Properties of Single Oxygen Vacancies on a Realistic (TiO$_2$)$_{84}$ Nanoparticle: A Challenge for Density Functionals

Ángel Morales-García, Oriol Lamiel-García, Rosendo Valero, Francesc Illas*

Departament de Ciència de Materials i Química Física & Institut de Química Teòrica i Computacional (IQTCUB), Universitat de Barcelona, c/ Martí i Franquès 1, 08028, Barcelona, Spain.

Abstract

Based on all electron relativistic density functional theory (DFT) calculations, the properties of single oxygen vacancies in TiO$_2$ nanoparticles have been obtained using a suitable representative model consisting of an octahedral (TiO$_2$)$_{84}$ nanoparticle of ~3 nm size terminated with (101) facets. This nanoparticle can be safely considered at the onset of the so-called scalable regime where properties scale linearly with size towards bulk like limit and hence results can be more directly compared to experiment. A set of reduced Ti$_{84}$O$_{167}$ nanoparticles are selected to investigate the geometrical, energetic and electronical properties by using PBE semi-local functional with three different amounts of Fock exchange 0% (PBE), 12.5% (PBEx) and 25% (PBE0). In particular, using the PBEx hybrid functional, previously validated for bulk anatase and rutile, it is predicted that the highly (three)-coordinated oxygen atom, located in the subsurface, and the least coordinated one at top sites are energetically the most suitable candidate for generating the oxygen vacancy. The subsurface case is in line with conclusions from experiments carried out on (101) single crystal anatase surfaces. The electronic structure of the reduced particles suggests that these would have better photocatalytic activity than their stoichiometric counterparts. Nevertheless, several properties of reduced TiO$_2$ NPs are strongly affected by the choice of the exchange-correlation functional implying that, in absence of validation by comparison to experiment, predictions must be taken with caution.
Introduction

Titanium dioxide (TiO$_2$), a semiconducting metal oxide, is widely used as a photocatalyst because of its chemical stability, low cost, and nontoxicity.$^{1-4}$ Both rutile and, especially, anatase nanostructures catalyze the overall splitting of water into molecular hydrogen and oxygen when illuminated with ultraviolet (UV) light.$^{5-8}$ This photocatalytic process represents a promising clean and sustainable alternative to fossil fuels.$^{9,10}$ However, applications of TiO$_2$ show a low quantum yield (QY) and limited harvesting of visible light which is a direct consequence of the too large band gap of this material. Moreover, fast recombination of photogenerated electron-hole pairs represents an additional problem. To tackle the above obstacles, several strategies have been applied to modify TiO$_2$, like depositing noble metals (e.g. Pt),$^{11}$ doping with transition-metal ions (e.g. Cu) or nonmetal elements (e.g. N),$^{12}$ adding electron carriers (e.g. graphene)$^{13}$ or coupling to small band gap quantum dots (e.g. PdS).$^{14}$ In addition, TiO$_2$ nanostructuring with specific morphologies and crystal facets has emerged as a promising way to improve the QY with visible light.$^{15,16}$

Alternatively, engineering defects on the TiO$_2$ nanostructures appears as another encouraging way to circumvent the aforementioned low QY of TiO$_2$.$^{17}$ It has been reported that creating a highly disordered surface layer and a large amount of oxygen vacancies (O$_v$) in TiO$_2$ not only improves visible and near-infrared light absorption but also leads to a superior activity in photocatalytic degradation of organic pollutants and H$_2$ evolution from H$_2$O.$^{18}$ From simple electron counting and assuming formal oxidation states, one neutral oxygen vacancy, O$_v$, provides two excess electrons to the TiO$_2$ system, and these electrons are available for the reduction of Ti$^{4+}$ to Ti$^{3+}$. Indeed, the formation of Ti$^{3+}$ center has been confirmed by experiments such as photoelectron spectroscopy and electron paramagnetic resonance (EPR).$^{19,20}$ Recently, it has been shown that Ti$^{3+}$/O$_v$ self-doped TiO$_2$ nanocrystals with a 1:80 Ti$^{3+}$:Ti$^{4+}$ ratio exhibit a remarkable enhancement of the photocatalytic efficiency; this is precisely ascribed to the presence of Ti$^{3+}$ centers and concomitant O$_v$ point defects.$^{21}$ The combination of O$_v$ and Ti$^{3+}$ centers leads to specific electronic structure features usually referred to as trapped electrons. These states appear around 1.0 eV below the conduction band minimum (CBM), which induces band gap narrowing. Engineering Ti$^{3+}$/O$_v$ sites can favorably promote CO$_2$ activation and conversion to CO under visible light.$^{22}$ Several computational studies have focused on these systems and reported the properties of O$_v$ in bulk, surface and subsurface TiO$_2$ sites.$^{23-25}$ Experiments and calculations show that formation energy of neutral oxygen vacancy is lower when defect site is located in the subsurface of anatase (101) and (001) surfaces, whereas the surface site is the most stable site in rutile (110) surface. Similar
information on TiO$_2$ nanostructures is so far lacking except for a recent study on a (TiO$_2$)$_{35}$ representation of an octahedral nanoparticle.$^{26}$ This is indeed the smallest TiO$_2$ NP compatible with Wulff construction, yet metastable with respect to non-crystalline structures$^{27}$ as further commented below.

