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# Origin of photocatalytic activity enhancement in Pd/Pt-deposited anatase N-TiO<sub>2</sub> – experimental insights and DFT study of the (001) surface†

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In pursuit of the ideal photocatalyst, cheap and stable semiconductor TiO<sub>2</sub> is considered to be a good choice if one is able to reduce its band gap and decrease the recombination rate of charge carriers. The approach that offers such improvements for energy conversion applications is the modification of TiO<sub>2</sub> with nitrogen and noble metals. However, the origin of these improvements and possibilities for further design of single-atom catalysts are not always straightforward. To shed light on the atomic-scale picture, we modeled the nitrogen-doped (001) anatase TiO<sub>2</sub> surface as a support for palladium and platinum single-atom deposition. The thermodynamics of various synthesis routes for Pd/Pt deposition and nitrogen doping is considered based on density functional theory (DFT)-calculated energies, highlighting the effect of nitrogen doping on metal dimer formation and metal–support interaction. XPS analysis of the valence band of the modified TiO<sub>2</sub> nanocrystals, and the calculated charge transfer and electronic structure of single-atom catalysts supported on the (001) anatase TiO<sub>2</sub> surface provide an insight into modifications occurring in the valence zone of TiO<sub>2</sub> due to nitrogen doping and Pd/Pt deposition at the surface. DFT results also show that substitutional nitrogen doping significantly increases metal–support interaction, while interstitial nitrogen doping promotes only Pt–support interaction.

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#### 1. Introduction

Surface properties play a crucial role in the reactivity of nanomaterials. Aiming for the successful photocatalytic applications of nanosized semiconductors, one is obliged to understand, model and modify their surface according to the desired behavior. Control over surface properties, and consequently reactivity, is of

significant interest for future applications, mainly in renewable energy<sup>1,2</sup> and environmental chemistry.<sup>3</sup>

TiO<sub>2</sub> is, due to its stability, low price, and good physicochemical characteristics, a material of the highest interest in semiconductor engineering. Especially, the anatase TiO2 phase has the highest electron mobility in comparison to the rutile and brookite phases.<sup>4</sup> Numerous studies have been performed aiming to reduce the TiO2 band gap and achieve wider utilization of the solar spectrum and thus improve its photocatalytic properties. A successful approach proved to be doping with non-metals,5-7 doping with metals such as Pt,8 or interfacing metals by deposition at the surface.9-12 The Pt ion at the TiO2 surface acts as a chargegeneration center, producing free and trapped charges. 13,14 Density functional theory (DFT)+U study revealed that N-doping promotes dissociative water adsorption, leading to favorable adsorbate-substrate interactions and stabilization of OH groups at the TiO2 surface. 15 Therefore, combined metal modification and non-metal doping is a good approach for catalyst tailoring. 16-18 This, however, should be treated carefully, since dopants introduced in the TiO<sub>2</sub> lattice can have states near the conduction band (CB), allowing the formation of mid-gaps that can accelerate electron-hole recombination rates and hinder photocatalytic performance. 19

Given that photocatalytic activity is considerably affected by the exposed facets, non-metal doping, for band gap optimization,

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combined with synthesis routes for facet selectivity, has been proposed in several studies.  $^{4,7,20,21}$  Although the surface energy of the (001) anatase surface is larger than that of the (101) surface,  $^4$  few experimental approaches exist to synthesize catalysts with the (001) facet exposed. For example, Zhang *et al.*  $^{22}$  obtained Ca and N co-doped TiO<sub>2</sub> sheets with the (001) facet exposed through a one-step hydrothermal process. The higher surface energy of the (001) surface is, in general, related to better photocatalytic reactivity.  $^7$ 

Doping of surface sites is, in fact, functionalization of the TiO<sub>2</sub> surface, and such a modified surface plays a catalytic role in complex photo-induced reactions enabling advanced technological applications. 1,23 The growth of metal nanoparticles on a TiO2 substrate has been widely studied; Pt nanoparticles on TiO<sub>2</sub> were encapsulated by Ti and O from the support, forming significantly anisotropic Pt@TiO2 interfaces along different directions.24 Cu clusters on TiO2 were studied as promising candidates for methanol synthesis reactions.<sup>25</sup> Ag supported on titania has been found to be efficient in the reduction of harmful gases such as nitrogen oxides.26 Lately, single-metal atom catalysts have been attracting attention as a way of obtaining targeted properties, while consuming significantly fewer resources.<sup>27</sup> Single-atom deposition on top of the N-TiO2 surface can be regarded as a method for fine-tuning of charge states of adsorbed metals, and, in this case, understanding the behavior of TiO2 as a support for metal deposition is also significant. For instance, in their work regarding Au and Ag deposition on TiO2, Schlexer et al. showed how non-metal dopants stimulate charge transfer between the metal and oxide surface.16

Our earlier work<sup>28</sup> showed that nitrogen-doping of a nanosized TiO<sub>2</sub> surface, combined with Pd or Pt deposition at the surface, generates unique properties of these materials, *e.g.* the combination of a suitable band gap (2.35 eV) and higher absorption in the visible light range as compared to N-doped TiO<sub>2</sub>, which enable better utilization of visible light. In this article, we extend previous research by reporting the X-ray photoelectron spectroscopy (XPS) analysis of the electronic structure of the valence zone, which outlines the synergetic effect of noble metals and nitrogen at the TiO<sub>2</sub> surface. Photocatalytic hydrogen evolution from methanol solution using Pd or Pt@N-TiO<sub>2</sub> was also demonstrated in the work of Chiu *et al.*<sup>29</sup>

To study the origin of such improvements, and to understand the fundamental properties of Pd or Pt modified N-TiO<sub>2</sub>, as well as the interaction between N, Pd, and Pt atoms and the support, we use DFT to model N-doping and deposition of noble metals (Pt and Pd) as single atoms. Deposition of two metal atoms and dimer formation on the (001) anatase surface are also addressed. The studied systems are a model of single atom catalysts (SACs), with catalytic performance highly dependent on the spatial configuration and the electronic states of the deposited metals that interact with the support. This interaction modifies the stability, dispersion, and the charge states of SACs, and the local electronic states of the support.<sup>30</sup> The DFT approach provides an atomic-level insight into the fundamental properties of materials, which can enable the understanding of material performances. Some examples include the achieved

synthesis of anatase TiO<sub>2</sub> with the exposed (001) facets<sup>22</sup> for hydrogen generation from methanol or ethanol solutions, and ultimately water, or the design of TiO<sub>2</sub>-based supported catalysts for proton-exchange membrane fuel cells.<sup>10</sup>

