Reduction of Hydrogenated ZrO₂ Nanoparticles by Water Desorption

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ABSTRACT: Reduction of zirconia by water desorption from a hydrogenated surface is the topic of this study. The focus is on the role of nanostructuring the oxide reducibility measured by the cost of formation of oxygen vacancies by water desorption. We have performed density functional theory calculations using the Perdew–Burke–Ernzerhof + U approach and including dispersion forces on the adsorption, dissociation, diffusion of hydrogen on the ZrO₂ (101) surface and on Zr₁₆O₃₂,Zr₄₀O₈₀, and Zr₈₀O₁₆₀ nanoparticles (NPs). The process involves the formation of a precursor state via diffusion of hydrogen on the surface of zirconia. The results show that O vacancy formation via H₂O desorption is more convenient than via direct O₂ desorption. The formation of an O₃H₂ surface precursor state to water desorption is the rate-determining step. This step is highly unfavorable on the ZrO₂ (101) surface both thermodynamically and kinetically. On the contrary, on zirconia NPs, characterized by the presence of low coordinated ions, water desorption becomes accessible such that even at temperatures close to 450 K the reaction becomes exergonic. The study shows the role of nanostructuring on the chemical and electronic properties of an oxide.

1. INTRODUCTION

Metal oxides represent a very wide class of heterogeneous catalysts. They can behave as inert supports for an active metal particle or can directly participate in the catalytic process via active surface species and/or basic/acid sites. The composition, structure, and size of the oxide catalyst determine its chemical activity. Zirconium dioxide (ZrO₂) has been extensively used in various catalytic processes. For example, in reactions involving hydrogen, zirconia is used in hydrogenation reactions of CO₂ and CO to synthesize organic molecules, as well as in solid oxide fuel cells for the electrochemical oxidation of H₂ thanks to its stability over a wide range of oxygen pressure and mechanical strength. In another context, zirconia is used as a catalyst for the ketonization reactions in the upgrading of bio-oil to reduce the oxygen content of the biomass after a pyrolysis treatment.

ZrO₂ is considered to be a nonreducible oxide, more similar to MgO and SiO₂ than to TiO₂ or CeO₂. This can be attributed to the high ionicity and wide band gap of ZrO₂. This is a property observed in large crystallites of zirconia, generally referred to as bulk zirconia. The nonreducible nature of zirconia emerges quite clearly when the material is exposed to hydrogen. The H₂ molecule absorbs and dissociates on extended zirconia surfaces through a heterolytic mechanism in which OH⁻ and ZrH⁺ surface groups are formed. This means that no oxide reduction takes place (no formation of adsorbed protons and of Zr³⁺ ions). Another indicator of the low reducibility of this oxide comes from the cost to create a surface O vacancy in the bulk or on the surface of the material (~6 eV). However, the physical and chemical properties of oxides can change dramatically when the material is prepared in the form of nanoparticles (NPs), nanowires, and thin films or in general is nanostructured. In fact, quantum confinement effects, undercoordination, and the special structural flexibility typical of many nanostructures, often referred to as fluxionality, result in novel properties that are not observed in the bulk regime. For instance, we have recently found that zirconia prepared in the form of 1–2 nm particles splits the H₂ molecule through a homolytic process, with the formation of Zₓ⁺ (4dₓ²) centers. This is the same mechanism observed on reducible oxides such as TiO₂ and CeO₂. In this process, two OH⁻ species form and two electrons (coming from the H₂ molecule) reduce two low coordinated Zr⁴⁺ to Zr³⁺ centers (giving rise to a magnetic ground state). In addition, the formation energy of a neutral O vacancy decreases by 2–3 eV when it is created on special sites of the NP. The reducible character of the nanostructures comes from the presence of low coordinated Zr sites in edges and corners, which introduce low-lying Zr 4d states that accept the charge released from the H₂ molecule or from the removed O atom.

