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Beyond Shape Engineering of TiO₂ Nanoparticles: Post-Synthesis Treatment Dependence of Surface Hydration, Hydroxylation, Lewis Acidity and Photocatalytic Activity of TiO₂ Anatase Nanoparticles with Dominant {001} or {101} Facets

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Abstract

TiO₂ anatase nanoparticles are among the relevant players in the field of light-responsive semiconductor nanomaterials used to face environmental and energy issues. In particular, shape-engineered TiO₂ anatase nano-sheets with dominant {001} basal facets gained momentum because of the possibility to exploit different and/or improved functional behaviors with respect to usual bipyramidal TiO₂ anatase nanoparticles, mainly exposing {101} facets. Nevertheless, such behavior depends in a significant extent on the physico-chemical features of surfaces exposed by nano-sheets. They can vary in dependence of the presence or removal of capping agents, namely fluorides, used for shape-engineering, and experimental investigations in this respect are still a few. Here we report on the evolution of interfacial/surface features of TiO₂ anatase nano-sheets with dominant {001} facets from pristine nanoparticles fluorinated both in the bulk and at their surface, to nanoparticles with F⁻ free surfaces by treatment in a basic solution and to totally F⁻ free nanoparticles by calcination at 873 K. The nanoparticles fluorine content and its subsequent evolution is determined by complementary techniques (ion chromatography, ToF-SIMS, XPS, AES, SEM-EDX), probing different depths. In parallel the evolution of the electronic properties and the Ti valence state is monitored by UV-Vis spectroscopy and XPS. The calcination treatment results in {001} facets poorly hydroxylated, hydrated and hydrophilic, which appear as surface features consequent to the expected (1x4) reconstruction. Moreover, IR spectroscopy of CO adsorbed as probe molecule indicates that the Lewis acidity of Ti⁺⁴ sites exposed on (1x4) reconstructed {001} facets of calcined TiO₂ nano-sheets is weaker than that of cationic centers on {101} facets of bipyramidal TiO₂ anatase nanoparticles. The samples have also been tested in phenol photodegradation highlighting that differences in surface hydration, hydroxylation and Lewis acidity between TiO₂ nanoparticles with nano-sheet (freed by F⁻ by calcination at 873 K) and bipyramidal shape have a strong impact on the photocatalytic activity, that is found to be quite limited for the nanoparticles mainly exposing (1x4) reconstructed {001} facets.

Keywords: shape-controlled TiO₂ nanoparticles, F⁻ doping, Ti³⁺, H₂O adsorption, CO adsorption, phenol photodegradation.

1. Introduction

In the last years semiconductor functional materials have attracted increasing attention for their widespread environmental and energy applications.¹⁻² In these fields crystal facet engineering of semiconductors nanoparticles (NPs), in particular of TiO₂, is emerging as an important strategy to finely tune their physicochemical properties to optimize surface reactivity and selectivity.³⁻⁵ For anatase TiO₂, the most widely employed semiconductor photocatalyst, the shape under equilibrium conditions predicted by the Wulff construction is a truncated tetragonal bipyramid, mainly exposing the {101} surfaces. However, starting from the seminal paper by Yang *et al.*,⁶ several researchers tried to obtain TiO₂ anatase nano-sheets (n-sh) with extended basal {001} surfaces, that have been proposed to possess superior photocatalytic activity.⁷⁻⁸ Since the {001} surfaces have a high surface energy, the preparation of TiO₂ n-sh requires the use of suitable adsorbates which can substantially lower their surface energy.⁹ In this respect, hydrofluoric acid is the most commonly employed capping agent since fluoride ions can preferentially adsorb and stabilize the {001} facets.⁹ After the hydrothermal synthesis, the surface of the anatase TiO₂ n-sh is thus terminated with Ti-F groups and F⁻ ions are expected to be present also in the bulk of the NPs. In some studies, as-prepared TiO₂ n-sh are used in photocatalytic reactions without further treatments,¹⁰ hindering the possibility to distinguish between the role of {001} surfaces and the electronic states resulting from the presence of bulk and surface fluorides. Indeed, it has been demonstrated that fluorides adsorption strongly affects interfacial hole and electron transfer in photocatalysis, as well as the extent of adsorption of substrates and O₂. This results in the modification of the mechanism and rate of substrate oxidation,¹¹⁻¹² as well as in the rate of O₂ reduction and carrier recombination.¹³ Moreover, also surface hydrophilicity is modified.¹⁴ To specifically investigate the effect of {001} facet engineering, fluorides must be thus removed and the common procedure is calcination at 873 K. Thermal treatments at lower temperatures (i.e. 773 K) were not completely effective in this respect.¹⁵ DFT calculations highlighted that such high temperature treatment should induce (1x4) reconstruction of TiO₂ (001) surfaces, that, conversely to what claimed for as-prepared TiO₂ n-sh particles, apparently leads to a weak reactivity.¹⁶ The atomic