The electronic properties of TiO$_2$ nanoparticles (NPs) have been discussed in the light of experiments reporting that the control of the shape and size of titania NPs modulates the photoactivity.$^{15}$ The understanding of TiO$_2$ NP properties is particularly attractive because its size and shape are appropriate to rationalize the experimental evidences.$^{28}$ However, experimentally, it is difficult to discern between the different effects of size and shape and those introduced by the synthetic conditions. The difficulties encountered by experiments to separate complex factors are not present when employing computational models in which one can represent different morphologies for a given composition or vary the composition for a given morphology.$^{27,29-31}$ For instance, the analysis of the atomic structure of TiO$_2$ small clusters and NPs of increasing size allows one to estimate the dimensions for emergence of crystallinity and scrutiny of electronic properties of well defined nanoparticles allows one to identify the size-limit of titania NP at which the properties show an asymptotic trend towards bulk-like behavior.$^{30}$ In particular, it has been shown that the realistic (TiO$_2$)$_{84}$ anatase NP is at the start of the scalable regime limit above which properties of titania NPs scale linearly towards bulk limit values. Assuming spherical shape$^{32}$ one would get an estimate size of 3 nm for the octahedral (TiO$_2$)$_{84}$ NP, a value which nicely fits with the predicted height from a linear relationship between the largest vertex to vertex distance in octahedral TiO$_2$ NPs and the cubic root of the number of TiO$_2$ units.$^{30}$ From both arguments one can conclude that (TiO$_2$)$_{84}$ anatase NP is an appropriate model system to investigate the changes induced by the generation of O$_v$ in their electronical and structural properties and hence in its photocatalytic activity.

Herein, using state of the art density functional theory (DFT) based methods, we systematically investigate the properties of oxygen vacancies in an initially stoichiometric (TiO$_2$)$_{84}$ anatase NP. To facilitate the study and handle the numerous possible isomers resulting from the creation of a single O$_v$ in this quite large NP, we follow the strategy developed in a preliminary study on a smaller (TiO$_2$)$_{35}$ anatase NP.$^{26}$ We have already mentioned that this is the smallest size for a bulk cut octahedral NP exhibiting (101) facets although in terms of total energy per TiO$_2$ unit it is clearly metastable.$^{27}$ This fact can affect the conclusions regarding reduction and ask for subsequent studies on a more realistic model. This is precisely the main reason to undertake the present work on the properties of oxygen vacancies on the (TiO$_2$)$_{84}$ NP. Several possible sites for
oxygen vacancy formation are evaluated according to the position and the coordination number (CN), using results for oxygen vacancy in (TiO$_2$)$_{35}$ as a convenient guide.$^{26}$

**Computational Details**

First of all, it must be recalled that standard density functionals within the Generalized Gradient Approach (GGA) fail to provide even a qualitatively correct description of electronic properties in stoichiometric and reduced TiO$_2$ systems due to an exceedingly large effect of the so-called self-interaction error inherent to the Kohn-Sham practical implementation of DFT.$^{33,34}$ To correctly describe the partial occupation and appropriate localization of states arising from Ti$^{3+}$ sites hybrid functionals are required where a given percent of Fock exchange is mixed into a given exchange GGA functional. Assuming that the Kohn-Sham energy levels provide an estimate of the binding energies of quasiparticles, previous studies have shown that modifying the PBE0 (25\% Fock)$^{35,36}$ functional so as to contain 12.5\% of Fock exchange (PBEx) reproduces successfully experimental band gap of anatase and rutile polymorphs whereas the commonly used PBE0 hybrid functional$^{35}$ significantly overestimates them.$^{23}$ In addition, PBEx functional describes properly the properties of the O$_v$ in both rutile and anatase TiO$_2$ bulk phases.$^{23}$

In the present study DFT based calculations have been carried with the PBEx hybrid functionals. However, in the view of the strong influence of the exchange-correlation functional on the outcome of the calculation, results from more traditional approaches such as PBE$^{36}$ and PBE0$^{35}$ are included for comparison. Another aspect regarding the accuracy of the density functional methods chosen concerns the contribution of dispersion terms. For the atomic structure of bulk anatase and rutile, Deringer and Scányi$^{37}$ found rather small changes in the predicted lattice parameters when taking into account these effects on top of results obtained from PBE. The changes are similar those encountered when going from PBE to PBEx. Consequently, the contribution of dispersion has not been further considered.

