 :49
10. Skorodumova NV, Simak SI, Lundqvist BI, Abrikosov IA, Johansson B (2002) Quantum origin of the oxygen storage capability of ceria. *Phys Rev Lett* 89 (16):166601
11. Conesa JC (1995) Computer Modeling of Surfaces and Defects on Cerium Dioxide. *Surf Sci* 339 (3):337-352
12. Hardacre C, Ormerod RM, Lambert RM (1994) Platinum-Promoted Catalysis by Ceria - a Study of Carbon-Monoxide Oxidation over Pt(111)/CeO₂. *J Phys Chem-U* 98 (42):10901-10905
13. Schierbaum KD (1998) Ordered ultra-thin cerium oxide overlayers on Pt(111) single crystal surfaces studied by LEED and XPS. *Surf Sci* 399 (1):29-38-X
14. Mullins DR, Radulovic PV, Overbury SH (1999) Ordered cerium oxide thin films grown on Ru(0001) and Ni(111). *Surf Sci* 429 (1-3):186-198
15. Eck S, Castellarin-Cudia C, Surnev S, Ramsey MG, Netzer FP (2002) Growth and thermal properties of ultrathin cerium oxide layers on Rh(111). *Surf Sci* 520 (3):173-185
16. Wilson EL, Chen Q, Brown WA, Thornton G (2007) CO adsorption on the model catalyst Pd/CeO₂-x(111)/Rh(111). *J Phys Chem C* 111 (38):14215-14222
17. Luches P, Pagliuca F, Valeri S (2011) Morphology, Stoichiometry, and Interface Structure of CeO₂ Ultrathin Films on Pt(111). *J Phys Chem C* 115 (21):10718-10726
18. Staudt T, Lykhach Y, Hammer L, Schneider MA, Matolin V, Libuda J (2009) A route to continuous ultra-thin cerium oxide films on Cu(111). *Surf Sci* 603 (23):3382-3388
19. Xiao WD, Guo QL, Wang EG (2003) Transformation of CeO₂(111) to Ce₂O₃(0001) films. *Chem Phys Lett* 368 (5-6):527-531
20. Lu JL, Gao HJ, Shaikhutdinov S, Freund HJ (2006) Morphology and defect structure of the CeO₂(111) films grown on Ru(0001) as studied by scanning tunneling microscopy. *Surf Sci* 600 (22):5004-5010
21. Grinter DC, Ithnin R, Pang CL, Thornton G (2010) Defect Structure of Ultrathin Ceria Films on Pt(111): Atomic Views from Scanning Tunnelling Microscopy. *J Phys Chem C* 114 (40):17036-17041

22. Flege JI, Kaemena B, Gevers S, Bertram F, Wilkens T, Bruns D, Batjer J, Schmidt T, Wollschlager J, Falta J (2011) Silicate-free growth of high-quality ultrathin cerium oxide films on Si(111). *Phys Rev B* 84 (23):235418
23. Pagliuca F, Luches P, Valeri S (2013) Interfacial interaction between cerium oxide and silicon surfaces. *Surf Sci* 607:164-169
24. Putna ES, Bunluesin T, Fan XL, Gorte RJ, Vohs JM, Lakis RE, Egami T (1999) Ceria films on zirconia substrates: models for understanding oxygen-storage properties. *Catal Today* 50 (2):343-352
25. vanWijck MAAM, Verhoeven MAJ, Reuvekamp EMCM, Gerritsma GJ, Blank DHA, Rogalla H (1996) CeO₂ as insulation layer in high T-c superconducting multilayer and crossover structures. *Appl Phys Lett* 68 (4):553-555
26. Overbury SH, Huntley DR, Mullins DR, Ailey KS, Radulovic PV (1997) Surface studies of model supported catalysts: NO adsorption on Rh/CeO₂(001). *J Vac Sci Technol A* 15 (3):1647-1652
27. Hardacre C, Roe GM, Lambert RM (1995) Structure, Composition and Thermal-Properties of Cerium Oxide-Films on Platinum(111). *Surf Sci* 326 (1-2):1-10