structure and reconstruction mechanism at elevated temperatures was recently investigated by *in situ* spherical aberration corrected scanning transmission electron microscopy,¹⁷ but, at the best of our knowledge, experimental insights, at the molecular level, on the evolution of surface chemical features of anatase TiO₂ n-sh from the pristine fluorinated to the bulk and surface F⁻ free form obtained by calcination, are lacking. Moreover, the effect of the substitution of only surface fluorides of TiO₂ n-sh with OH⁻ by washing with a basic solution (of NaOH, in the present case) is also of interest.

In this respect, here we report a contribution to the elucidation of three aspects of TiO₂ n-sh in their pristine, surface F⁻ free and bulk and surface F⁻ free forms thought to be relevant for the surface and interfacial behavior of titania nanoparticles as photocatalysts: (i) hydroxylation (by mid-IR spectroscopy in controlled atmosphere); (ii) hydration (by both microgravimetry and mid- and near-IR spectroscopy of adsorbed water), which plays a significant role in photocatalytic processes at titania surfaces, not only in liquid-solid regime, but also in gas-solid regime in ambient air;¹⁸⁻¹⁹ and (iii) Lewis acidity of Ti⁴⁺ surface sites, as probed by IR spectroscopy of adsorbed CO. For the sake of comparison, truncated bipyramidal anatase TiO₂ nanoparticles, mainly exposing {101} facets,²⁰ were also considered. To strengthen the comparative analysis of the surface features of TiO₂ n-sh in dependence on the F⁻ removal treatments and with respect to truncated bipyramidal TiO₂ NPs, combined structural (by X-ray diffraction), textural (in terms of specific surface area) and morphological (by both scanning and high-resolution transmission electron microscopy, SEM and HRTEM, respectively) investigations were carried out, targeting in particular the relative amount of {001} and {101} facets exposed by TiO₂ n-sh. Indeed, the interface and relative amount of co-exposed {001} and {101} facets have been reported to crucially influence the physico-chemical and functional properties.²¹⁻²² Moreover, special care was devoted to monitor the evolution of the F⁻ content. On one hand, quantitation was pursued by complementary techniques as chemical analysis of dissolved samples and of washing solutions, and analytical inspection at different depth in the TiO₂ nanoparticles by energy X-ray dispersive spectroscopy (EDX) coupled to SEM, Auger electron spectroscopy (AES), X-ray photoelectron spectroscopy (XPS) and time-of-flight secondary ion mass spectrometry (ToF-SIMS). On the other

hand, the fading by fluoride removal of electronic states due to the presence of bulk and surface F⁻ ions was monitored by diffuse reflectance UV-Vis-NIR and mid-IR transmission spectroscopy. Moreover, the evolution of the Ti valence state has been investigated by XPS.

The differences in surface chemical features of TiO₂ nanosheets freed from both bulk and surface fluorides by calcination, mainly terminated by {001} (1x4) reconstructed surfaces, and truncated bipyramidal TiO₂ nanoparticles mainly exposing {101} facets are also discussed in terms of performance in the photocatalytic degradation of phenol, one of the most studied substrate in photocatalysis and with known degradation mechanisms.²³⁻²⁴

2. Experimental

2.1 Preparation of TiO₂ nanoparticles

TiO₂ nano-sheets. The preparation of TiO₂ nano-sheets was carried out with a solvothermal method following an established literature procedure.²⁵⁻²⁶ In a typical synthesis 25 ml of Ti(OBu)₄ (Aldrich reagent grade 97%) was poured in a 150 ml Teflon lined stainless steel reactor and 3.5 ml of concentrated hydrofluoric acid (Aldrich reagent grade 47%) was added dropwise under stirring. The reactor was sealed and kept under stirring at 523 K for 24 hours. The resulting bluish paste was centrifuged and washed with acetone to remove the residual organics and then with water (Milli-Q). Finally, the obtained aqueous suspension was freeze-dried obtaining a bluish powder. The sample in this form is labelled as TiO₂ n-sh throughout the text.