All calculations explicitly include all electrons and the electron density is described through a numerical atom-centered (NAO) orbital basis set, as implemented in the Fritz Haber Institute *ab initio* molecular simulations (FHI-aims) package.$^{38}$ The light grid and tier-1 basis set are selected, the numerical accuracy of this basis set for TiO$_2$ systems is similar to that of a valence triple-$\zeta$ plus polarization Gaussian type orbitals (GTO) basis.$^{30}$ Recent studies have further assessed the quality of the light tier-1 NAO basis set on the stability and electronic properties of TiO$_2$ NPs.$^{31}$ The convergence threshold for atomic forces in relaxation of pristine and reduced (TiO$_2$)$_{84}$ NPs is set to $10^{-2}$ eV Å$^{-1}$. The presence of a transition element like Ti requires the inclusion of relativistic effects
to properly ensure sufficient accuracy for calculated properties. In the present work, relativistic effects are included through the zero order regular approximation (ZORA).\textsuperscript{39,40}

A series of reduced Ti\textsubscript{84}O\textsubscript{167} NPs has been considered which differ in the position where the oxygen vacancy has been created. The considered NPs have all formally two Ti\textsuperscript{3+} centers and hence involve two excess electrons leading to different electronic states—closed shell singlet, open shell singlet and triplet—depending on the spin coupling. Hereby, closed-shell singlet and open-shell triplet states are considered as in previous work.\textsuperscript{26} The oxygen vacancy formation energy, $E_f^0$, is calculated as:

$$E_f^0 = E(Ti_{84}O_{167}) + \frac{1}{2}E(O_2) - E(Ti_{84}O_{168})$$  \hspace{1cm} (1)

where, $E(O_2)$ stands for the energy of O\textsubscript{2} molecule in its triplet ground state and $E(Ti_{84}O_{167})$ and $E(Ti_{84}O_{168})$ correspond to the total energies of a given reduced Ti\textsubscript{84}O\textsubscript{167} NP and of the pristine (TiO\textsubscript{2})\textsubscript{84} one. In all cases, the structure of the NP is fully relaxed for each one of the functionals employed in this study. According to definition of Eq. (1), a positive value of $E_f^0$ is associated with the extra energy required for removing a neutral single oxygen. We also define the vertical energy ($E_v$) which is calculated as in Eq. (1) but using the energy of the non-relaxed reduced Ti\textsubscript{84}O\textsubscript{167} NP and the relaxation energy ($E_{rel}$) defined as $E_{rel} = E_v - E_f^0$. Further details about the definition of energies are schematically given in Supporting Information.

**Results and Discussion**

It is necessary to insist on the fact that the contribution of non-local, exact, Fock exchange to the exchange-correlation potential has a marked influence in predicted properties of oxides in general and of TiO\textsubscript{2} related systems, in particular. Precisely, this is an especially delicate issue for the properties of O\textsubscript{v} containing systems exhibiting unpaired electrons strongly localized at Ti\textsuperscript{3+} centers but with the degree of localization depending on the choice of the exchange-correlation potential. To rigorously tackle this problem and to avoid any particular bias, this section is divided into two parts. In the first one, properties of O\textsubscript{v} on reduced Ti\textsubscript{84}O\textsubscript{167} NPs are discussed in light of results obtained with the PBE\textsubscript{Ex} functional which, as pointed out above, properly describes stoichiometric and reduced bulk TiO\textsubscript{2}. In the second part the influence of the Fock exchange contribution in the exchange-correlation potential is analyzed in some details so as to reach unbiased conclusions regarding the interpretation on vacancy defect states in the Ti\textsubscript{84}O\textsubscript{167} NPs. To this end, results have been obtained from the PBE and PBE\textsubscript{0} functionals. Results from the later are explicitly discussed here whereas the former are reported in the Supporting Information.
Properties of \( \text{O}_v @ \text{Ti}_{84}\text{O}_{168} \) as described from the PBEx functional