# 2. Experimental methods

Nitrogen-doped TiO<sub>2</sub> (N-TiO<sub>2</sub>) was synthesized *via* a solvothermal method<sup>31</sup> using urea (Acros Organics, purity 99%) as a nitrogen source and titanium(v) isopropoxide (Acros Organics, purity 98%) as a titanium source (resulting in the molar ratio N/Ti = 0.1). The final solution was evaporated to dryness in a steam bath, and subsequently heated at 400 °C for 4 hours in a furnace (Termolab Fornos electronics LDA). Palladium and platinum were deposited on the N-TiO<sub>2</sub> surface by ultra-violet reduction from chloride solutions, targeting a 0.10 wt% Pt loading (Pt/N-TiO<sub>2</sub>) and 0.05 wt% Pd loading (Pd/N-TiO<sub>2</sub>), using a photo-reactor with a 450 W Hg immersion lamp (ACE Glass Incorporated, NJ) and irradiation time of 60 min at a constant temperature (30–40 °C). The solution was filtered, dried and finally calcined at 440 °C for 1 h to obtain Pt, Pd/N-TiO<sub>2</sub> nanoparticles.

XPS analysis of the valence band (VB) was carried out in a SPECS customized ultra-high vacuum surface analysis system containing a PHOIBOS 100 spectrometer for energy analysis, a dual-anode Al/Ag monochromatic source and an electron flood gun. XPS spectra were obtained in fixed analyzer transmission mode using a monochromatic Al K $\alpha$  line (photon energy of 1486.74 eV). Survey and high resolution spectra were obtained with the pass energy of 40 eV (energy step 0.5 eV) and 20 eV (energy step 0.1 eV), respectively. The position of the Ti  $2p_{3/2}$  line, expected to be at 458.6 eV in the case of anatase, <sup>32</sup> was used as a binding energy reference for all samples.

#### 3. Theoretical calculations

The effects of nitrogen doping and metal deposition on the (001) surface of anatase  ${\rm TiO_2}$  were studied using spin-unrestricted DFT-based calculations, with the Perdew–Burke–Ernzerhof general gradient approximation (PBE-GGA)<sup>33</sup> and a pseudopotential plane wave approach, as implemented in the Quantum ESPRESSO package. The value of the calculations: 3s, 3p, 3d, and 4s states for Ti, 2s and 2p states for O and N, 5d, 6s, and 6p states for Pt and 4d and 5s states for Pd. Plane waves were expanded using a kinetic energy cut-off of 30 Ry, while for charge density, a 16 times larger cut-off was adopted. An irreducible Brillouin zone was sampled using Monkhorst–Pack sampling and a 2 × 2 × 1 grid of k-points.

To correct the shortcomings of the GGA functional when addressing the electronic structure of the studied surfaces, the U value of 3 eV was applied to the 3d states of Ti, as it was not shown to significantly affect the ordering of vacancy and dopant states in the gap.<sup>36</sup> On the other hand, very large and unphysical U values (U = 6 eV) for the on-site correction of Ti 3d states were required to reproduce the experimental band gap;<sup>23</sup>

therefore, the value used in our work improves slightly the band gap given by GGA, and is also shown to be optimal for the study of metal deposition on a TiO2 surface, giving satisfactory precision of total energy differences for these applications. 16,36 Additionally, doping energies calculated with and without U correction were compared, and aside from the energy shift when U is applied, no change in general trend is seen.

The anatase TiO<sub>2</sub> bulk structure was initially optimized, giving lattice parameters a = 3.816 Å and c = 9.513 Å, which are in agreement with the experimentally reported values (3.796 Å and 9.444 Å).<sup>37</sup> The (001) surface of TiO<sub>2</sub> was modeled using a (3  $\times$  3) supercell, containing four Ti layers and twice as many O layers in a slab, with a 15 Å vacuum separating the surface slabs. A section of the relaxed (001) slab containing the model cell is shown in Fig. 1. The resulting model cell contains 96 atoms (32 Ti, 64 O), and it was relaxed with the two bottom Ti layers and corresponding O layers fixed at their bulk positions, until the forces acting on the atoms were less than 0.001 eV Å<sup>-1</sup> in any Cartesian direction.

According to the standard notation in the literature, 23 coordinately unsaturated surface Ti and O atoms were labeled Ti<sub>5c</sub> and O<sub>2c</sub>, respectively, while fully coordinated O atoms in the subsurface layer were labeled  $O_{3c}$ . Following the supercell relaxation, Ti<sub>5c</sub>-O<sub>2c</sub> surface bonds were found to be nonequivalent, with bond distances of 1.74 Å and 2.26 Å, in agreement with previous reports (1.8 Å and 2.2 Å).<sup>23</sup> The number of slab layers in the model cell was sufficient to assure consistency of the calculated surface energy for the (001) anatase surface (0.98 J m<sup>-2</sup>) with the value of 1.01 J m<sup>-2</sup> obtained by Liu et al.38

An identical relaxation procedure was applied while modeling the nitrogen doping and metal deposition on top of the relaxed

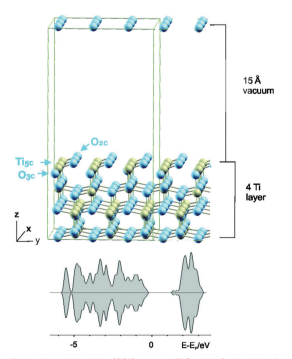


Fig. 1 Relaxed model of the (001) anatase TiO<sub>2</sub> slab (unit cell is framed); blue and gray spheres denote oxygen and titanium, respectively. At the bottom of the image, the resulting total density of states is shown.

(001) TiO<sub>2</sub> surface. Two types of nitrogen doping were simulated in our study, interstitial doping (Nint-TiO2) and substitutional doping (N<sub>sub</sub>-TiO<sub>2</sub>). Interstitial doping was modeled by placing a N atom on the surface in plane with the other atoms, between the first and the second Ti layers, and between the second and the third layers. After relaxation of the supercells, the N atom was positioned above the surface, see Fig. 5a; regardless, we refer to this configuration as interstitial. For the substitutional doping, we addressed two cases, i.e. nitrogen substituting O3c and O2c atoms (see Fig. 1) in the surface layer. In the case of metal deposition on the nitrogen-doped TiO<sub>2</sub> surface, the most stable interstitial and substitutional N-TiO2 surfaces were employed as the supports for metal atoms. This is a simplification of the correlated nitrogen-metal effect, but it was done with the aim to simulate the experimental procedure used in this work. However, some implications of other synthesis routes are discussed based on the total energy differences provided in Section 4.4. Deposition of Pt or Pd was modeled by adding metal atoms onto various surface sites next to nitrogen on the (001) surface - on top of Ti and at the bridge site. All structures were relaxed and optimized according to the procedure used for the starting slab. In addition, the deposition of metals away from the surface nitrogen was tested (labeled in text as next), as well as the deposition of metal atoms on top of the relaxed surface of the clean (pristine) (001) anatase TiO2 surface. To model metal dimer formation at the surface during deposition, two metal atoms were placed next to each other at the surface, and the obtained structures were fully relaxed.