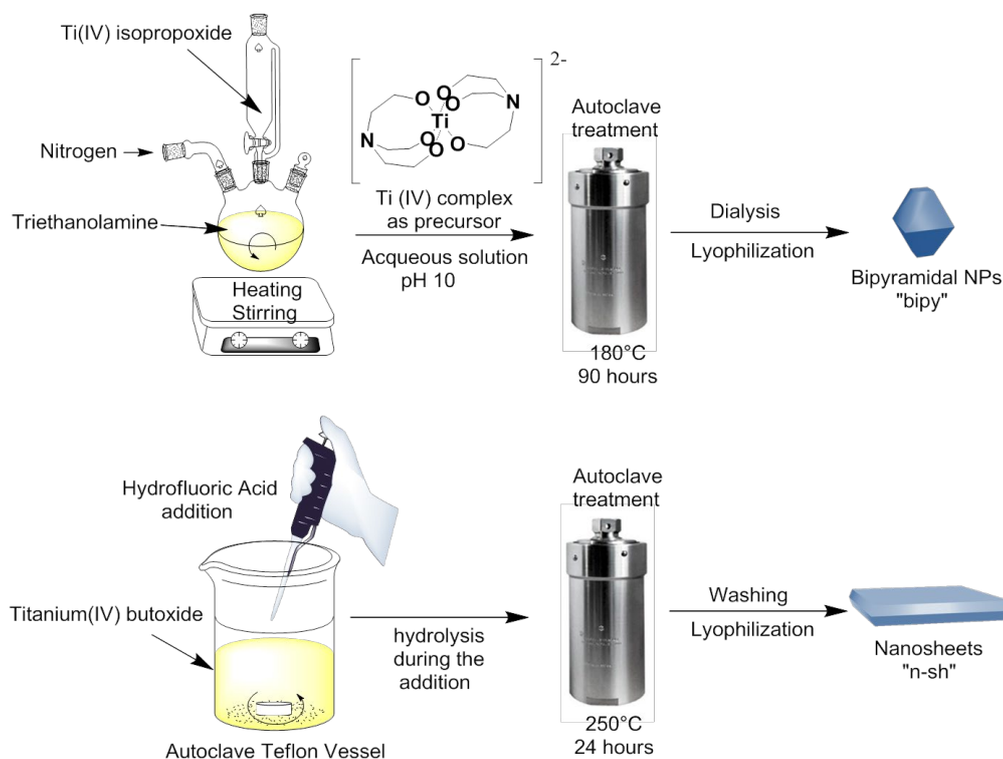
To attain the removal of the fluorides from the surface, the as-prepared TiO₂ n-sh was washed with 0.1 M NaOH (10 mg TiO₂/ml, 2 hours under stirring at ~300 K).¹⁴ The suspension was then centrifuged (5000 rpm, 15 min), and the supernatant kept for quantification of fluorides and possibly solubilized Ti species, by ion chromatography and ICP-MS, respectively. The paste of TiO₂ nanoparticles recovered by centrifugation and then again centrifuged in 0.1 M HNO₃ (40 ml, 5000 rpm, 15 min) and ultrapure water (MilliQ, Millipore; 40 ml, 5000 rpm, 15 min) to remove Na⁺ ions from the surface. The

effectiveness of this treatment was assessed by Auger electron spectroscopy and ToF-SIMS analyses (*vide infra*). The sample resulting from these treatments is hereafter referred to as TiO₂ n-sh_NaOH.

The removal of the fluorides from both the bulk and the surface of TiO₂ n-sh was achieved by calcination in air at 873 K. In a typical treatment, 200 mg of TiO₂ n-sh powder were heated up to 873 K at 6.5 K/min, kept for 60 min at the indicated temperature, and then cooled down to RT in the closed furnace in approximately 10 hours. This sample will be hereafter referred to as TiO₂ n-sh_873K.

Truncated bipyramidal TiO₂ nanoparticles. The material named TiO₂ bipy NPs were obtained by forced hydrolysis of 40 mM aqueous solution of Ti(Teoah)₂ complex (Teoah = triethanolamine; initial pH 10), carried out by hydrothermal treatment at 453 K for 90 hours in autoclave. This procedure is similar to that developed by Sugimoto, but without the intermediate gelation step.²⁷ Further details on the preparation of these NPs, hereafter referred to as TiO₂ bipy, which show a bipyramidal shape, mainly limited by {101} facets, can be found in our previous publication.²⁰

Scheme 1. Synthesis of TiO₂ nano-sheets and TiO₂ bipyramidal nanoparticles.