In general, the interest toward novel forms of reduced ZrO₂ is increasing. For example, it has been demonstrated that O-deficient ZrO₂₋ₓ NPs display better catalytic activity in reactions of transformation of biomass into fuels.
enthalpy and energy barrier explain the experimental and dissociation, showing that zirconia NPs behave in a presence of oxygen vacancies.26

Apart from direct interaction with H2, there are several mechanisms for the reduction of the oxide surface. In the presence of deposited metal particles, O surface ions can spill over the metal, leaving a surface O vacancy (reverse spillover). This process has been estimated to cost only 0.2 and 0.6 eV on a Ru10 cluster supported on TiO2 and ZrO2, respectively.27 The cost is expected to be even lower at surface steps and edges and is further reduced when the metallic clusters are supported on an oxide NP.30 The oxide surface can also be reduced by a hydrogenation process and subsequent desorption of water, leaving behind an O vacancy and its associated charge (H2 + O2− → H2O + 2e− (Vo)). In the latter case, the presence of a supported metal particle can facilitate the dissociation of H2 and the spillover of hydrogen on the surface to form OH groups and reduced metal cations. Alternatively, O surface ions can spill over the metal particle and react with the adsorbed H atoms to form H2O, leaving behind an O vacancy.31–33

In the absence of supported metal particles, the direct reduction process ZrO2 + H2 → ZrO2−x + H2O is highly endothermic for extended zirconia surfaces because of the low reactivity toward H2 and the high cost to create an O vacancy (~6 eV).12 Hofmann et al. found an enthalpy of +2.71 eV for the reduction reaction on the (101) surface of tetragonal zirconia from density functional theory (DFT) Perdew–Burke–Ernzerhof (PBE) calculations.34 A higher value, +3.57 eV, has been found by our group using the PBE + U approach that improves the description of the zirconia band gap. The process involves an energy barrier of 2.55 eV.35 The large enthalpy and energy barrier explain the experimental observation that H2 desorption instead of H2O desorption takes place from the ZrO2 surface when the hydroxylated Ru/ZrO2 system is heated to 370 K.36 This study follows other studies from our group on the topic of zirconia NPs and related properties. We first analyzed the general features and the electronic structure of zirconia NPs of different sizes, elucidating the role of low coordinated ions and the corresponding presence of new defect states in the band gap of the material.37–39 In this work, we also considered the cost of formation of oxygen vacancies via O2 desorption, as this is a possible route to generate reduced samples. In a subsequent study, we have considered metal adsorption on the ZrO2 NP, in particular, the interaction of Au atoms and their ability to exchange charge with the oxide support as a function of the particle size.40 In a third study, we considered H2 adsorption and dissociation, showing that zirconia NPs behave in a completely different way compared to bulk zirconia as far as interaction with hydrogen is concerned.41 In the present study, the work has been extended toward another important topic: the formation and desorption of water from hydrogenated ZrO2 NPs. In particular, we have investigated two steps of the reduction process on zirconia NPs from both thermodynamic and kinetic points of view. First, the H2 molecule is dissociated on the NP surface (step I). Second, one H atom diffuses into the OH group to form the O2H2 surface complex that is then desorbed as H2O (step II).

I—hydrogenation ZrO2 + H2 → H2ZrO2

II—dehydration H2ZrO2 → ZrO2−x + H2O

In addition, we have investigated the barriers for diffusion of the H atoms into distant adsorption sites, a process that contrast the formation of an O2H2 surface complex. In general, water desorption from a hydroxylated oxide surface is a viable mechanism, leading to the reduction of an oxide. The present work shows the relevant role of nanostructuring in favoring the process that can result in substantial differences in the chemical behavior of the same material when prepared in a bulk form or in the form of nanostructures.