#### 4. Results and discussion

#### 4.1. The motivation for follow-up research

Our earlier work<sup>28</sup> addressed Pd or Pt deposition at the surface of nanosized N-TiO2, which proved to be an efficient catalyst in photocatalytic hydrogen production from ethanol. A wide range of applications of N-doped TiO2 led to the development of many approaches, such as sputtering, ion implantation, pulse-laser deposition, sol-gel, oxidation of TiN, hydrothermal processing and NH<sub>3</sub> annealing<sup>39,40</sup> for N-TiO<sub>2</sub> synthesis. Also, various precursors were used, e.g. N-TiO2 nanocatalysts with a homogeneous anatase structure were successfully synthesized through a microemulsion-hydrothermal method by using some organic compounds, such as triethylamine, urea, thiourea, and hydrazine hydrate (the optimal mole ratios of N to Ti for triethylamine, urea, thiourea, and hydrazine hydrate were 2, 1, 0.5, and 0.5, respectively).41

Having in mind this perspective, the approach we used for obtaining Pd, Pt/N-TiO2 with a dominant anatase structure is just one possibility. However, the synthesis procedure and conditions for obtaining N-TiO2 determine not only the dominant TiO<sub>2</sub> phase but also defect concentration and nitrogen position (substitutional and interstitial).<sup>39,42</sup> These structural variations lead to enhanced properties of the photocatalysts, but they also, given that they are a combination of many factors, hinder our understanding of the underlying interactions between nitrogen,

metals, and the TiO<sub>2</sub> surface. Our intention in this follow-up study was to understand the origin of the photocatalytic enhancement of nitrogen-doped metal-deposited anatase TiO2, by looking at the atomic-scale picture of the TiO2 surface based on the results of DFT calculations. In this work, earlier experimental findings regarding bonding and electronic structure<sup>28</sup> are viewed in correlation with the calculated thermodynamic and bonding properties. Also, additional XPS measurements of the valence band of the studied samples are done, given that the underlying structures of the VB and CB are straightforwardly obtained in the DFT calculations.

#### 4.2. Experimental study of Pd, Pt surface modification and N-doping of anatase TiO2 nanoparticles

Based on the X-ray diffraction (XRD) analysis (Supplementary data 1, ESI†), the studied materials have a dominant structure of the anatase phase, with a small percentage of the rutile phase (3.8%), and crystallite sizes of  $12 \pm 2$  nm. <sup>28</sup> Our previous XPS study<sup>28</sup> also resolved the surface composition of the samples based on the intensities of characteristic photoelectron lines recorded in high resolution and the atomic sensitivity factors provided by the manufacturer of the XPS system. Relative elemental concentration of the atoms on the surface are: 0.4% nitrogen in N-TiO2, 0.4% Pd in Pd/N-TiO2 and 2.1% Pt in Pt/N-TiO<sub>2</sub>. The majority of nitrogen is ascribed to interstitial doping, based on the evidence of N-O bond formation (67.3%), while some of the nitrogen is found to substitute the oxygen in the TiO<sub>2</sub> lattice (21.5%) (XPS spectrum shown in Supplementary data 2 (ESI†) and discussed in detail elsewhere<sup>28</sup>). Additionally, chemical states of the studied noble metals were addressed; as shown in Fig. 3, Pt is found to be present in a partially oxidized state. Three contributions to the Pt  $4f_{7/2}$  line at 70.8 eV (49.1%), 72.5 eV (36.6%), and 74.7 eV (13.4%) were observed, giving evidence of metallic platinum, presumably in the Pt-Ti-O bond, and a small amount of PtO2.28

The Pd 3d<sub>5/2</sub> line situated at 336.6 eV has a unique contribution fully consistent with the PdO phase,28 showing that all of the palladium found at the surface is in the Pd<sup>2+</sup> form, see Fig. 2.

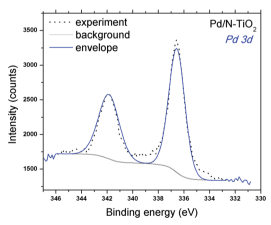


Fig. 2 XPS spectrum of the inelastic contribution of Ti 3s and Pt 4f lines from the Pt/N-TiO<sub>2</sub> sample (reproduced from ref. 28 with permission from the PCCP Owner Societies).

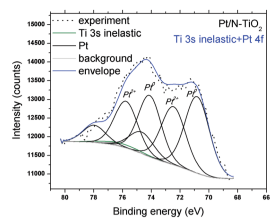


Fig. 3 Detailed XPS spectrum of the Pd 3d line from the Pd/N-TiO<sub>2</sub> sample (reproduced from ref. 28 with permission from the PCCP Owner Societies).

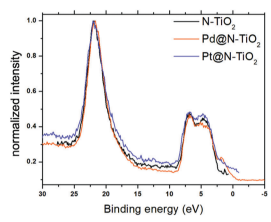


Fig. 4 XPS spectra of VB (intensity normalized to the maximum of the O 2s line) for N-TiO<sub>2</sub>, Pd/N-TiO<sub>2</sub>, and Pt/N-TiO<sub>2</sub>, recorded with a pass energy of 20 eV (FAT 20 regime).

Valence bands of the reference material (N-TiO2) and Pdand Pt-loaded N-TiO2 were recorded; Fig. 4 shows the normalized VB spectra. The position of the O 2s line at 21.8 eV overlaps for the three samples, justifying the usage of the Ti 2p<sub>3/2</sub> line as a binding energy reference. The valence band shape is the result of two peaks at 8 eV and 6 eV, resulting from  $\pi$  and  $\sigma$  O 2p orbitals, respectively, 43 confirming the dominant anatase TiO2 structure.

Based on the intersection of the valence band edge and background asymptotes, we find the top of the valence band of the reference sample to be around 2.2 eV below the Fermi level. Having in mind the XPS line broadening due to the nonuniform charging of the samples, we expected to find the VB edge at higher binding energies as compared to those of the theory<sup>44</sup> and UPS (ultraviolet photoelectron spectroscopy) experiments. 45 In the case of Pt- and Pd-loaded samples, the situation is significantly changed: the top of the VB moved toward higher energies, see Fig. 4. For Pd-deposition, a unique level inside the energy gap of the initial N-TiO2 is detected above the VB top (shoulder at 1.8 eV). A similar effect, although

less pronounced, is seen in the case of Pt deposition, where the widening of the VB top is in the shape of an exponentially declining function.