2.2 Characterization methods

High resolution transmission electron microscopy (HR-TEM) images of the materials (powder grains “dry” dispersed on lacey carbon Cu grids) were obtained using a JEOL 3010-UHR microscope operated at 300 kV.

Adsorption of N₂ at 77 K was exploited to measure the specific surface area of the samples with a Micromeritics ASAP 2020 sorption analyzer. Before the measurements samples were outgassed at r.t., until the attainment of a residual pressure of 1×10^{-3} mbar. Data were treated by applying the Brunauer–Emmett–Teller (BET) equation.

X-ray powder diffraction patterns of the various types of TiO₂ nanoparticles were recorded with a PANalytical X’Pert Pro powder diffractometer, equipped with an X’Celerator detector, using Cu K_α radiation generated at 45 kV and 40 mA. The 2θ range was from 10° to 80° with a step size of 0.01° and a counting time of 0.6 s per point. The 004 and 200 anatase reflections were considered to calculate the average size of crystal domains (d) along the [001] and [100] directions, respectively, by

the Scherrer equation:
$$d = \frac{K \lambda}{(B_{exp} - B_{ins}) \cos \theta}$$

where K is a shape factor (assumed to be 0.9 in the present case), λ is the X-ray wavelength, θ is the measured Bragg angle, B_{exp} and B_{ins} are the experimental and instrumental broadening, respectively. The instrumental broadening was evaluated using a Si powder standard.

IR spectroscopy in controlled atmosphere. An aliquot of each type of TiO₂ nanoparticles was pressed in self-supporting pellets (“optical density of ca. $10 \text{ mg} \cdot \text{cm}^{-2}$) and placed in quartz cells equipped with KBr windows designed to carry out spectroscopic measurements at the beam temperature (b.t.; ca. 323 K) or at low temperature (i.e. ~ 100 K) by cooling with liquid N₂.²⁰ The cells were connected to a conventional vacuum line (residual pressure: 1×10^{-5} mbar) allowing all thermal treatments and adsorption-desorption experiments to be carried out *in situ*. A Bruker IFS 28 spectrometer (resolution: 2 cm^{-1} ; detector: MCT) was employed for the spectra collection at b.t and at ca. 100 K, averaging 128 scans. H₂O and D₂O (99% D atoms, Eurisotop) were introduced onto the

samples after several freeze-pump-thaw cycles. To attain a complete H/D isotopic exchange of surface hydroxy groups and water molecules, admission (15 mbar)/outgassing (residual pressure as low as $\times 10^{-5}$ mbar) of D₂O vapor was repeated until invariance of spectra. For CO adsorption, the samples were previously outgassed at 873 K for 120 min and contacted with 20 mbar of O₂ at the same temperature. Then, the pellets were cooled to 373 K in O₂ and then cooled to r.t. under outgassing. Finally, they were cooled down to *ca.* 100 K by filling with liquid N₂ a reservoir in contact with the sample holder trough a metallic seal. For the collection of the spectra of CO adsorbed at 60 K, the self-supporting pellets were activated as described above in a properly modified Oxford CCC 1204 cryostatic chamber, inserted in a Bruker Equinox 55 spectrometer (resolution: 2 cm⁻¹; detector: MCT; 128 scans).

Diffuse reflectance (DR) UV-Vis-NIR spectra were recorded with a Cary 5000 Varian spectrophotometer equipped with an integrating sphere with an inner coating of Spectralon®. This material was used also as reference. For the collection of spectra, thick self-supported pellets of TiO₂ nanoparticles were placed in a cell with an optical quartz window designed to carry out the measurements in controlled atmosphere, by connection to a conventional vacuum line (as for IR spectroscopy). For the sake of clarity, the spectra are reported only in the 380-2500 nm range, starting, on the short wavelength side, from the onset of the edge-shaped inter-gap absorption typical of TiO₂ anatase,²⁸ extended towards the UV, which dominates the spectral profile by far.

Determination of fluoride by ion chromatography. In order to evaluate the total amount of fluorides a complete mineralization in microwave was performed heating the samples for 90 min at 473 K under vigorous stirring in NaOH 10 M. After the treatment, a dilution 1:10 of the solution was analyzed by Ion Chromatography (IC). The instrument (DIONEX DX 500) is equipped with a GP40 pump, AS9-HC column (Dionex, length 25 cm and internal diameter of 4 mm), Suppressor ASRS-Ultra II autosuppression mode, electrochemical detection system ED40, LC30 column thermostat (temperature 303 K). The mobile phase consists of 9 mM K₂CO₃ and the eluent flow is equal to 1 mL min⁻¹.