The optimized structure of the realistic \((\text{TiO}_2)_{84}\) NP obtained from a Wulff construction\(^{41,42}\) and featuring the most stable (101) facets\(^{43}\) is shown in Figure 1. To systematically examine the effect of a single oxygen vacancy one would need to consider all possibilities which, even exploiting symmetry, faces a combinatorial explosion. Therefore, we rely on results obtained from an earlier systematic study on the smaller \((\text{TiO}_2)_{35}\) NP\(^{26}\) and choose the most favorable sites for \(\text{O}_v\) as candidates for \((\text{TiO}_2)_{84}\) NP. This is justified because both NPs have the same octahedral shape, symmetry and exposed (101) facets.\(^{26,28}\) Figure 1 displays the fully optimized atomic structure of the stoichiometric \((\text{TiO}_2)_{84}\) and of different reduced \(\text{Ti}_{84}\text{O}_{167}\) NPs that are obtained after removal of a selected single oxygen atom. The vacancy sites considered are classified according to the following notation: top (T), edge (E), facet (F), and inside (I) types which clearly define the position of the removed oxygen. To further identify each single \(\text{O}\) vacancy, we use the \(X\)-m notation where \(X\) and \(m\) designate domain type (T, E, F or I) and CN (1, 2 or 3) respectively. For instance, F-3 corresponds to the removing an oxygen atom coordinated to three Ti atoms (CN = 3) in the facet (F) domain. Note also that an additional digit (1 or 2) is used to distinguish two I-3 analog sites.

We start analyzing two different optimized structures of the pristine \((\text{TiO}_2)_{84}\) NP, which just depend on the orientation of apical oxygen atoms: \((i)\) almost linear, straight tip, or \((ii)\) bent to opposite sides, bent tip (see Figure 3 in Ref. [30]). The bent tip structure depicted in Fig. 1a is energetically more stable by around 0.011 eV per \(\text{TiO}_2\) unit (0.96 eV in absolute terms) for all employed DFT functionals, consistent with the previous analysis over \((\text{TiO}_2)_{35}\) NP where a similar favorable energetic stability for bent tip structure was found (1.06 eV in absolute terms).\(^{26}\) We assume that the difference between straight and bent tip structures tends to be negligible by increasing the size of \(\text{TiO}_2\) NP although this is not an important issue. In fact, despite their slight structural difference, the electronic band gap \(E_{\text{gap}}\), evaluated as the difference between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) is almost the same. This value, however, strongly depends on the exchange-correlation potential used to describe the electronic structure and PBEx functional nicely reproduces\(^{23}\) the experimental \(E_{\text{gap}}\) of 3.20 eV of bulk anatase phase.\(^{44}\) For the stoichiometric \((\text{TiO}_2)_{84}\) NP, the PBEx functional calculated \(E_{\text{gap}}\) is 3.49 and 3.61 eV for straight and bent tip structures, respectively. Here, it is noted that for small nanoparticles \(E_{\text{gap}}\) depends significantly on the geometrical structure\(^{29}\) although the effect fades away for larger particles in the scalable regime.\(^{30}\)

To avoid any possible artifacts, the properties of \(\text{O}_v\) in \((\text{TiO}_2)_{84}\) NP are studied by removal of one neutral oxygen atom from the bent tip structure (Figure 1). Six oxygen sites are selected and
systematically removed one at a time. This set of Ti$_{84}$O$_{167}$ reduced nanoparticles represents the structures that have the lowest $E_f^O$ in each one of the domains based on our previous study of (TiO$_2$)$_{35}$ NP$^{26}$ having the same morphology and exhibiting the same facets as commented above. Table 1 reports the values of $E_v$, $E_{rel}$, $E_f^O$ and the energy level of the oxygen vacancy defect state below LUMO ($S_{v-1}$ and $S_{v-2}$) calculated at the PBEx level. Note that the oxygen vacancy sites with just a $S_{v-1}$ energy correspond to a closed-shell singlet state which is found to be the electronic ground state after geometry optimization. This is quite an unexpected result since spin polarized solutions are the ground state for O$_v$ in bulk anatase and comes from the differences in structural relaxation that are possible in the NP as compared to the bulk or extended surfaces. In fact, the rather broad range of $E_f^O$ values for the different sites appears to be due to differences in the structural relaxation of the reduced Ti$_{84}$O$_{167}$ NPs. The vacancy formation is largely determined by the structural features around the oxygen vacancy site. For the PBEx functional, $E_f^O$ appears in the 3.56–4.27 eV range, a prediction fully consistent with those previously reported using the same PBEx functional for anatase bulk phase, with $E_f^O$ values in the 4.21–4.65 eV range, and for the (TiO$_2$)$_{35}$ NP, although in this case $E_f^O$ involves a broader range (2.27–5.15 eV).$^{23,26}$