2. COMPUTATIONAL DETAILS

DFT-based calculations were performed with the Vienna Ab-initio Simulation Package (VASP 5.3),40 with plane waves as basis sets with a kinetic energy cutoff of 400 eV. The PBE exchange–correlation functional, within the generalized gradient approximation (GGA), was applied.41 The GGA + U approach implemented by Dudarev et al.42,43 was used to partly correct the self-interaction error intrinsic in GGA functionals. An on-site Coulomb correction, U eff = U − J, was set to 4 eV for the 4d states of Zr atoms. This value provided lattice parameters for tetragonal ZrO2: a0 = 3.662 Å and c0 = 5.223 Å, in good agreement with the experimental ones, 3.64 and 5.27 Å, respectively.44 The calculated Kohn–Sham band gap, 4.5 eV, is still considerably smaller than the experimental value 5.78 eV.45

The DFT + U approach is not free from limitations and should be considered as a pragmatic way to partly remove the self-interaction error inherent to DFT. Furthermore, U values depend on coordination and might change during the transition-state search. In this respect, the results presented in the following must be considered as qualitative rather than quantitative. However, the main purpose of this work is to compare, using the same approach, the properties of zirconia in an extended or a nanostructured form. In this respect, because some of the energies discussed for the two sets of systems differ substantially, the results and the general conclusions are not expected to depend on the choice of the U parameter.

Long-range dispersion forces are included by means of the D2' method46 on top of the PBE functional; this is a slight modification of the original parametrization of the D2 approach proposed by Grimme47 in which we changed the parameters C6 and R0 of the DFT-D2 approach, as suggested by Tosoni and Sauer.

The (101) surface of tetragonal ZrO2 was modeled with a five-layer 3 × 2 supercell containing a total of 180 atoms (Zr60O120). The lattice parameters were fixed at the bulk t-ZrO2 optimized structure obtained with a plane-wave basis set with a kinetic energy of up to 600 eV and with an 8 × 8 × 8 Monkhorst–Pack k-point grid. We modeled octahedral-based zirconia NPs with compositions Zr60O120, Zr60O80 and Zr60O160 having diameters of 0.9, 1.5, and 1.9 nm, respectively.7,8 The models were cut from bulk tetragonal ZrO2, exhibiting {101} surfaces in the octahedral facets and {100} surfaces in the corner, which are the two most stable surfaces for this phase. The geometry optimizations of both slab and NPs were carried out at the Γ-point, up to ionic forces smaller than 0.05 eV/Å and a threshold of 10−5 eV for the electronic self-consistent cycles.

The reaction energy for the hydrogenation (step I) and water desorption (step II) has been calculated according to
\[ \Delta E_1 = E(H_2ZrO_2) - E(ZrO_2) - 1/2E(H_2) \] (1)

\[ \Delta E_II = E(ZrO_2_{x-v}) + E(H_2O(g)) - E(H_2ZrO_2) \] (2)

where \( E(H_2ZrO_2) \) is the energy of the hydrogenated zirconia, \( E(ZrO_2) \) is the energy of clean zirconia, \( E(ZrO_2_{x-v}) \) is the energy of zirconia with one O vacancy, and \( E(H_2) \) and \( E(H_2O(g)) \) are the energies of \( H_2 \) and \( H_2O \) molecules in the gas phase, respectively. The same equations are used to calculate the reaction Gibbs free energy, \( \Delta G \), of the reaction steps. The free energies \( G(ZrO_2) \) and \( G(ZrO_2_{x-v}) \) are approximated to the PBE + U energy, and those for \( H_2 \), \( H_2O \), and \( H_2ZrO_2 \) are defined according to

\[ G(H_2(g), H_2O(g)) \approx E_{PBE} + ZPE - T\cdot S \] (3)

\[ G(H_2ZrO_2) \approx E_{PBE} + ZPE \] (4)

where ZPE is the zero-point energy calculated as \( \left( \frac{\sum \nu \cdot \hbar}{2} \right) \), for which only O–H and Zr–H vibrations (\( \nu \)) are considered, \( T \) is the temperature, and \( S \) is the total entropy. In the \( G(H_2ZrO_2) \) term, the vibrational internal energy, \( U_{vib} \), and the vibrational entropy, \( S_{vib} \), have been neglected. This is justified because OH and ZrH vibrations, in the range of 3000–3700 and 1400 cm\(^{-1} \), respectively, yield \( U_{vib} \) and \( S_{vib} \) terms on the order of \( 10^{-6} \) eV only. The entropy of \( H_2 \) and \( H_2O \) molecules in the gas phase is calculated as the sum of the translational, \( S_{trans} \), vibrational, \( S_{vib} \), and rotational, \( S_{rot} \), contributions according to eqs S–7