Determination of soluble Ti species by ICP-MS. Inductively Coupled Plasma – Mass Spectrometry (ICP-MS) analyses were carried out using a Thermo Scientific ICP-MS ICAP-Qs model, equipped with a quadrupole mass analyzer and a flatpole quadrupole collision/reaction cell. The instrument is calibrated with 1% nitric acid standard for Ti prepared by diluting standard reference certificates at 1000 mg L⁻¹ using ⁴⁵Sc, ⁸⁹Y, ¹⁵⁹Tb (100 ppb) as internal standards. The calibration curve was constructed with 5 points and linear correlation coefficient > 0.99. Interference due to polyatomic ions is eliminated by operating the collision cell in He mode with kinetic energy discrimination (He - KED). ⁴⁸Ti was used for quantitation.

SEM/EDX. For SEM and EDX measurements aluminum holders were used, which were cleaned by sonication in ethanol and subsequently sample powder was pressed on the Al-holder with spatula. High-resolution electron micrographs and EDX spectra have been collected with a SEM of type Zeiss Supra 40 (Carl Zeiss, Oberkochen, Germany), equipped with a Schottky field emitter and having attached a silicon drift detector energy dispersive X-ray spectrometer (SDD-EDS) from Thermo Fisher Scientific (energy resolution at Mn K α of 128 eV and 100 mm² crystal active area). SEM images were acquired operating the microscope at 10 kV.

Auger electron spectroscopy (AES). The powder was deposited on a cleaned Si-wafer (sonicated in isopropanol) by slightly pressing the powder with a spatula on the wafer surface. AES analysis was carried out with a PHI 700 Auger Scanning Probe (ULVAC-PHI Inc.) equipped with a coaxial cylindrical mirror analyzer. Auger electrons spectra were excited by the primary electron beam at different locations on the sample surface. To evaluate the presence of Fluorine on the TiO₂ nanoparticles surface, AES survey spectra as well as window spectra of energy region corresponding to F KLL were recorded (5 keV, 20 nA and 20 keV, 1 nA).

X-ray photoelectron spectroscopy (XPS). The XPS investigations were performed with a KRATOS Axis Ultra DLD. Monochromated Al K α with an energy of 1486.6 eV was used for the excitation of the photo- and Auger electrons. The binding energy scale of the spectra were referred to C1s at 285 eV. Quantitative analysis was performed with Casa XPS software package using the background-

corrected area of the main peaks of the elements and sensitivity factors and the spectrometer-specific transmission function. The peaks were fitted with a Gaussian-Lorentzian profile.

Time-of-flight secondary ion mass spectrometry (ToF-SIMS). Samples were prepared on silver foil as well as on silicon wafers by slightly pressing the powder with a spatula on the wafer surface. The substrates were cleaned by sonication in isopropanol prior to sample deposition. ToF-SIMS measurements were performed on a ToF-SIMS IV instrument (ION-TOF GmbH) of the reflectron type, equipped with a 25 keV bismuth liquid metal ion gun as a primary ion source.

Photocatalytic activity assessment. The photocatalytic activity of TiO₂ was evaluated by irradiating the suspensions of titania nanoparticles (loading 100 mg l⁻¹; pH=6.5) in cylindrical Pyrex glass cells under magnetic stirring (4 cm diameter, 2.5 cm height, area of 1.26 x 10⁻³ m², cut-off 295 nm, 5 ml suspension volume), using a fluorescent source with $\lambda_{\text{max}} = 365$ nm (Philips PL-S 9W BLB, integrated irradiance = 10 W m⁻²). Substrate absorption at this wavelength is negligible. The use of this wavelength assures minimal impact of nanoparticle agglomeration on photoactivity during the test.²⁴ Moreover, to mitigate the influence of large agglomerates formation during the experiment, the irradiation time was limited to less than one half-life of phenol. Photocatalytic activity was measured as the initial rate of phenol phototransformation, by fitting disappearance curves to an exponential decay. Incident radiant power was measured in the range 290-400 nm with an Oceans Optics USB2000 spectrophotometer equipped with a cosine corrected optical fiber probe, spectroradiometrically calibrated with a NIST traceable DH-2000 CAL UV-Vis source (Ocean Optics). The initial concentration of phenol is 0.1 mM. Time profiles of phenol decay were obtained as average of three irradiation runs.