28. Berner U, Schierbaum KD (2002) Cerium oxides and cerium-platinum surface alloys on Pt(111) single-crystal surfaces studied by scanning tunneling microscopy. *Phys Rev B* 65 (23):235404
29. Luches P, Pagliuca F, Valeri S (2014) Structural and morphological modifications of thermally reduced cerium oxide ultrathin epitaxial films on Pt(111). *Phys Chem Chem Phys* 16 (35):18848-18857
30. Matolin V, Libra J, Matolinova I, Nehasil V, Sedlacek L, Sutara F (2007) Growth of ultra-thin cerium oxide layers on Cu(111). *Appl Surf Sci* 254 (1):153-155
31. Yang F, Graciani J, Evans J, Liu P, Hrbek J, Sanz JF, Rodriguez JA (2011) CO Oxidation on Inverse CeO_x/Cu(111) Catalysts: High Catalytic Activity and Ceria-Promoted Dissociation of O-2. *J Am Chem Soc* 133 (10):3444-3451
32. Ma SG, Rodriguez J, Hrbek J (2008) STM study of the growth of cerium oxide nanoparticles on Au(111). *Surf Sci* 602 (21):3272-3278
33. Škoda M, Libra J, Šutara F, Tsud N, Skála T, Sedláček L, Cháb V, Prince KC, Matolín V (2007) A resonant photoemission study of the Ce and Ce-oxide/Pd(1 1 1) interfaces. *Surf Sci* 601 (21):4958-4965
34. Grinter DC, Yim CM, Pang CL, Santos B, Mentis TO, Locatelli A, Thornton G (2013) Oxidation State Imaging of Ceria Island Growth on Re(0001). *J Phys Chem C* 117 (32):16509-16514
35. Suchorski Y, Wrobel R, Becker S, Weiss H (2008) CO Oxidation on a CeO_x/Pt(111) Inverse Model Catalyst Surface: Catalytic Promotion and Tuning of Kinetic Phase Diagrams. *J Phys Chem C* 112 (50):20012-20017
36. Eck S, Castellarin-Cudia C, Surnev S, Prince KC, Ramsey MG, Netzer FP (2003) Adsorption and reaction of CO on a ceria–Rh(1 1 1) “inverse model catalyst” surface. *Surf Sci* 536 (1–3):166-176
37. Rodriguez JA, Graciani J, Evans J, Park JB, Yang F, Stacchiola D, Senanayake SD, Ma S, Pérez M, Liu P, Sanz JF, Hrbek J (2009) Water-Gas Shift Reaction on a Highly Active Inverse CeO_x/Cu(111) Catalyst: Unique Role of Ceria Nanoparticles. *Angewandte Chemie International Edition* 48 (43):8047-8050

38. Dvorak F, Stetsovykh O, Steger M, Cherradi E, Matolinova I, Tsud N, Skoda M, Skala T, Myslivecek J, Matolin V (2011) Adjusting Morphology and Surface Reduction of CeO₂(111) Thin Films on Cu(111). *J Phys Chem C* 115 (15):7496-7503
39. Luches P, Pagliuca F, Valeri S, Boscherini F (2013) Structure of Ultrathin CeO₂ Films on Pt(111) by Polarization-Dependent X-ray Absorption Fine Structure. *J Phys Chem C* 117 (2):1030-1036
40. Szabova L, Stetsovykh O, Dvorak F, Camellone MF, Fabris S, Myslivecek J, Matolin V (2012) Distinct Physicochemical Properties of the First Ceria Monolayer on Cu(111). *J Phys Chem C* 116 (11):6677-6684
41. Spiel C, Blaha P, Suchorski Y, Schwarz K, Rupprechter G (2011) CeO₂/Pt(111) interface studied using first-principles density functional theory calculations. *Phys Rev B* 84 (4):045412
42. Nilius N, Kozlov SM, Jerratsch JF, Baron M, Shao X, Vines F, Shaikhutdinov S, Neyman KM, Freund HJ (2012) Formation of One-Dimensional Electronic States along the Step Edges of CeO₂(111). *Acs Nano* 6 (2):1126-1133