Figure 2 displays trends for $E_f^O$ in the Ti$_{84}$O$_{167}$ nanoparticle, I-3-2 site has the lowest $E_f^O$ value of 3.56 eV (see Table 1). Note, however, that the T-1 domain is energetically competitive showing an energy difference of 0.02 eV only. Another highlighted energetic parameter is the vertical energy $E_v$ (Table 1). It is observed that O$_v$ formation at I-3-2 site features the lowest vertical energy value (4.29 eV) whereas the largest one (5.26 eV) corresponds to the formation of the unrelaxed O$_v$ at F-2 site. On the other hand, for the I-3-2 and F-2 site, $E_{rel}$ values of ~0.72 and ~1.31 eV, respectively are obtained. Note that, except for the F-2 site, $E_{rel}$ varies in a narrow range of 0.7–0.9 eV so that, within 0.2 eV, the stability of the different possible single oxygen vacancies is hence dominated by the $E_v$ energy. This energetic analysis confirms that the O$_v$ formation is preferentially located inside the nanoparticle in the subsurface region instead of directly at surface terraces.

According to experimental evidences, surface oxygen vacancies are not present on freshly cleaved anatase (101) samples.$^{45,46}$ A reduced anatase (101) crystal shows isolated as well as ordered intrinsic subsurface defects in scanning tunneling microscopy (STM), consistent with DFT calculations for surface slab models which predict that O$_v$ at subsurface and bulk sites are significantly more stable than on the surface.$^{45,46}$ Note that O$_v$ at T-1 site is essentially as stable as at I-3-2 site but, it is a feature that (i) is not present in extended crystalline (101) anatase surfaces and (ii) unique to the morphology of the nanoparticle, thus it would not be found in extended surfaces.
The prediction from the present all electron relativistic DFT based calculations for realistic TiO$_2$ NP are in agreement with the experimental evidence and provides further support to the claim that (TiO$_2$)$_{84}$ corresponds to a particle size in the scalable regime and hence representative of the large NPs used in the experiments. This provides additional support to the choice of the PBEx hybrid functional as appropriate to investigate the properties of stoichiometric and reduced realistic TiO$_2$ NP as suggested from previous studies.$^{23,26,31}$

We discuss now in some detail the structural changes around the first neighbor titanium atoms to each of the investigated O$_v$ sites (Figure 3). The reorganization of the structure around the vacant space and the presence of the Ti$^{3+}$ sites arising from the two excess electrons are the major factors in the relaxation of the reduced Ti$_{84}$O$_{167}$ NPs. To understand the quantitative analysis of $E_f^D$ (Table 1), we carefully analyze the geometrical differences between pristine and reduced (TiO$_2$)$_{84}$ anatase NPs for each one of the O$_v$. Oxygen vacancy at the T-1 and I-3-2 sites show the lowest $E_f^D$, 3.56 and 3.58 eV respectively, that is consistent with the negligible variation of Ti-O distance once the oxygen vacancy is created (Figures 3a, 3e). Contrarily, O$_v$ at the I-3-1 site (Figure 1a) shows noticeable structural changes requiring 0.47 eV more than the O$_v$ at the I-3-2 site although this energy difference is already present in the values of $E_v$. Accordingly, this extra energy requirement cannot be attributed to the noted changes in the structural properties (Figure 3a). Assuming these arguments, the most favorable domains to accommodate O$_v$ are mainly dictated by the initial structure with the atomic reorganization around the vacant space playing a minor role. Considering both effects, the ranking of site for O$_v$ ordered from the lowest to the highest oxygen formation energy is I-3-2 ~ T-1 < F-2 < E-2 < I-3-1 < F-3.

Finally, we consider the effect of the presence of Ti$^{3+}$ ions in reduced TiO$_2$ NP in the resulting electronic structure. The existence of Ti$^{3+}$ can be regarded as one type of n-type doping of TiO$_2$ samples. From a theoretical point of view, the description of Ti$^{3+}$ centers in TiO$_2$ systems is not straightforward since PBE predicts physically meaningless fully delocalized whereas hybrid functionals predict a variety of near degenerate solutions with different degrees of localization as described in the literature.$^{47}$ Nevertheless, upon O$_v$ formation a band gap narrowing is observed in all cases.$^{19,21,26}$ In bulk samples, experiments show that the defect state appears at around 1 eV below CBM, a feature which is reproduced by the PBEx functional.$^{22}$ Figure 4 presents the PBEx molecular orbital energy level diagram of Ti$_{84}$O$_{167}$ NP for a total of six O$_v$ sites. Note that here all energy (quasi band) levels are referenced with respect to the vacuum.$^{48}$ The defect states are reported as singly (triplet state) or doubly (singlet state) occupied molecular orbitals. Only spin-up eigenvalues are shown because molecular energy levels of spin-up and spin-down singlet and triplet
states are similar. The spin-unrestricted triplet state is competitive with the closed-shell singlet one in reduced TiO$_2$ NPs, a feature which is specific of the reduced NPs and does not appear in the bulk anatase. It is clearly shown that the defect states appear between HOMO and LUMO and that, not surprisingly, the $E_{\text{gap}}$ and $S_v$ values vary with O$_v$ site. The presence of the O$_v$ defect also affects the position of the HOMO and LUMO energy levels so that new electronic transitions both from HOMO and defect states to LUMO and from HOMO to defect states become possible.