HPLC determination of phenol has been carried out with an Agilent Technologies HPLC chromatograph 1200 Series equipped with a diode array detector, binary gradient high-pressure pump and an automatic sampler. Isocratic elution was carried out with a mixture of 15/85 acetonitrile/formic acid aqueous solution (0.05 % w/v), flow rate 0.5 ml min⁻¹, injection volume 20 μ l. The column used was a Kinetex C18 150-2 (150 mm length, 2 mm I.D., 2.6 μ m core-shell particles, Phenomenex).

3. Results and discussion

3.1 Structural and morphological characterization of the TiO₂ nanoparticles

Figure 1A shows a representative SEM image of the as-prepared TiO₂ n-sh, appearing as sheets 75 ± 25 nm wide in the basal plane and 9 ± 1.6 nm thick (see Table 1). In HR-TEM images (Figure 1B), nanoparticles properly oriented with respect to the impinging electron beam show lattice fringes 0.2370 nm apart ($d_{004} = 0.2378$ nm, ICDD PDF 00-021-1272) parallel to the lateral view of the basal facets, and related FT (inset) confirm that these facets are normal to the *c*-axis (reported in the image) and thus correspond to {001} surfaces. The lateral boundaries of the nanoparticle imaged in Figure 1B form an angle of $\sim 130^\circ$. This is compatible with the projection of intersecting (101) and (10 $\bar{1}$) surfaces for a NP with the *a*-axis forming a small angle with the image plane since the angle between intersecting (101) and (10 $\bar{1}$) surfaces should be 136° , considering a tetragonal unit cell with lattice parameters $a = 3.78$ Å and $c = 9.51$ Å. The TiO₂ n-sh_NaOH sample, resulting from the washing of TiO₂ n-sh in a NaOH solution to remove surface fluorides shows a similar morphology (see Figure 1C and Figure S1 in the Supporting Information), that, conversely, was almost lost by subsequent calcination at 873 K (Figure 1D). Therefore, the calcined TiO₂ n-sh_NaOH will no longer be considered. Noteworthy, direct calcination of TiO₂ n-sh does not lead to significant morphological changes (sample TiO₂ n-sh_873K), as from both SEM and HR-TEM imaging (Figure 1E and 1F, respectively).

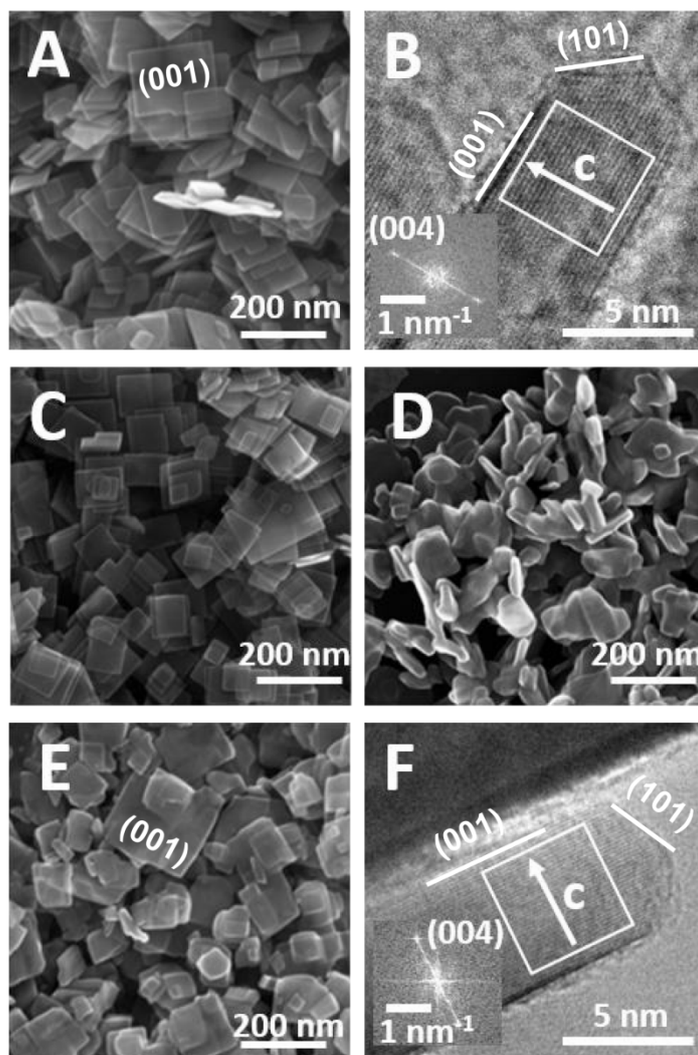


Figure 1. SEM (panels A,C,D,E) and HR-TEM (panels B,F; inset: FT of the image portion within the with frame) images of: A,B) as-prepared TiO₂ n-sh; C) TiO₂ n-sh _NaOH; D) TiO₂ n-sh _NaOH calcined at 873 K; E,F) TiO₂ n-sh _873 K.