\[ S_{trans} = k_B \left( \frac{5}{2} + \ln q_{trans} \right) \] (5)

\[ S_{vib} = k_B \left[ \frac{\theta / T}{e^{\theta / T} - 1} - \ln(1 - e^{-\theta / T}) \right] \] (6)

\[ S_{rot} = k_B \left[ 1 + \ln \left( \frac{IT}{I} \right) + \ln \left( \frac{8\pi^2 k_B}{h^3} \right) \right] \] (7)

where \( q \) is the partition function, \( \theta \) is the vibrational temperature (\( h\nu/k_B \)), and \( I \) is the inertia moment. At 298 K and 1 bar, our calculated entropies for \( H_2 \) and \( H_2O \) are 131 and 189 J/mol-K, respectively, perfectly agree with those reported in the literature.

Finally, transition states are computed by means of the nudged elastic band method using the climbing image approach\(^{39} \) and fully characterized by pertinent vibrational analysis.

### 3. RESULTS AND DISCUSSION

#### 3.1. Thermodynamics of Hydrogenation and Dehydration of ZrO\(_2\) NPs

The first step of the reduction process investigated is the adsorption and dissociation of a \( H_2 \) molecule on the zirconia surface (hydrogenation). As mentioned above, bulk ZrO\(_2\) and ZrO\(_2\) (101) surfaces are nonreducible. The regular (101) surface dissociates the \( H_2 \) molecule through a heterolytic mechanism into \( H^+ \) and \( H^- \) ions, Figure 1a, with a slightly exothermic process, \(-0.06\) eV, Table 1. The Gibbs free energy for the dissociation is endergonic at 298 K and 1 bar, with \( \Delta G_1 = +0.34 \) eV, because of the entropy of \( H_2 \) gas that hinders the adsorption and dissociation.

A completely different situation is obtained on NPs. An important aspect to consider in the study of ZrO\(_2\) NPs is how the chosen structures are representative of real NPs. First of all, we decided to consider three stoichiometric particles of different sizes: \( Zr_{80}O_{160}, Zr_{40}O_{80} \) and \( Zr_{16}O_{32} \). The work performed so far shows that more than the size what counts are the morphological aspects and the presence of low coordinated sites. In this respect, the smallest and largest particles considered, \( Zr_{80}O_{160} \) and \( Zr_{40}O_{80} \) are electronically similar and behave differently from the zirconia (101) surface, whereas the intermediate size \( Zr_{40}O_{80} \) has a general behavior, which is reminiscent of that of the bare zirconia surface. We modeled adsorption of the two \( H_2 \) on the corners of the NPs as these sites are expected to be more reactive because of their low coordination. The dissociation of \( H_2 \) on these sites is largely exothermic, Table 1. The reaction energy of the hydrogenation step is \(-1.79, -1.80, \) and \(-1.62 \) eV, respectively, on \( Zr_{80}O_{160}, Zr_{40}O_{80} \) and \( Zr_{16}O_{32} \), Table 1. We have recently reported such high reactivity of nanostructured zirconia compared to extended flat and stepped zirconia surfaces (for a more detailed description of this process, the reader is referred to ref\(^{20} \)). \( Zr_{80}O_{160} \), and \( Zr_{40}O_{80} \) split the \( H_2 \) molecule through a heterolytic mechanism, which implies the direct reduction of the oxide via formation of \( Zr^3+ \) centers (Figure 1b,d) and adsorbed protons. The heterolytic dissociation is energetically unfavorable compared to the homolytic process. On the contrary, \( Zr_{40}O_{80} \) splits \( H_2 \) heterolytically, as for the regular (101) surface, with no reduction of the oxide (Figure 1c). In this case, the homolytic process lies higher in energy. We attribute this difference to the different shape, and consequently to the different electronic structure, of the three NPs. In particular, the presence of low-lying \( Zr^4+ \) states in the energy gap of \( Zr_{80}O_{160} \) and \( Zr_{16}O_{32} \), owing to the low coordinated \( Zr \) ions at corners and edges (Figure 2b,d), favors the reduction of \( Zr^4+ \) to \( Zr^3+ \). By contrast, the energy gap of \( Zr_{40}O_{80} \) (Figure 2c) does not exhibit such defect states, in analogy with the (101) surface (Figure 2a).