43. Torbrugge S, Cranney M, Reichling M (2008) Morphology of step structures on CeO₂(111). *Appl Phys Lett* 93 (7):073112
44. Jerratsch JF, Shao X, Nilius N, Freund HJ, Popa C, Ganduglia-Pirovano MV, Burow AM, Sauer J (2011) Electron Localization in Defective Ceria Films: A Study with Scanning-Tunneling Microscopy and Density-Functional Theory. *Phys Rev Lett* 106 (24):246801
45. Esch F, Fabris S, Zhou L, Montini T, Africh C, Fornasiero P, Comelli G, Rosei R (2005) Electron localization determines defect formation on ceria substrates. *Science* 309 (5735):752-755
46. Ganduglia-Pirovano MV, Da Silva JLF, Sauer J (2009) Density-Functional Calculations of the Structure of Near-Surface Oxygen Vacancies and Electron Localization on CeO₂(111). *Phys Rev Lett* 102 (2):026101
47. Loschen C, Bromley ST, Neyman KM, Illas F (2007) Understanding ceria nanoparticles from first-principles calculations. *J Phys Chem C* 111 (28):10142-10145
48. Kaemena B, Senanayake SD, Meyer A, Sadowski JT, Falta J, Flege JI (2013) Growth and Morphology of Ceria on Ruthenium (0001). *J Phys Chem C* 117 (1):221-232
49. Pfau A, Schierbaum KD (1994) The Electronic-Structure of Stoichiometric and Reduced CeO₂ Surfaces - an Xps, Ups and Hreels Study. *Surf Sci* 321 (1-2):71-80
50. Matolin V, Libra J, Skoda M, Tsud N, Prince KC, Skala T (2009) Methanol adsorption on a CeO₂(111)/Cu(111) thin film model catalyst. *Surf Sci* 603 (8):1087-1092
51. Senanayake SD, Stacchiola D, Evans J, Estrella M, Barrio L, Perez M, Hrbek J, Rodriguez JA (2010) Probing the reaction intermediates for the water-gas shift over inverse CeO_x/Au(111) catalysts. *J Catal* 271 (2):392-400
52. Castellarin-Cudia C, Surnev S, Schneider G, Podlucky R, Ramsey MG, Netzer FP (2004) Strain-induced formation of arrays of catalytically active sites at the metal-oxide interface. *Surf Sci* 554 (2-3):L120-L126
53. Duchon T, Dvorak F, Aulicka M, Stetsovykh V, Vorokhta M, Mazur D, Veltruska K, Skala T, Myslivecek J, Matolinova I, Matolin V (2014) Ordered Phases of Reduced Ceria As Epitaxial Films on Cu(111). *J Phys Chem C* 118 (1):357-365

54. Wilkens H, Schuckmann O, Oelke R, Gevers S, Reichling M, Schaefer A, Baumer M, Zoellner MH, Niu G, Schroeder T, Wollschlager J (2013) Structural transitions of epitaxial ceria films on Si(111). *Phys Chem Chem Phys* 15 (42):18589-18599
55. Stetsovych V, Pagliuca F, Dvorak F, Duchon T, Vorokhta M, Aulicka M, Lachnitt J, Schernich S, Matolinova I, Veltruska K, Skala T, Mazur D, Myslivecek J, Libuda J, Matolin V (2013) Epitaxial Cubic Ce₂O₃ Films via Ce-CeO₂ Interfacial Reaction. *J Phys Chem Lett* 4 (6):866-871
56. Kozlov SM, Demiroglu I, Neyman KM, Bromley ST (2015) Reduced ceria nanofilms from structure prediction. *Nanoscale* 7 (10):4361-4366
57. Yang F, Choi Y, Agnoli S, Liu P, Stacchiola D, Hrbek J, Rodriguez JA (2011) CeO₂ <-> CuO_x Interactions and the Controlled Assembly of CeO₂(111) and CeO₂(100) Nanoparticles on an Oxidized Cu(111) Substrate. *J Phys Chem C* 115 (46):23062-23066
58. Stetsovych O, Beran J, Dvorak F, Masek K, Myslivecek J, Matolin V (2013) Polarity driven morphology of CeO₂(100) islands on Cu(111). *Appl Surf Sci* 285:766-771