#### 4.3. Nitrogen doping and single-metal atom deposition on the (001) anatase TiO<sub>2</sub> surface – spatial configuration and energetics

We addressed both interstitial doping (N<sub>int</sub>-TiO<sub>2</sub>) and substitutional doping (N<sub>subs</sub>-TiO<sub>2</sub>) of the (001) anatase surface. The former one is of higher significance in relation to our experimental findings, given that previous research28 demonstrated that, following the used synthesis procedure, the majority of nitrogen is relaxed into interstitial positions (see Supplementary data 2, ESI†).

The energy of interstitially doped nitrogen on the (001) TiO<sub>2</sub> surface is calculated with reference to N2(g):

$$\Delta E(N_{int}) = E(N_{int}/TiO_2^{slab}) - 1/2E(N_2) - E(TiO_2^{slab})$$
 (1)

where the energy of the reconstructed surface is taken for the energy of TiO<sub>2</sub><sup>slab</sup>.

The findings of Ortega et al. and Finazzi et al., regarding the stability of nitrogen on the (101) anatase surface, 6,18 also indicate that the most stable structure is interstitially doped nitrogen on the surface of the TiO2 slab. Based on the smallest value of the implantation energy for the (001) surface, the results obtained in our simulations coincide with the previously mentioned ones (sub-surface implantation energy is 2.32 eV larger). The relaxed N<sub>int</sub>-TiO<sub>2</sub> slab is shown in Fig. 5a and the corresponding energetics is available in Table 1. Significant rearrangement of the surface structure is seen, with formation of a short N-O bond; the calculated N-O distance is 1.27 Å, which is in accordance with earlier reports (1.3-1.4 Å). 18

The energy of nitrogen substituting oxygen on the (001) anatase TiO<sub>2</sub> surface, i.e. the substitution energy, is determined as:

$$\Delta E(N_{subs}) = E(TiO_{2-x}^{slab}N) - 1/2E(N_2) + 1/2E(O_2) - E(TiO_2^{slab})$$
(2)

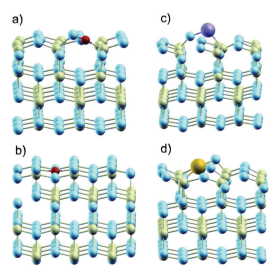


Fig. 5 Energetically preferred arrangements of the modified (001) anatase  $TiO_2$  surface; (a)  $N_{int}$ - $TiO_2$ , (b)  $N_{subs}$ - $TiO_2$ , (c)  $Pt@TiO_2$ , and (d)  $Pd@TiO_2$ . Blue, gray and red small spheres denote oxygen, titanium, and nitrogen, while large purple and yellow spheres denote Pt and Pd, respectively.

The relaxed N<sub>subst</sub>-TiO<sub>2</sub> structure is shown in Fig. 5b. The substitution of the O<sub>3c</sub> atom is found to be more energetically favorable ( $\Delta E = +4.60 \text{ eV}$ ) than substitution of the  $O_{2c}$  atom  $(\Delta E = +4.77 \text{ eV})$  by 0.17 eV, which is in accordance with the threefold coordination of the N atom and corresponds to earlier reports for other anatase surfaces. 6,18 Earlier DFT studies showed that nitrogen sub-surface or bulk substitutions are slightly favored over surface substitutions. 18 However, to study the interaction of Pt and/or Pd with N atoms, we restrict our calculations to the most stable structures: interstitial nitrogen on the surface (Nint-TiO2), and the more stable of the two surface substitutional cases - nitrogen substituting O<sub>3c</sub> (N<sub>subs</sub>-TiO<sub>2</sub>). Both interstitial and substitutional nitrogen doping energies are positive, showing that nitrogen doping by molecular N<sub>2</sub>(g) is thermodynamically unfavorable.

Table 1 also presents the calculated energies of metal deposition on the clean (reconstructed) TiO2 surface, calculated as:

$$E(\mathbf{M}_x) = E(\mathbf{M}_x \circledast \text{TiO}_2^{\text{slab}}) - E(\text{TiO}_2^{\text{slab}}) - xE(\mathbf{M}); \quad x = 1, 2; \quad \mathbf{M} = \text{Pd}, \text{ Pt}$$
(3)

where E(M) is the energy of an isolated noble metal atom. The final structures of these two cases are shown in Fig. 5c and d. The relaxed structure of Pt doped (001) anatase TiO2 is in agreement with the DFT study of Li et al.11

In addition to our results, Table 1 summarizes previously published energies of nitrogen doping and metal deposition on the (101) anatase TiO<sub>2</sub> surface, which is energetically the most favorable anatase TiO2 surface. Although the previous DFT results and model surface sizes somewhat differ, a general trend regarding the energetics of nitrogen doping and platinum deposition is qualitatively consistent for the (001) and (101) surfaces. Positive nitrogen doping energies indicate an endothermic reaction for adsorption of nitrogen from the gas phase, which is expected given the high stability of molecular nitrogen. 18 Nevertheless, calculating the energy with reference to the standard state of nitrogen (N2) provides an easy comparison of the nitrogen-surface interaction and is widely accepted in the literature. Synthetic routes use other nitrogen sources and external conditions in the synthesis, and the final nitrogendoped slab is stable (negative energy).

Charge transfer for the clean (001) anatase TiO2 surface and the N-doped surface is always from Pd to the support and from the support to Pt (Supplementary data 3, ESI,† see detailed

Table 1 Calculated (PBE-GGA) energies of nitrogen doping and energies of single metal atom deposition ( $\Delta E$ ) on the (001) anatase TiO<sub>2</sub> surface. Literature data regarding nitrogen doping and noble metal deposition on the (001) and (101) anatase TiO2 surfaces are provided for comparison

	$\Delta E$ (eV)			
Dopant	$(001)~{ m TiO}_2$	(101) TiO <sub>2</sub>		
N <sub>int</sub>	2.07	2.71 <sup>18</sup>		
$N_{ m subs}$	4.60	$4.92^{18}$		
Pt	-3.49	$-2.84^{46}$		
	$-2.93^{8}$			
Pd	-3.17	$-0.39^{47}$		
Au		$-0.61^{16}$		
Ag		$-1.04^{16}$		

Discussion of charge transfer in Section 4.6). Although charge transfer from metal to support is not typical, similar behavior to that of palladium is seen in the case of Au deposition on TiO<sub>2</sub>. 18

#### 4.4. Structure and energetics of Pd and Pt deposition on the N-doped anatase (001) TiO2 surface

After considering the mono-atomic deposition of N, Pt, or Pd on top of the clean (001) TiO2 surface, a synergy of nitrogen doping and metal deposition on the surface is considered. In Table 2, we summarize the stabilities of Pt@N-TiO2 and Pd@N-TiO2 for both interstitially and substitutionally doped N-TiO2. To illustrate the experimental case of metal deposition, the energy of M@N-TiO<sub>2</sub> is referenced to the energy of the corresponding N-TiO<sub>2</sub>. Additionally, to study other possible doping/deposition strategies, the energy of M@N-TiO2 is evaluated with respect to the energy of the metal-deposited (M@TiO2) surface or clean TiO2 surface.