The exothermicity of the hydrogenation on \( Zr_{80}O_{80} \) comes from the special structural flexibility that enhances its reactivity rather than from electronic effects due to low coordination of the \( Zr \) ions.

The entropic terms also make the \( H_2 \) dissociation less exergonic, resulting in Gibbs free energies of \(-1.19, -1.33, \) and \(-1.01 \) eV on \( Zr_{80}O_{160}, Zr_{40}O_{80} \) and \( Zr_{16}O_{32} \), respectively. Note that these values are qualitatively different from those corresponding to the (101) surface, where the reaction has a positive \( \Delta G \), Table 1. In addition, the OH and ZrH stretching frequencies from both the homolytic and heterolytic mechanisms are significantly blue-shifted in the NPs (\(-3700 \) cm\(^{-1} \)) compared to those on the regular (101) surface (\(-3000 \) cm\(^{-1} \)). This can be attributed to longer OH and ZrH bonds in the

Figure 1. Structures of hydrogenated zirconia: (a) \( ZrO_2 \) (101) surface; (b) \( Zr_{80}O_{160} \) (c), (d) \( Zr_{40}O_{80} \) and (d) \( Zr_{16}O_{32} \). Zr: blue atoms; O: red atoms; and H: white atoms. The reduction of two \( Zr^4+ \) to \( Zr^3+ \) in \( Zr_{80}O_{160} \) and \( Zr_{16}O_{32} \) is clearly seen from the spin density plots, indicated in yellow (\( \rho_{\text{spin}} = 0.01 \) e/\( \text{Å}^3 \)).
The second step implies the desorption of a water molecule from the hydrogenated surface with the subsequent formation of an O vacancy (dehydration). This process involves the diffusion of a H atom on the surface and its binding to an OH group with the formation of an OsH2 unit. In the (101) surface, this step has an energy cost of 3.55 eV. In this case, a slightly lower Gibbs free energy is obtained because the entropy of the H2O gas favors desorption (3.28 eV at 298 K and 1 bar, Table 1), but the process remains significantly endergonic. In the NPs, ΔGI is also endergonic but considerably smaller, about 1.5 eV for both Zr80O160 and Zr16O32 (Table 1); on the contrary, the cost remains high on Zr40O80 (ΔEI = 3.08 eV and ΔGI = 2.74 eV) where a heterolytic dissociation of hydrogen has occurred, Table 1. The reduction of the energy cost comes from the stabilization of the O vacancies in the NPs.

In the (101) surface, the creation of a vacancy costs around 6 eV with respect to the formation of 1/2O2, whereas in Zr80O160, Zr40O80, and Zr16O32, the cost is 2.77, 3.82, and 2.96 eV, respectively.16 It is important to note that the results show that the creation of an O vacancy in ZrO2 by desorption of H2O has a lower cost than via desorption of oxygen and formation of 1/2O2. This holds true for both NPs and surfaces.