59. Pan Y, Niluis N, Stiehler C, Freund HJ, Goniakowski J, Noguera C (2014) Ceria Nanocrystals Exposing Wide (100) Facets: Structure and Polarity Compensation. *Adv Mater Interfaces* 1 (9):1400404
60. Noguera C, Goniakowski J (2013) Polarity in Oxide Nano-objects. *Chem Rev* 113 (6):4073-4105
61. Maeda T, Kobayashi Y, Kishi K (1999) Growth of ultra-thin titanium oxide on Cu(100), Fe/Cu(100) and ordered ultra-thin iron oxide studied by low-energy electron diffraction and X-ray photoelectron spectroscopy. *Surf Sci* 436 (1-3):249-258
62. Passeggi MCG, Vergara LI, Mendoza SM, Ferron J (2002) Passivation and temperature effects on the oxidation process of titanium thin films. *Surf Sci* 507:825-831
63. Finetti P, Caffio M, Cortigiani B, Atrei A, Rovida G (2008) Mechanism of growth and structure of titanium oxide ultrathin films deposited on Cu(001). *Surf Sci* 602 (5):1101-1113
64. Atrei A, Bardi U, Rovida G (1997) Structure and composition of the titanium oxide layers formed by low-pressure oxidation of the Ni₉₄Ti₆(110) surface. *Surf Sci* 391 (1-3):216-225
65. Atrei A, Ferrari AM, Finetti P, Beni A, Rovida G (2009) LEED and DFT Study of the Quasihexagonal TiO₂ Structure on Cu(001). *J Phys Chem C* 113 (45):19578-19584
66. Atrei A, Ferrari AM, Szieberth D, Cortigiani B, Rovida G (2010) Lepidocrocite-like structure of the TiO₂ monolayer grown on Ag(100). *Phys Chem Chem Phys* 12 (37):11587-11595
67. Atrei A, Cortigiani B, Ferrari AM (2012) Epitaxial growth of TiO₂ films with the rutile (110) structure on Ag(100). *J Phys-Condens Mat* 24 (44):445005
68. Orzali T, Casarin M, Granozzi G, Sambri M, Vittadini A (2006) Bottom-up assembly of single-domain titania nanosheets on (1x2)-Pt(110). *Phys Rev Lett* 97 (15):156101
69. McCavish ND, Bennett RA (2003) Ultra-thin film growth of titanium dioxide on W(100). *Surf Sci* 546 (1):47-56
70. Bennett RA, Mulley JS, Newton MA, Surman M (2007) Spectroscopy of ultrathin epitaxial rutile TiO₂(110) films grown on W(100). *J Chem Phys* 127 (8):084707

71. Pang CL, Grinter DC, Matharu J, Thornton G (2013) A Scanning Tunneling Microscopy Study of Ultrathin Film Rutile TiO₂(110) Supported on W(100)-O(2 x 1). *J Phys Chem C* 117 (48):25622-25627
72. Oh WS, Xu C, Kim DY, Goodman DW (1997) Preparation and characterization of epitaxial titanium oxide films on Mo(100). *J Vac Sci Technol A* 15 (3):1710-1716
73. Chen MS, Goodman DW (2008) Ultrathin, ordered oxide films on metal surfaces. *J Phys-Condens Mat* 20 (26):264013
74. Guo Q, Oh WS, Goodman DW (1999) Titanium oxide films grown on Mo(110). *Surf Sci* 437 (1-2):49-60
75. Lai X, Guo Q, Min BK, Goodman DW (2001) Synthesis and characterization of titania films on Mo(110). *Surf Sci* 487 (1-3):1-8
76. Chen MS, Goodman DW (2004) The structure of catalytically active gold on titania. *Science* 306 (5694):252-255
77. Matsumoto T, Batzill M, Hsieh S, Koel BE (2004) Fundamental studies of titanium oxide-Pt(100) interfaces - I. Stable high temperature structures formed by annealing TiO_x films on Pt(100). *Surf Sci* 572 (2-3):127-145