In Table 2, we see the small energy differences calculated among various reactions in some of the discussed cases; therefore, other factors including geometry, kinetics or experimental conditions during synthesis might lead to different deposition priorities.19

By comparing the presented energies with energies of deposition on the clean (001) surface, we concluded that doping anatase surface with nitrogen significantly increases metal-support interaction, i.e. promotes platinum deposition. In comparison to the deposition on clean TiO2, metal deposition energy is lower for substitutionally nitrogen-doped surface by 2.32 eV, and for interstitially nitrogen-doped by 0.97 eV. The same trend was shown for Au deposition alongside N-doped 101 anatase TiO2 in the study by Ortega et al. 18 On the other hand, deposition energy of Pd on N<sub>int</sub>-TiO<sub>2</sub> is similar to the deposition energy of Pd on clean TiO<sub>2</sub>, while N<sub>subs</sub>-TiO<sub>2</sub> promoted this action by lowering the deposition energy by 1.23 eV. These trends are further discussed based on the charge transfer in the next section. Looking at the synthesis strategies simultaneous nitrogen doping and metal deposition could be achieved from a thermodynamic point of view. Nitrogen doping of metal-modified TiO2 surface from the gas phase is still thermodynamically unfavorable (positive doping energies), although in some cases much more feasible as compared to doping of clean TiO2 surface - this is especially true for interstitial nitrogen doping of platinum-deposited surface (nitrogen implantation energy lowered from 2.07 to 0.75 eV).

In summary, doping the anatase surface with nitrogen lowers metal adsorption energy in the case of substitutional nitrogen doping, showing that such nitrogen implantation increases metal-support interaction for both metals. On the other hand, interstitial nitrogen doping promotes Pt-support interaction, while for Pd no significant change in deposition energy is seen. However, one must keep in mind that substitutional doping is much less favored than interstitial doping (Table 1).

#### 4.5. Energetics of metal dimers on pristine and N-doped anatase (001) TiO2 surface

Beside a single-atom surface deposition, understanding the growth of metal clusters is also valuable for catalytic applications, as it can lead to new mechanisms that accelerate the catalytic reaction. 48,49 On the other hand, for some applications, including single-atom heterogeneous catalysis, deposition of a single-atom is essential and clustering must be avoided. 50,51 To simulate the initial step of the small cluster formation process at the (001) anatase TiO2 surface (represented by the deposition of two metal atoms) and quantify the probability of small metal cluster formation on nitrogen-doped (001) anatase TiO2, we compare the energetics of a single metal atom deposition on the surface with the energetics of metal dimer deposition. Metal dimer deposition energy on the clean and Nint-doped (001) anatase  $TiO_2$  surface is calculated by taking x = 2 in eqn (3), meaning that here we refer to the energy released upon deposition of two separate metal atoms next to each other to form a dimer on the surface as the metal dimer deposition energy. Results are presented in Table 3.

By comparing the energy released during the deposition of one (Table 1) and two metal atoms (Table 3) on the pristine (001) anatase TiO<sub>2</sub> surface, one can argue that clustering is probable in the case of Pt, due to a large energy release for the second atom deposition. The identical behavior for Pt deposition at the (101) anatase surface is shown in the work of Han et al., 46 where they demonstrated the growth of small Pt clusters. Vittadini and Selloni showed, in a DFT study<sup>47</sup> of the interaction of small Au clusters (Au<sub>n</sub>, n = 1-3) with a stoichiometric and reduced (101) TiO2 surface, that gold clustering on the TiO2 surface is also favored. However, in the latter case, this conclusion holds for the case of a low concentration of vacancies on reduced (101) TiO<sub>2</sub> as they serve as nucleation centrers.<sup>47</sup> In the case of Pd deposition on pristine TiO2, Nint-TiO2 and Pd@Nint-TiO2, dimer

 $\textbf{Table 2} \quad \textbf{Calculated reaction energy } (\Delta E) \text{ and difference of reaction energy and corresponding energy on clean TiO}_2 (\Delta E - \Delta E^{\text{pristine}}) \text{ of (1) single-metal energy on clean TiO}_2 (\Delta E - \Delta E^{\text{pristine}}) \text{ of (2) single-metal energy ene$ atom deposition on the (001) facet of nitrogen-doped TiO<sub>2</sub>; (2) simultaneous procedure of metal deposition and nitrogen doping of the (001) anatase TiO<sub>2</sub> surface; (3) nitrogen doping of the metal deposited (001) TiO<sub>2</sub> surface

			$\Delta E$ (eV)		$\Delta E - \Delta E^{\text{pristine}}$ (eV)	
Type of reaction		N position	M = Pt	M = Pd	M = Pt	M = Pd
(1)	$M + N-TiO_2^{slab} \rightarrow M@N-TiO_2^{slab}$	N <sub>int</sub>	-4.46	-3.03	-0.97	+0.14
((a)	$M + 1/2N_2 + TiO_2^{slab} \rightarrow M@N-TiO_2^{slab}$	$N_{ m subs}$	-5.81	-4.40	-2.32	-1.23
((2)	$M + 1/2N_2 + 11O_2^{-11} \rightarrow M(a)N-11O_2^{-11}$	$ m N_{int} \ N_{subs}$	$-2.38 \\ -1.21$	$-0.96 \\ 0.2$	_	
(3)	$1/2N_2 + M@TiO_2^{slab} \rightarrow M@N-TiO_2^{slab}$	N <sub>int</sub>	0.83	2.22	-1.24	+0.15
		$N_{ m subs}$	2.28	2.90	-2.32	-1.70

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Table 3 Calculated energy (ΔΕ) of (1) metal dimer deposition on clean (001) anatase TiO<sub>2</sub> surface; (2) metal dimer deposition on nitrogen-doped (001)

anatase N<sub>int</sub>-TiO<sub>2</sub> surface; (3) secondary metal deposition on single-metal deposited and N-doped (001) anatase surface (M@N<sub>int</sub>-TiO<sub>2</sub>). For N-doped surface, reaction energy is compared with the energy of same reaction on the clean surface ( $\Delta E - \Delta E^{\text{pristine}}$ )