From the thermodynamic data, we have calculated the reaction conditions at which the Gibbs free energy becomes negative (exergonic); hence, the process becomes thermodynamically spontaneous. For that, we have considered the entropy of the H2 and H2O gases at different temperatures between 50 and 1600 K and for pressures of 1 bar (ambient) and 10−12 bar (ultrahigh vacuum, UHV) according to eqs 5–67. Thus, a trend ΔG versus T is constructed for each pressure and reaction step. The temperatures at which the condition ΔG = 0 is satisfied are reported in Table 2. For the hydrogenation process, H2 dissociates spontaneously on the (101) surface at 1 bar and below 70 K because of the low reactivity of the extended zirconia toward H2 dissociation (for higher temperatures, H2 desorption dominates). On the contrary, the NPs can be hydrogenated at ambient pressure in a wide range of temperatures, up to 800−900 K (Table 2) because of the high reactivity toward H2 (exergonic adsorption free energies of −1/−1.3 eV, Table 1). Above this temperature range, the entropy...
contribution of H₂ prevents its adsorption to the surface and favors desorption. The desorption of water from the heterolytically dissociated 2H/ZrO₂ (101) surface at ambient pressure requires temperatures above 1500 K because of the high endergonic character of the process (ΔG²⁺ = +3.28 eV) or above 800 K when the partial pressure of water decreases to UHV conditions (10⁻¹² bar). As in the initial hydrogenation step, NPs can undergo dehydration at milder conditions compared to the regular surface. In particular, water can start desorbing from the surface, creating an O vacancy at temperatures above 450 K, a temperature at which several catalytic processes involving oxides start to occur. In summary, NPs of zirconia can dissociate hydrogen and desorb water at milder reaction conditions compared to the extended surface, changing completely the landscape for the chemistry of these systems. In the special case of zirconia, this leads a nonreducible oxide to become reducible when prepared in a nanostructured form.

3.2. Kinetic Aspects (Energy Barriers). In this section, we investigate the complete reaction energy path, from the adsorption of H₂ to the formation and desorption of H₂O, including the reaction barriers. This can be divided into two steps, as mentioned above. The first step involves the physisorption of H₂ followed by its dissociation (homo- or heterolytic); the second step is the diffusion of hydrogen on the surface and the formation of an OH₂ surface complex; and the last step is related to the desorption of H₂O from the surface, leaving behind an O vacancy. First, on the ZrO₂ (101) surface, the H₂ molecule is physisorbed with an energy of −0.11 eV; it dissociates heterolytically into H⁺ and H⁻ species through a small energy barrier of 0.28 eV and a small binding of −0.06 eV (see above). A similar barrier for the heterolytic dissociation was recently published by our group (0.24 eV). Experimentally, such a barrier has been estimated to be around 0.4 eV. To induce water desorption, the H⁻ ion must diffuse into the O−H⁺ group and form a O₂H₂ unit (this can also be seen as a H₂O molecule adsorbed on a surface O vacancy). This process forces the system to become reduced. In fact, when the two H atoms are bound to the same oxygen (O₃H₂ surface complex), the excess of charge is localized on two Zr³⁺ centers in a triplet state (Figure 3); it is the same configuration obtained when two H atoms are adsorbed on two different oxygen sites, resulting in a homolytic cleavage of the H−H bond (excited state). This state is energetically highly unfavorable, and in fact, this process is noncompetitive with respect to H₂ desorption. Still, it is a necessary step for water desorption. When the O₂H₂ surface complex is formed, two electrons are transferred to two Zr ions (Zr²⁺). This is different from the case of O-deficient zirconia (O vacancy) where electrons localize in the vacancy and give rise to a singlet ground state (Figure 3) because of the morphological change in the structure connected to the O removal. The formation of the O₂H₂ complex on the regular surface by H diffusion is thus highly endothermic, ΔE = +2.79 eV, and the barrier is obviously even higher, 2.93 eV. A lower barrier, 2.55 eV, has been reported in a previous study by our group using a slightly different computational approach. Once the O₂H₂ surface complex is formed, it desorbs with an energy cost of 0.72 eV. Therefore, the highest barrier in the entire process is the diffusion of a H atom to form a O₂H₂ precursor state of the desorbing water molecule; neglecting prefactors, this is the rate-determining step with a very high energy barrier, close to 3 eV. This process will hardly occur on the bare zirconia surface as H₂ desorption is clearly preferred.