**Effect of the amount of Fock exchange on the Properties of O$_v$@Ti$_{84}$O$_{168}$**

We discuss now in detail the effect of percentage of Fock contribution on vacancy defect states comparing PBEx and PBE0 results. The pure GGA type PBE functional has also been considered for completeness although given the shortcomings of this approach in describing the electronic structure of oxides, the corresponding results are reported in the Supporting Information.

We start by describing the effect of the functional on the calculated values of $E_v$, $E_{\text{rel}}$ and $E_f^0$. The contribution of Fock exchange has a relatively low influence on $E_v$ as one can readily see by comparing PBEx and PBE0 results in Tables 1 and 2. The largest $E_v$ difference between these two hybrid functionals is of 0.7 eV and corresponds to the F-3 and I-3-2 domains. On the other hand, significantly larger differences are encountered for $E_{\text{rel}}$ ranging now between 2.44 and 3.34 eV which make the $E_f^0$ values calculated by the PBE0 functional significantly smaller than those obtained from calculations using PBEx functional. The fact that PBE0 functional with a larger (25%) contribution of the Fock exchange involves a larger contribution to the energy gain upon relaxation of the reduced Ti$_{84}$O$_{167}$ NP is consistent with a larger degree of localization of trapped electrons in the Ti$^{3+}$ centers with a concomitant larger structural relaxation; for instance, the F-3 domain increases in 0.7 and 2.18 eV for $E_v$ and $E_{\text{rel}}$, respectively. This is indeed the main energetic term that leads to $E_f^0$ below 3.0 eV (see Figure 2 and Table 2). Although PBE0 calculated $E_f^0$ values are smaller than those predicted from PBEx, the stability of the different possible single oxygen vacancies continues to be governed by the $E_v$ quantity following the same argument explained at PBEx level earlier. The PBE0 calculations predict that the most preferential site to form the O$_v$ is located at tip (T-1) and facet domain (F-2). The T-1 case is not so surprising as this is also the preferred site for O$_v$ in ceria nanoparticles,$^{49,50}$ the larger localization induces a larger relaxation and the final result is perhaps as expected, the lowest coordinated O atom is the easiest to remove. This is also in agreement with the results obtained with the PBEx functionals which predicts that O$_v$ is almost equally favorable at T-1 and I-3-2 sites. The F-2 case is harder to understand and it is not consistent with the experimental findings for extended surfaces perhaps due
to its proximity to the nanoparticle edge (see Figure 1). In any case, it is important to point out that the differences in $E_P^0$ for the different most stable sites are of the order of 0.3 eV only and this is perhaps in the error bar of this type of DFT based calculations for this property. Nevertheless, the final $E_P^0$ calculated values are much smaller with PBE0 functional implying that TiO$_2$ appears to be more reducible than predicted with PBEx. Unfortunately, without a guide from experiment it is hard to assess which of the two functionals better describes TiO$_2$ reduced nanoparticles and, in the view of the better performance of PBEx functional for stoichiometric and reduced rutile and anatase one is tempted to suggest that this will be also the case for the reduced NPs even if some feature is counterintuitive.

Although the experimental indications are not completely corresponding with predictions from the PBE0 functional, this approach reproduces the degree of localization on the neighboring Ti atoms, which are reduced to Ti$^{3+}$ in the I-3-2 site, which is not always observed by using PBEx functional (see Figure 4). Note that whereas PBEx functional gives the most favorable site at I-3-2 site consistent with the experiments for extended anatase (101) surfaces, it does not appropriately reproduce the oxygen vacancy defect state below LUMO (Figure 4) promoting a doubly (singlet state) occupied molecular orbital. This issue is solved when the percentage of Fock contribution increases (PBE0 functional) giving singly (triplet state) occupied molecular orbitals as shown in Figure 5.