		$\Delta E$ (eV)		$\Delta E - \Delta E^{\text{pristine}}$ (eV)	
		M = Pt	M = Pd	M = Pt	M = Pd
(1) Metal dimer on TiO <sub>2</sub>	$2M + TiO_2^{slab} \rightarrow M_2 @TiO_2^{slab}$	-8.20	-5.21		
(2) Metal dimer on N <sub>int</sub> -doped TiO <sub>2</sub>	$2M + N_{int}\text{-TiO}_2^{slab} \rightarrow M_2@N_{int}\text{-TiO}_2^{slab}$	-8.72	-6.23	-0.52	-1.02
(3) Metal deposition on M@N $_{\rm int}$ -doped TiO $_2$	$\mathrm{M} + \mathrm{M}@\mathrm{N}_{\mathrm{int}}\text{-}\mathrm{TiO}_{2}^{\mathrm{slab}} \to \mathrm{M}_{2}@\mathrm{N}_{\mathrm{int}}\text{-}\mathrm{TiO}_{2}^{\mathrm{slab}}$	-4.26	-3.20	+0.45	-1.16

formation is energetically quite similar, ranging from -3.03 eV to -3.20 eV (Tables 1-3).

The calculated energy released upon deposition of a second Pt atom at Nint-TiO2 (Table 3, the last row) is less compared to the energy released for the first atom (Table 2, the last row); however, it is larger than the deposition energy on the clean TiO<sub>2</sub>. Based on the calculated values presented in the tables, the energy released upon single metal deposition, on the clean or modified TiO<sub>2</sub>, follows the trend: Pt@Pt/TiO<sub>2</sub> > Pt@N<sub>int</sub>-TiO<sub>2</sub> > Pt@Pt/N<sub>int</sub>-TiO<sub>2</sub> > Pt@TiO<sub>2</sub> for platinum deposition, and  $Pd@TiO_2 \approx Pd@N_{int}\text{-}TiO_2 \approx Pd@Pd/N_{int}\text{-}TiO_2 > Pd@Pd/N_{int}$ TiO<sub>2</sub> for palladium deposition.

Now, let us consider Pt and Pd deposition on the N-TiO2 surface. If we start with very low concentrations, in the case of Pt, the most favorable adsorption site is the N-site. It is almost 1 eV more favorable than adsorption on the pristine surface (i.e. far from N, see Tables 2 and 3). So, single Pt atoms will initially deposit on N sites. If a higher Pt loading is considered, new Pt atoms arriving on the surface can choose between Pt@N-TiO2 and free N-sites. The latter ones are energetically more favorable and also if the concentration of Pt@N-TiO<sub>2</sub> is still low, the probability of making a dimer is low. As the concentration increases further, Pt atoms previously adsorbed at N-sites will serve as nucleation centers for dimer formation and further growth of larger particles. Pt atoms will not favor pristine (001) terrace sites as the adsorption energy is much smaller compared to that of other configurations. So, in the case of a low concentration of Pt, single Pt atoms are likely to be anchored to N-sites at the surface. Now, let us consider the same scenario for Pd deposition. At a very low concentration of Pd, pristine (001) terrace sites are slightly more favored than N-sites (Tables 1 and 2). So, there should be basically no preference between available sites (the difference is only 0.14 eV) and at a low concentration of Pd, single Pd atoms should be adsorbed on anatase, which was, just recently, confirmed experimentally.<sup>52</sup> As one starts to increase the concentration of Pd, dimers will start to form. Dimer formation is, however, favored on N-sites, so these sites can be considered as the nucleation centers for larger Pd particles. Nevertheless, agglomeration will take place even if there is no nitrogen dopant, if the concentration increases over a certain limit.<sup>52</sup> When larger particles grow at high metal concentrations, one can consider that cohesive energy is released per atom and both Pd and Pt have high cohesive energies.

To summarize so far, nitrogen interstitial doping can, therefore, be seen as a way of additional stabilization of the singleatom surface deposition, when platinum concentrations are comparable with the concentration of surface nitrogen. For both Pt and Pd, a larger energy for dimer deposition is obtained at Nint-TiO2 as compared to that at the clean TiO2 surface, suggesting that when depositing metals in higher concentration, N-doped sites also serve as nucleation centers for dimer formation and possibly formation of larger metal clusters. However, in the case of Pd deposition, dimer formation is energetically the least favorable on the pristine TiO2, while on the nitrogen-doped surface, no significant difference in deposition energy per atom is seen when depositing one or two atoms. Based on the obtained results, single atom dispersions of both Pt and Pd can be achieved on N-doped anatase (001) TiO2, but in the case of platinum, doping plays the crucial role from the energetic point of view. However, the presence of surface N affects the electronic structure of both deposited metals, as shown below.

#### 4.6. Electronic structure trends

The density of states (DOS) for mono- and di-atomic metal deposition on a clean (001) anatase TiO<sub>2</sub> surface is presented in Fig. 6. Deposited metals contribute with their electronic states mainly in the band gap energy range, either on top of the VB or, in some cases, the bottom of the CB. Having in mind that p-type impurities have their states mainly located near the top of the VB, while n-type impurities have their states near the CB edge, the DOS shown in Fig. 6 implies that even atoms of the same type will behave differently if surface clustering occurs. Given the targeted electronic structure for photocatalytic water splitting, to achieve overpotentials for both proton reduction and oxygen species oxidation (i.e. H<sub>2</sub> and O<sub>2</sub> evolution reactions), the occurrence of the additional states at the top of the VB is desirable. Adding one more metal atom to form a surface dimer reflects in extra electronic states and further narrowing of the band gap. In all cases, there is no polarization of charge on either metal or substrate atoms.

Doping the (001) surface with nitrogen introduces an empty N-2p state in the band gap, which is consistent with every studied N position (surface, subsurface...). 16 Some difference is observed between substitutional and interstitial nitrogen implantation; this is discussed for bulk doping in our previous work.<sup>28</sup> Given that the majority of surface nitrogen is in the form of interstitial nitrogen, 28 we further analyze the details of the electronic structure modifications resulting from a combined interstitial nitrogen doping and metal deposition. In agreement with other studies, 18 interstitial nitrogen introduces three N-p orbitals; one free above the Fermi energy. In Fig. 7, a detailed analysis of metal deposition and nitrogen-doping shows how Pd

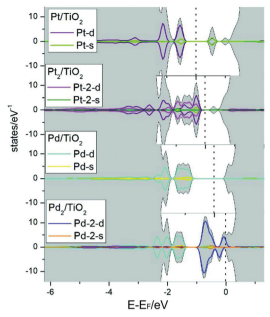


Fig. 6 Total density of states for the studied cases of single metal atoms and metal dimers on the clean TiO2 surface (gray) with angular momentum-resolved contributions for metals (for the dimer cases, labels for the first atom correspond to the mono-doped case). Fermi energy is highlighted for each surface using a vertical dotted line

and Pt states overlap with N states. Overall, this results in a continual population on top of the VB, as opposed to discrete dopant levels seen in the case of nitrogen doping. Additional charge is transferred to the N atom when Pd is deposited next to NO; this effect is seen as occupation of previously empty nitrogen states, comparing N<sub>int</sub>-TiO<sub>2</sub> and Pd@N<sub>int</sub>-TiO<sub>2</sub> in Fig. 7. This is also evident in the calculated charge transfer, presented in Fig. 9. Deposition further from N (labeled next) does not lead to significant coupling of nitrogen and metal states (Fig. 7, Pdnext@Nint-TiO2).