Now, we consider zirconia NPs where lower energy barriers are expected for the formation and desorption of the H₂O molecule. In Figure 4, the reaction energy profiles for the dehydration step from the hydrogenated NPs are compared with those from the (101) surface. We did not study in detail the physisorption and the barrier for dissociation of the H₂.
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1, the small changes in H 2 physisorption and barriers for
where the barrier for a similar process has been estimated to be
above that on this NP the thermodynamics for the reduction
path for the dehydration of 2H/Zr40O80 has some resemblance
OsH2 precursor is lower (2.24 eV) because of the higher
energy barrier decreases from 2.93 eV on the extended surface
to 1.77 eV in Zr80O160 NPs and to 1.83 eV in Zr16O32 NPs. This
is a very significant reduction of more than 1 eV. It has the
important consequence that H diffusion competes with H2
desorption, differently from the bare surface. The subsequent
desorption of H2O has a cost of 0.85 eV in Zr80O160 and 1.11
eV in Zr16O32. Thus, also on the NPs, the highest barrier is not
associated with the water desorption but rather with the hydrogen
diffusion and with the formation of the precursor
state. Despite the large decrease in the barrier, this does not
reach the low value found for TiO2, a typical reducible oxide,
where the barrier for a similar process has been estimated to be
around 0.3 eV.33

Things are partly different for the Zr80O160 NP. We have seen
above that on this NP the thermodynamics for the reduction
reaction is similar to that of the (101) surface because of a
similar electronic structure with the absence of defective states
in the band gap (Figure 2c). Accordingly, the reaction energy
path for the dehydration of 2H/Zr80O160 has some resemblance
with that of the extended surface; the energy cost to form the
O=H2 precursor is lower (2.24 eV) because of the higher
structural flexibility of the nanostructure, and so is the barrier
(2.27 eV). From this state, water desorbs to the gas phase with
a cost of 0.84 eV. Thus, also in this case, the highest barrier is
associated with hydrogen diffusion. The barrier is considerably
lower than that on the regular surface (2.27 vs 2.93 eV) but still
much larger than those on the other two zirconia NPs
considered (about 1.8 eV) where the mechanism of H2
dissociation is different.

These results show the positive effect of nanostructuring on
the reducibility of zirconia by water removal. The special
fluidity of the NPs contributes to lowering not only the
energy costs but also the energy barriers involved in the
process, thanks to a larger atomic relaxation around the created
O vacancy. This structural flexibility is much smaller or even
absent in bulk ZrO2 and ZrO2 surface. However, comparing the
different behaviors of Zr80O160 and Zr16O32 NPs with that of
Zr80O160 NPs, one can conclude that the key factor is the presence
of low-lying acceptor states introduced in the electronic structure by low coordinated Zr sites in corners
and edges, which strongly depends on the morphology of the NP.

There is a final aspect that needs to be discussed and that,
one more, shows a different behavior for the extended zirconia
surface compared to that of the NPs. We are referring to the
process for H diffusion on the hydrogenated surface. In
the initial state, the two H atoms remain adsorbed on vicinal sites
(e.g., a Zr and an O ion, in the case of heterolytic splitting). On
the ZrO2 (101) surface, the displacement of H+ and H− ions
to more distant O and Zr sites has an energy cost of 0.64 and 0.73
eV, respectively, and it implies to overcome a barrier of 1.50
(H+) and 1.42 (H−) eV, Table 3. The fact that separating the

Table 3. Diffusion Energy, ΔE_diff and Energy Barrier, ΔE‡,
for the Diffusion of a H Atom into Adsorption Sites
Distant from the OH− Precursor for the H2O Formation

| support             | ΔE_diff (eV) | ΔE‡ (eV) |
|----------------------|-------------|----------|
| ZrO2 (101)           | OH− to O’   | 0.64     | 1.50     |
| ZrH+ to Zr’         |             | 0.73     | 1.42     |
| Zr80O160             | O2cH+ to O3c| 0.69     | 1.73     |
| ZrH+ to O3c         |             | 0.84     | 1.58     |
| ZrH+ to Zrfacet      |             | 0.24     | 2.15     |
| Zr16O32              | O2cH+ to O3c| 0.06     | 1.72     |
| ZrH+ to O2c         |             | 0.50     | 1.54     |

2 eV, not very di
fusion and water desorption become competitive processes.