78. Papageorgiou AC, Cabailh G, Chen Q, Resta A, Lundgren E, Andersen JN, Thornton G (2007) Growth and reactivity of titanium oxide ultrathin films on Ni(110). *J Phys Chem C* 111 (21):7704-7710
79. Papageorgiou AC, Pang CL, Chen Q, Thornton G (2007) Low-dimensional, reduced phases of ultrathin TiO₂. *Acs Nano* 1 (5):409-414
80. Bennett RA, Poulston S, Stone P, Bowker M (1999) STM and LEED observations of the surface structure of TiO₂(110) following crystallographic shear plane formation. *Phys Rev B* 59 (15):10341-10346
81. Dulub O, Hebenstreit W, Diebold U (2000) Imaging cluster surfaces with atomic resolution: The strong metal-support interaction state of Pt supported on TiO₂(110). *Phys Rev Lett* 84 (16):3646-3649
82. Bennett RA, Pang CL, Perkins N, Smith RD, Morrall P, Kvon RI, Bowker M (2002) Surface structures in the SMSI state; Pd on (1 x 2) reconstructed TiO₂(110). *J Phys Chem B* 106 (18):4688-4696
83. Matsumoto T, Batzill M, Hsieh S, Koel BE (2004) Fundamental studies of titanium oxide-Pt(100) interfaces - II. Influence of oxidation and reduction reactions on the surface structure of TiO_x films on Pt(100). *Surf Sci* 572 (2-3):146-161
84. Boffa AB, Galloway HC, Jacobs PW, Benitez JJ, Batteas JD, Salmeron M, Bell AT, Somorjai GA (1995) The Growth and Structure of Titanium-Oxide Films on Pt(111) Investigated by Leed, Xps, Iss, and Stm. *Surf Sci* 326 (1-2):80-92
85. Sedona F, Rizzi GA, Agnoli S, Xamena FXLI, Papageorgiou A, Ostermann D, Sambri M, Finetti P, Schierbaum K, Granozzi G (2005) Ultrathin TiO_x films on Pt(111): A LEED, XPS, and STM investigation. *J Phys Chem B* 109 (51):24411-24426
86. Sedona F, Agnoli S, Granozzi G (2006) Ultrathin wagon-wheel-like TiO_x phases on Pt(111): A combined low-energy electron diffraction and scanning tunneling microscopy investigation. *J Phys Chem B* 110 (31):15359-15367

87. Finetti P, Sedona F, Rizzi GA, Mick U, Sutara F, Svec M, Matolin V, Schierbaum K, Granozzi G (2007) Core and valence band photoemission Spectroscopy of well-ordered ultrathin TiO_x films on Pt(111). *J Phys Chem C* 111 (2):869-876
88. Barcaro G, Agnoli S, Sedona F, Rizzi GA, Fortunelli A, Granozzi G (2009) Structure of Reduced Ultrathin TiO_x Polar Films on Pt(111). *J Phys Chem C* 113 (14):5721-5729
89. Schoiswohl J, Surnev S, Netzer FP, Kresse G (2006) Vanadium oxide nanostructures: from zero- to three-dimensional. *J Phys-Condens Mat* 18 (4):R1-R14
90. Barcaro G, Cavaliere E, Artiglia L, Sementa L, Gavioli L, Granozzi G, Fortunelli A (2012) Building Principles and Structural Motifs in TiO_x Ultrathin Films on a (111) Substrate. *J Phys Chem C* 116 (24):13302-13306
91. Li F, Parteder G, Allegretti F, Franchini C, Podloucky R, Surnev S, Netzer FP (2009) Two-dimensional manganese oxide nanolayers on Pd(100): the surface phase diagram. *J Phys-Condens Mat* 21 (13):134008
92. Surnev S, Ramsey MG, Netzer FP (2003) Vanadium oxide surface studies. *Prog Surf Sci* 73 (4-8):117-165