On the other hand, Pt doping does not affect the occupation of empty N orbitals, given that platinum orbitals are lower in energy. This is the main difference in the behavior of palladium and platinum as surface modifiers; this trend is also present in the calculated charge transfer, shown in Fig. 9 and 10. Experiments by Pepin et al. 53 demonstrated the photocatalytic enhancement in the activity for photocatalytic reactions of acetaldehyde due to platinum deposition on TiO2. As observed in the mentioned work, oxygen species adsorbed at the surface react with the photogenerated conduction band electrons captured by the Pt deposits, which prevents recombination of the photogenerated charge carriers. Having this in mind, nitrogen doping does not seem to significantly hinder photocatalytic enhancement given that Pt deposited at the surface of N-TiO2 still has available empty electronic states.

Changes in electronic structure due to metal dimer deposition on N<sub>int</sub>-TiO<sub>2</sub> are presented in Fig. 8. When depositing a Pd dimer on the surface of N<sub>int</sub>-TiO<sub>2</sub>, empty states of the second Pd atom shift above the Fermi energy, implying that if Pd clustering was achieved, it could serve as a center for electron transfer between

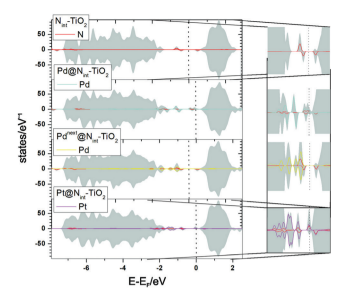


Fig. 7 Total density of states of single metal atoms deposited on the N<sub>int</sub>-TiO<sub>2</sub> surface (gray) with metal-resolved contributions (red line corresponds to the DOS of nitrogen in all graphs); enlarged view of band gap region is shown for each case on the right; Fermi energy is highlighted for each surface using a vertical dotted line.

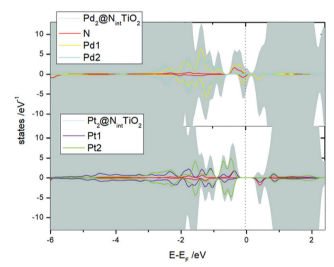
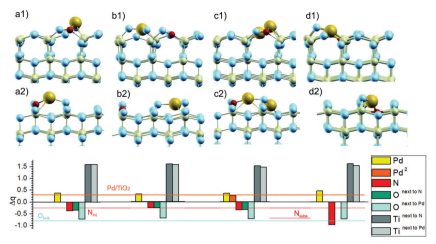


Fig. 8 Total density of states of metal dimers on the N<sub>int</sub>-TiO<sub>2</sub> surface with metal-resolved contributions, and nitrogen contribution shown in red; Fermi energy is highlighted for each surface using a vertical dotted line.

the species involved, for example, in water splitting (proton reduction and the oxygen species oxidation). In the cases of both Pd- and Pt-dimers there is a strong overlap between the states of the two metal atoms and nitrogen, indicating chemical bonding, which can also be observed in the structures presented in Fig. 9. Clearly, the tuning of the electronic structure and the introduction of new states at the edges of the CB and VB could have a significant impact on the reactivity and catalytic activity of these systems. Recently, it was shown that very small Pt clusters with less than two atomic layers are efficient photoelectron collectors from



 $\textbf{Fig. 9} \hspace{0.2cm} \textbf{(a)} \hspace{0.2cm} Pd@N_{int}-TiO_2\text{, (b)} \hspace{0.2cm} Pd_{next}@N_{int}-TiO_2\text{, (c)} \hspace{0.2cm} Pd_2@N_{int}-TiO_2\text{, and (d)} \hspace{0.2cm} Pd@N_{subs}-TiO_2\text{; (1)} \hspace{0.2cm} \textbf{and (2)} \hspace{0.2cm} denote \hspace{0.2cm} views \hspace{0.2cm} from \hspace{0.2cm} different \hspace{0.2cm} slab \hspace{0.2cm} sides; \hspace{0.2cm} under \hspace{0.2cm} each \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} Pd@N_{subs}-TiO_2\text{; (1)} \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} denote \hspace{0.2cm} views \hspace{0.2cm} from \hspace{0.2cm} different \hspace{0.2cm} slab \hspace{0.2cm} sides; \hspace{0.2cm} under \hspace{0.2cm} each \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} Pd@N_{subs}-TiO_2\text{; (1)} \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} denote \hspace{0.2cm} views \hspace{0.2cm} from \hspace{0.2cm} different \hspace{0.2cm} slab \hspace{0.2cm} sides; \hspace{0.2cm} under \hspace{0.2cm} each \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} Pd@N_{subs}-TiO_2\text{; (1)} \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} denote \hspace{0.2cm} views \hspace{0.2cm} from \hspace{0.2cm} all \hspace{0.2cm} each \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} Pd@N_{subs}-TiO_2\text{; (1)} \hspace{0.2cm} and \hspace{0.2cm} (d) \hspace{0.2cm} Pd@N_{subs}-TiO_2\text{; (2)} \hspace{0.2cm} all \hspace{0.2cm} a$ model cell is a corresponding Löwdin charge chart presenting the calculated charge with reference to the charge of an isolated atom ( $\Delta q$ ).