Therefore, one must conclude that the appropriate selection of the exchange-correlation functional to investigate the properties of reduced TiO$_2$ NPs requires a thorough examination of a range of properties and to analyze how the amount of Fock exchange affects them. In the case of O$_v$ in TiO$_2$ NPs there is a certain degree of incertitude on determining exactly the most stable one although one can for sure make more reliable predictions by focusing on a range of formation energies including various possibilities. This is supported by analysis of Figure 2 clearly showing that PBEx and PBE0 functionals exhibit similar trends regarding the energy formation of different O$_v$ and that this is clearly different from the trend predicted by PBE functional. Also, describing properly the stability and the electronic structure of O$_v$ related states is far from being a solved problem since the amount of Fock exchange required to accurately describe the gap is insufficient to produce the expected localization. The latter remains, nevertheless, to be confirmed by experiments on suitable nanoparticles.
Conclusions

The properties of neutral single oxygen vacancy in realistic anatase nanoparticles have been investigated by choosing a suitable model and making use of all electron relativistic DFT based calculations within the PBEx hybrid functional previously optimized to reproduce the structure and electronic properties of bulk, stoichiometric and reduced, anatase and rutile. The model chosen is a \((\text{TiO}_2)_{84}\) anatase nanoparticle of \(~3\) nm size terminated with (101) facets which is at the onset of scalable regime and a total of six different sites have been systematically investigated to analyze the structural and electronic properties of various isomers of the reduced \(\text{Ti}_{84}\text{O}_{167}\) nanoparticle.

The oxygen vacancy formation and the energy levels of the defect state are quite dependent on the site from which the oxygen atom is removed. From energy arguments, the most favorable oxygen vacancy is the three-coordinated domain located at the subsurface region, being in nice accordance with experimental evidences observed in anatase well defined surfaces. The latter is an additional argument in support of the claim that this particle size is well in the scalable regime and can be taken as representative of larger nanoparticles for future studies involving for instance hydroxylation and/or doping. Nevertheless, the formation of a vacancy produced by removing the least coordinated O atom is almost as favorable and becomes the most favorable one according to calculations with the PBE0 functional.

The present calculations predict a HOMO-LUMO gap for the pristine nanoparticle of 3.61 eV and the presence of defect states in the reduced nanoparticle in the 0.49–1.68 eV range below the LUMO energy level, which are mostly due to the presence of \(\text{Ti}^{3+}\) cations. In some cases, the PBEx electronic ground state of the reduced nanoparticles appears to be a closed singlet, even optimizing structures with atomic displacements around \(\text{O}\) (see I-3-2 site in Figures 3 and 4) which is at variance with the behavior found for bulk anatase and its extended surfaces. Here, PBE0 functional predicts that the electronic ground state is a triplet with two singly occupied orbitals located in two \(\text{Ti}^{3+}\) cations more in line with expectations from chemical intuition and with experiments in bulk and extended surfaces. Therefore, although we rely mostly on the conclusions based on the results obtained from calculations with the PBEx hybrid functional, one must admit that the influence of the functional is large and, in some cases, the site predicted to lead to the most stable reduced nanoparticle and the character of the oxygen vacancy containing nanoparticles electronic ground state can vary with the amount of Fock. The fact that PBEx functional reproduces most of the quantities related to stoichiometric and reduced anatase and rutile provides support to the present conclusions for the reduced nanoparticles but also points out that the choice of the DFT based method is far from being a settled problem. The positive part is, however, that many
qualitative trends are less dependent on the functional. The overall reduction of the band gap of the reduced nanoparticle together with the presence of defect states suggest that reduced nanoparticles would be better suited to harvest light in the visible. However, the standard redox analysis does not clearly confirm that reduced TiO$_{2-x}$ NPs can be applied to visible light absorbing photocatalyst for the water splitting photocatalytic process.

Although the realistic TiO$_2$ NP studied here seems appropriate to unravel the features of reduced nanoparticles, further studies are required to shed light on the photoreactivity of reduced TiO$_2$ NPs towards water splitting considering for instance multi-oxygen vacancy effects, migration of oxygen vacancies and solvation effects. Note that the present study may be extended to investigate reduced photocatalysts which show ability to directly utilize solar energy for producing solar fuels such as H$_2$, CH$_4$ or CH$_3$OH.$^{51}$

ASSOCIATED CONTENT
Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI:
The description of energetic scheme is shown in Figure S1. Oxygen vacancy formation energies, Kohn-Sham orbital energy level diagrams and structural properties are described in Table S1 and Figures S2 and S3 for PBE level. Whereas, structural properties at PBE0 level are depicted at Figure S4.

AUTHOR INFORMATION

Corresponding Author
* E-mail: francesc.illas@ub.edu

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Table 1. Vertical energy ($E_v$), relaxation energy ($E_{rel}$), oxygen vacancy formation energy ($E_f^O$) and energy level of oxygen vacancy defect state below LUMO (S$_{v-1}$ and S$_{v-2}$) as predicted from DFT based calculations with the PBEx hybrid functional. All values are in eV units and the lowest $E_f^O$ values are highlighted in bold type. “CN” is number of first neighbor titanium atoms to remove oxygen atom and “site” is the combination of position and CN number.