Figure 2A shows a representative SEM image of the as-prepared TiO₂ bipy, which appear as truncated bipyramids 30 ± 5 nm wide in the basal plane and 40 ± 9 nm thick along the *c*-axis (see Table 1). In the HR-TEM image (Figure 2B) we can see in greater detail a single particle with well-defined borders, which appears as a bidimensional projection of a truncated bipyramid with the *c*-axis almost perpendicular to the electron beam. From the SEM and HR-TEM images reported in Figure 2C-D we can see that calcination at 873 K does not induce significant morphological changes.

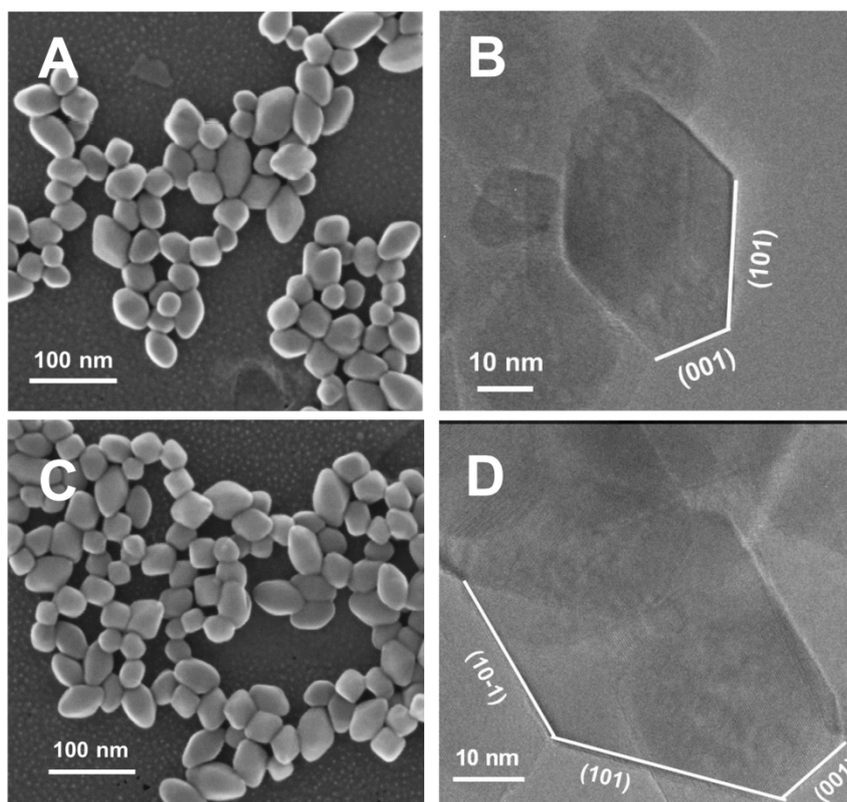


Figure 2. SEM (panels A, C) and HR-TEM (panels B, D) images of: A,B) TiO₂ bipy and C,D) TiO₂ bipy_873 K.

All samples were found to be pure anatase TiO₂ phase by XRD (Figure S2). No significant variations of lattice parameters due to the incorporation of F⁻ were observed. The Scherrer analysis, applied as an integral method, highlights that the average crystallite dimension in the [001] direction is approximately doubled for TiO₂ n-sh_873K with respect to TiO₂ n-sh and TiO₂ n-sh_NaOH (Table 1), suggesting that the primary nanosheets might have aggregated in pairs along the *c*-axis, as schematized in Figure S3. This stacking through basal facets should be the responsible for the decrease of the specific surface area (SSA_{BET}) from 53 to 34 m²/g, because no significant change in size and shape of primary nanoparticles was observed neither in SEM nor in HR-TEM images. Conversely, very limited changes in crystal size along the [001] direction and in SSA_{BET} occurred for TiO₂ bipy when passing from the pristine to the calcined at 873 K form, in agreement with the electron microscopy results (Figure 2). The pairing of nano-sheet particles, which has been already proposed in previous studies,²⁹

should have a relevant impact on the relative amount of lateral {101} and basal {001} surfaces of TiO₂ n-sh_873K. By a geometric modeling of nano-sheet pairing (details in Figure S4 and related comment), the {101}:{001} ratio was estimated to change on an average from 20:80 for TiO₂ n-sh to 40:60 for TiO₂ n-sh_873K (Table 1). The resulting relative change in calculated specific surface area was in excellent agreement with the experimental one (see SI for details).