93. Surnev S, Kresse G, Ramsey MG, Netzer FP (2001) Novel interface-mediated metastable oxide phases: Vanadium oxides on Pd(111). *Phys Rev Lett* 87 (8):086102
94. Schoiswohl J, Sock M, Eck S, Surnev S, Ramsey MG, Netzer FP, Kresse G (2004) Atomic-level growth study of vanadium oxide nanostructures on Rh(111). *Phys Rev B* 69 (15):155403
95. Schoiswohl J, Surnev S, Sock M, Eck S, Ramsey MG, Netzer FP, Kresse G (2005) Reduction of vanadium-oxide monolayer structures. *Phys Rev B* 71 (16):165437
96. Schoiswohl J, Kresse G, Surnev S, Sock M, Ramsey MG, Netzer FP (2004) Planar vanadium oxide clusters: Two-dimensional evaporation and diffusion on Rh(111). *Phys Rev Lett* 92 (20):206103
97. Guimond S, Gobke D, Romanyshyn Y, Sturm JM, Naschitzki M, Kuhlenbeck H, Freund HJ (2008) Growth and characterization of ultrathin V(2)O(y) (y approximate to 5) films on Au(111). *J Phys Chem C* 112 (32):12363-12373
98. Guimond S, Sturm JM, Gobke D, Romanyshyn Y, Naschitzki M, Kuhlenbeck H, Freund HJ (2008) Well-ordered V₂O₅(001) thin films on Au(111): Growth and thermal stability. *J Phys Chem C* 112 (31):11835-11846
99. Petukhov M, Rizzi GA, Granozzi G (2001) Spectroscopic and structural characterisation of a VO_x (x approximate to 1) ultrathin epitaxial film on Pt (111). *Thin Solid Films* 400 (1-2):154-159
100. Petukhov M, Rizzi GA, Granozzi G (2001) Ultrathin film growth and spectroscopic characterization of VO_x (0.8 <= x <= 1.3) on Pt(111). *Surf Sci* 490 (3):376-384
101. Surnev S, Kresse G, Sock M, Ramsey MG, Netzer FP (2001) Surface structures of ultrathin vanadium oxide films on Pd(111). *Surf Sci* 495 (1-2):91-106
102. Jiang ZQ, Zhou WP, Tan DL, Zhai RS, Bao XH (2004) Evidence for perimeter sites over SmO_x-modified Rh(100) surface by CO chemisorption. *Surf Sci* 565 (2-3):269-278

103. Kuriyama T, Kunimori K, Nozoye H (1998) Interaction of CO with the SmOx/Ru(001) interface. Surf Sci 402 (1-3):178-181
104. Zhu JF, Ma YS, Zhuang SX (2009) Exploring the role of samarium in the modification of rhodium catalysts through surface science approach. Surf Sci 603 (10-12):1802-1811
105. Jhang JH, Schaefer A, Cartas W, Epuri S, Baeumer M, Weaver JF (2013) Growth and Partial Reduction of Sm₂O₃(111) Thin Films on Pt(111): Evidence for the Formation of SmO(100). J Phys Chem C 117 (41):21396-21406
106. Cartas W, Rai R, Sathe A, Schaefer A, Weaver JF (2014) Oxidation of a Tb₂O₃(111) Thin Film on Pt(111) by Gas-Phase Oxygen Atoms. J Phys Chem C 118 (36):20916-20926
107. Schaefer A, Gevers S, Zielasek V, Schroeder T, Falta J, Wollschlager J, Baumer M (2011) Photoemission study of praseodymia in its highest oxidation state: The necessity of in situ plasma treatment. J Chem Phys 134 (5):054701
108. Gevers S, Weisemoeller T, Schaefer A, Zielasek V, Baumer M, Wollschlager J (2011) Structure of oxygen-plasma-treated ultrathin praseodymia films on Si(111). Phys Rev B 83 (19):193408
109. Wilkens H, Gevers S, Rohe S, Schaefer A, Baumer M, Zoellner MH, Schroeder T, Wollschlager J (2014) Structural Changes of Ultrathin Cub-PrO₂(111)/Si(111) Films Due to Thermally Induced Oxygen Desorption. J Phys Chem C 118 (6):3056-3061