| Position | CN | Site | $E_v$ | $E_{rel}$ | $E_f^O$ | S$_{v-1}$ | S$_{v-2}$ |
|----------|----|------|-------|-----------|--------|-----------|-----------|
| Tip      | 1  | T-1  | 4.48  | 0.89      | 3.58   | 0.64      | 0.49      |
| Edge     | 2  | E-2  | 4.71  | 0.70      | 4.01   | 1.68      | 0.53      |
| Facet    | 2  | F-2  | 5.26  | 1.31      | 3.95   | 1.15      | 0.69      |
|          | 3  | F-3  | 5.02  | 0.75      | 4.27   | 0.56      | 0.53      |
| Inside   | 3  | I-3-1| 4.73  | 0.69      | 4.03   | 1.63      | -         |
|          | 3  | I-3-2| 4.29  | 0.72      | 3.56   | 0.37      | -         |
Table 2. Vertical energy ($E_v$), relaxation energy ($E_{rel}$), oxygen vacancy formation energy ($E_f^O$) and energy level of oxygen vacancy defect state below LUMO ($S_{v-1}$ and $S_{v-2}$) as predicted from DFT based calculations with the PBE0 hybrid functional. All values are in eV units and the lowest $E_f^O$ values are highlighted in bold type. “CN” is number of first neighbor titanium atoms to removed oxygen atom and “site” is the combination of position and CN number.

| Position | CN | Site  | $E_v$ | $E_{rel}$ | $E_f^O$ | $S_{v-1}$ | $S_{v-2}$ |
|----------|----|-------|-------|-----------|---------|-----------|-----------|
| Tip      | 1  | T-1   | 4.35  | 2.86      | 1.48    | 0.51      | 0.28      |
| Edge     | 2  | E-2   | 4.87  | 2.88      | 1.99    | 2.94      | 2.43      |
| Facet    | 2  | F-2   | 5.27  | 3.34      | 1.92    | 2.69      | 2.21      |
|          | 3  | F-3   | 5.72  | 2.93      | 2.78    | 1.88      | 1.70      |
| Inside   | 3  | I-3-1 | 4.73  | 2.44      | 2.28    | 2.60      | -         |
|          | 3  | I-3-2 | 4.99  | 2.80      | 2.19    | 1.74      | 1.57      |
Figure 1. (a) Side and top views of the pristine (TiO$_2$)$_{84}$ NP. (b) Side and top views of the different reduced Ti$_{84}$O$_{167}$ NPs. Blue, red and black spheres denote titanium, oxygen and first neighbor titanium atoms to the O$_v$ site, respectively. Note that the reduced NP labeled as I-3-2 is slightly tilted to show clearly the region of O$_v$ site in the subsurface.
Figure 2. Oxygen vacancy formation energy ($E_f^O$) for single oxygen vacant Ti$_{84}$O$_{167}$ nanoparticle at PBE, PBEx and PBE0 levels depicted with black, red and blue navy colors, respectively. Note that dashed gray line indicates that the oxygen vacancy in I-3-2 site is the most favorable domain to create the oxygen vacancy at PBEx level. $E_f^O$ values for PBE, PBEx and PBE0 functionals are compiled in Tables S1, 1 and 2, respectively.
Figure 3. Stoichiometric and reduced $(\text{TiO}_2)_{84}$ NP motifs that correspond to the first neighbor titanium atoms to the O$_v$ site depicted in black at Figure 1: (a) T-1 and I-3-1, (b) E-2, (c) F-3, (d) F-2 and (e) I-3-2 and calculated at PBEx level. Blue and red spheres denote titanium and oxygen atoms, respectively.
Figure 4. Kohn-Sham orbital energy level diagram of Ti$_{84}$O$_{168}$ and reduced Ti$_{84}$O$_{167}$ NPs at obtained from calculations with the PBEx functional. The triplet state energy levels are for spin-up eigenvalues. The red dotted lines correspond to the standard redox potentials for water splitting at pH = 0 (H$^+$/H$_2$ = -4.44 eV and O$_2$/H$_2$O = -5.67 eV). The singly and doubly occupied defect states are denoted in blue and green for triplet and singlet states, respectively. Other occupied and unoccupied molecular orbitals are represented with black lines.
**Figure 5.** Kohn-Sham orbital energy level diagram of Ti$_{84}$O$_{168}$ and reduced Ti$_{84}$O$_{167}$ NPs at PBE0 level. The triplet state energy levels are for spin-up eigenvalues. The red dotted lines correspond to the standard redox potentials for water splitting at pH = 0 (H$^+$/H$_2$ = -4.44 eV and O$_2$/H$_2$O = -5.67 eV). The singly and doubly occupied defect states are denoted in blue and green for triplet and singlet states, respectively. Other occupied and unoccupied molecular orbitals are represented with black lines.
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