4. CONCLUSIONS

We have performed, by means of DFT-based calculations with
the PBE + U exchange–correlation functional and D2′
dispersion, a comparative study of the reduction reaction
ZrO2 + H2 → ZrO2x−y + H2O in nanostructured zirconia and in
the regular (101) surface. This has been separated into two
steps: hydrogenation of the surface (step 1) and subsequent

DOI: 10.1021/acsomega.7b00799
ACS Omega 2017, 2, 3878−3885
removal of a water molecule with the formation of an O vacancy (step II).

Because of the low reducibility of ZrO$_2$, the reduction of the extended (101) surface is unfavorable, with energies $\Delta E_I = -0.06$ eV ($\Delta G_I = 0.34$ eV) and $\Delta E_H = 3.55$ eV ($\Delta G_H = 3.28$ eV). Zirconia NPs are more reactive, resulting in significantly lower reaction energies, $\Delta E_I = -1.6$ to $-1.8$ eV ($\Delta G_I = -1$ to $-1.2$ eV) and $\Delta E_H = 2-3$ eV ($\Delta G_H = 1.5-2.7$ eV), because of their special structural flexibility and electronic structure. In a real system, there will be a random distribution of Zr$^{3+}$ ions on the surface of the NPs. The disorder introduced by this distribution of sites will provide a configurational entropy contribution that is going to provide an extra stabilization for the reduced NPs compared to the bare surface.

However, the mechanism for H$_2$ splitting is determined by the electronic structure. On the (101) surface, H$_2$ is dissociated heterolytically (formation of OH$^+$ and ZrH$^+$ species). On Zr$_{50}$O$_{160}$ and Zr$_{16}$O$_{32}$ NPs, the presence of low-lying Zr 4d acceptor states from low coordinated Zr ions at corners and edges favors the homolytic H$_2$ splitting, which implies the direct reduction of the oxide (formation of two OH$^+$ species and two Zr$^{3+}$ centers). By contrast, such states are not present in Zr$_{80}$O$_{160}$ where the H$_2$ molecule dissociates heterolytically as on the (101) surface. This demonstrates that not only the dimensions of the NPs count but also their shape and surface morphology.

From the thermodynamic data, we conclude that the hydrogenation reaction at ambient pressure (1 bar) can be performed at any temperature up to 900 K on the NPs, whereas on the (101) surface, it is possible only below 70 K (the competitive mechanism is H$_2$ desorption).

For the dehydrogenation step, the energy barrier to form a H$_2$O molecule on the oxidized surface decreases from 2.93 eV on the (101) surface to 2.27 in Zr$_{50}$O$_{160}$ (which is more “bulklike”) and to 1.77 and 1.83 eV in Zr$_{50}$O$_{160}$ and Zr$_{16}$O$_{32}$, respectively. As a consequence, under UHV conditions, the dehydrogenation of the (101) surface requires temperatures above 800 K, whereas on the NPs, it is possible above 450 K, a typical reaction temperature in many catalytic processes. The results presented refer to the cases where only one H$_2$ molecule has been adsorbed on the surface of the zirconia NPs. However, under working conditions, the particle will be fully hydrogenated, and coverage effects will play a role. In particular, we expect lower barriers for hydrogen migration to form the O$_2$H$_2$ surface complex; also, the presence of excess electrons due to the high hydrogen coverage can facilitate water desorption. In general, this work shows the importance of nanostructuring for the chemistry of oxides as this completely modifies the oxide reducibility.

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