TiO<sub>2</sub> bulk; by contrast, larger particles with more than two Pt layers are the active sites for hydrogen coupling to catalyze HER.<sup>54</sup>

Having in mind the energetics of single metal atom deposition discussed in Section 4.5, and the detailed DOS shown for metal deposition on clean and N-doped TiO<sub>2</sub>, we can expect that additional structures formed on the top of the VB in metaldeposited samples seen in the XPS spectra (Fig. 4) are due to the interaction of Pt states with nitrogen (for Pt/N-TiO2) or Pd states with the support and nitrogen (for Pd/N-TiO2). Pt dimer formation also leads to additional states on top of the VB of TiO<sub>2</sub> and N-TiO<sub>2</sub> (see Fig. 6 and 8), consistent with the XPS spectra (Fig. 4). In our experimental study of Pt/N-TiO<sub>2</sub> nanocrystals, <sup>28</sup> we found 3 contributions to the Pt line, one of them corresponding to the significant contribution of metallic Pt (Fig. 2), i.e. evidence of Pt-Pt bond formation. Given that DFT results showed the highest energy released for Pt dimer formation on pristine TiO<sub>2</sub>, we expect that clustering of Pt occurred on the TiO2 support, and this might be due to the significantly smaller concentrations of surface nitrogen as compared to the surface Pt found in the studied samples.

The activity of the studied nanocatalysts was determined by the charge state of the metal atom. Charge state also provides an insight into the behavior of a metal during the synthesis; for example, stronger metal-support interactions can be used to avoid sintering during reaction that inactivates the metal. We discuss charge transfer based on the calculated Löwdin charges (valence charge states). The resulting charges presented here are referred to as the charge of isolated atoms, and a negative Löwdin charge indicates excess electrons compared to an isolated atom. For deep-lying Ti and O ions, calculated valence charge states amounted to +1.6 and -0.8, giving somewhat lower values as compared to the Bader charges<sup>8</sup> of +2.6 and −1.3 calculated based on atoms in molecules (AIM). The results obey the stoichiometry and correctly add up for bulk neutrality of the anatase TiO2. In Fig. 9 and 10, these values are represented by the background lines as a reference for discussion of

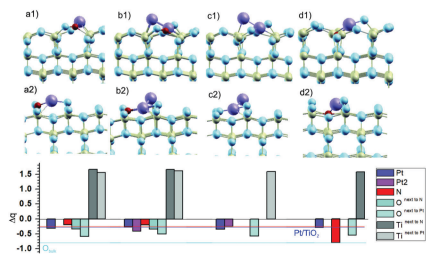


Fig. 10 (a) Pt@N<sub>int</sub>-TiO<sub>2</sub>, (b) Pt<sub>2</sub>@N<sub>int</sub>-TiO<sub>2</sub>, (c) Pt<sub>2</sub>@TiO<sub>2</sub>, and (d)Pt@N<sub>subs</sub>-TiO<sub>2</sub>; (1) and (2) denote views from different slab sides; under each model cell is a corresponding Löwdin charge chart presenting the calculated charge with reference to the charge of an isolated atom  $(\Delta q)$ .

charge transfer, which occurred due to doping. Lines representing the charge of mono-doped nitrogen, and single-atom deposited palladium and platinum are also shown. Calculated Löwdin charges for both substitutional nitrogen (-0.76) and interstitial nitrogen (-0.26) are in excellent agreement with the reported Bader charges  $(-0.89 \text{ and } -0.30, \text{ respectively}).^{18}$ 

Fig. 9 and 10 show relaxed slabs of the energetically most favorable structures for single atom and dimer metal deposition. Based on the charts presented in Fig. 9, we can highlight how co-doping with N and Pd affects charge transfer (Table S1 with the values for the studied cases given in the Supplementary data 3, ESI†).

In all studied cases, charge transfer is directed from palladium to the support, which is a less frequent case of charge transfer. This agrees with our experimental findings for anatase TiO2 nanocrystals,<sup>28</sup> which point out that all of the deposited palladium is in the form of PdO. Unique contributions found for the Pd line in our XPS measurements (Fig. 3) indicate that all Pd atoms are deposited with the same charge transfer towards the matrix and any presence of metallic palladium is excluded. Having in mind the calculated dimer formation energies, we can conclude that our synthesis approach led to the deposition of small amounts of Pd, for which, as discussed in Section 4.5, dimer, and subsequently cluster, formation is not favored. If the metal is in proximity of the surface nitrogen, nitrogen further promotes the charge transfer from palladium. Thus, among the studied cases, the strongest metalnitrogen interaction is observed when Pd is deposited on top of nitrogen, which is substituting for oxygen at the surface.

As can be seen in Fig. 10, Pt is negatively charged when deposited on the (001) anatase TiO2 surface, which agrees with the results of Xia et al. 10 and experimental findings regarding a strong interaction with the support.9 Lewera et al. found in their XPS study an increased electron density on Pt, which was reflected in the asymmetry of the Pt 4f XPS signal.<sup>55</sup>

## 5. Conclusions

Experimental XPS analysis of the valence band of a metal (Pd or Pt)-deposited surface of N-TiO2 nanocrystals is compared to that of N-TiO<sub>2</sub>, showing a shift of the top of the VB toward higher energy. For the Pd-modified sample, an additional level inside the energy gap of the initial N-TiO<sub>2</sub> is detected above the top of the valence zone (shoulder at 1.8 eV).

The structural and electronic properties of the (001) anatase TiO<sub>2</sub> surface, clean and doped with nitrogen, used as a support for Pd and Pt deposition, are studied using DFT+U. The thermodynamics of various synthesis routes for metal-deposited and nitrogen-doped (001) anatase TiO2 surfaces is considered, highlighting the effect of nitrogen doping on metal clustering and metal-support interactions. According to the presented results, the nitrogen substituting oxygen  $(O_{3c})$  significantly increases the metal-support interaction for both Pd and Pt, while interstitial nitrogen doping promotes only the Pt-support interaction. Analysis of diatomic deposition of noble metals on the top of the clean and N-doped surfaces indicates that from the energetic point of view,

clustering can be avoided and single-atom deposition can be achieved for either metal on the Nint-TiO2 surface when depositing in small concentrations.

Electronic structure analysis shows the overlapping of metal and nitrogen states, resulting in a continual population on the top of the VB, as opposed to discrete dopant levels seen in the case of nitrogen doping. Additional charge is transferred to the N atom when Pd is deposited next to NO at the surface, while Pt deposition does not influence the filling of empty N orbitals since Pt orbitals are lower in energy. This is the key difference in the behavior of Pd and Pt single-atoms as surface modifiers, and this general trend is present in the calculated charge transfer results. Clearly, the combination of TiO<sub>2</sub> doping with metal deposition provides an effective way for creation of new reactive centers and tuning the surface properties of the described materials. By a careful control of the metal deposition conditions, it seems that one can control the electronic structure of deposited atoms and the formation of metal aggregates. These results could be a guide for the further design of new TiO2-based (photo)catalysts, while chemical and catalytic properties are yet to be carefully addressed.

### Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

#### **Abbreviations**

VB Valence band CBConduction band

DFT Density functional theory

DOS Density of states

XPS X-ray photoelectron spectroscopy

SAC Single-atom catalyst

# Conflicts of interest

There are no conflicts to declare.

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