Table 1. Average dimension of crystal domains obtained by Scherrer analysis of the 004 and 200 XRD peaks, particle thickness (T) and basal length (L) measured from SEM images, specific surface area and relative change, calculated average percentage of {001} facets and relative change in SSA (see SI for calculation details).

Sample	d ₀₀₄ (nm)	d ₂₀₀ (nm)	T (nm)	L (nm)	SSA _{BET}		calculated	
					(m ² /g)	relative change	%{001}	SSA relative change
n-sh	18	> 100	9.4 ± 1.6	75 ± 25	53	1	78	1
n-sh_NaOH	21	> 100	8.4 ± 1.7	73 ± 27	57	~1	79	1
n-sh_873K	46	> 100	10 ± 2	64 ± 25	34	0.64	58	0.62
bipy	36	29	40 ± 9	30 ± 5	43	1	10	-
bipy_873K	32	26	35 ± 5	45 ± 9	37	0.86	13	-

3.2 Determination of the F⁻ content

The fluorine content and its subsequent evolution were determined by complementary techniques (Table 2), also probing different depths (details in Figure S5 and related comment). By ion chromatography, pristine TiO₂ n-sh is found to contain 16 mg F⁻/g TiO₂, decreasing to 7 mg F⁻/g TiO₂ for TiO₂ n-sh_NaOH, where ion-exchange is expected to remove fluorides only from the surface. The absence of significant dissolution of surface layers was confirmed by the trace amount of titanium found in the supernatant, corresponding to less than 0.015 atoms of Ti for nm² of TiO₂. As a result, an initial F⁻ surface concentration of ~5 F⁻/nm² is calculated. This surface fluorination is relevant since the Ti⁴⁺ density is 7 and 5 nm⁻² on {001} and {101} surfaces, respectively.³⁰ The significant decrease in

the ToF-SIMS signal of F⁻ in the mass spectra of TiO₂ n-sh_NaOH sample, normalized to the TiO₂⁻ intensity, and in the XPS F1s spectra, normalized to the Ti2p signal, (see Table 2) supports partial removal of fluorine as found with IC, i.e. by ‘losing’ F⁻ ions from the outermost surface of nano-sheets particles by ion exchange. Note that AES (probing 3-4 nm depth, i.e. reaching the bulk of the NPs which are ~ 9 nm thick) as well as EDX (probing ~1 μm depth) cannot distinguish between the levels of fluorine signals as detected in the two samples TiO₂ n-sh and TiO₂ n-sh_NaOH (details in the SI). By combining these results, one can thus infer that TiO₂ n-sh_NaOH does not contain F⁻ ions on its surfaces, while they are still present in the bulk (Figure S3). A reliable quantification into absolute chemical composition of fluorine with all four techniques, ToF-SIMS, XPS, AES and EDX is impossible mainly due to the strong morphological effects typical for nanopowders (heterogeneous distribution of solids/voids, heterogeneous orientation of nanoparticles, see SI).³¹

All methods agreed in indicating the removal of fluorine in the nano-sheets calcined at 873 K down to or below the detection level of each method (see Table 2), from both the bulk and the surface.

Table 2. Fluoride concentration for the different samples obtained by ion chromatography, and fluorine signals as detected by ToF-SIMS, XPS, AES and EDX.

Sample	IC	ToF-SIMS	XPS	AES	EDX
	(mg F/g TiO ₂)	I _F -/I _{TiO₂} - (a. u.)	I _F /I _{Ti} (a. u.)	I _F KLL (a. u.)	I _F Kα (a. u.)
n-sh	16.1 ± 1.2 (5 F ⁻ /nm ²)*	73 ± 6	0.20 ± 0.04	Yes	Yes
n-sh_NaOH	7.0 ± 0.8	54 ± 4	0.05 ± 0.02	Yes	Yes
n-sh_873K	< 0.4	3.6 ± 0.4**	0.03 ± 0.01	≅LoD	≅LoD
bipy	< 0.4	5.7 ± 0.6**	≅LoD***	≅LoD	≅LoD

* F⁻ surface density, see text for details

**This concentration level, different from zero also for the originally F-free TiO₂ bipy, was found even in the Si (100) blank substrate used for analysis, demonstrating that this level of F⁻ represents the background contamination present at the ToF-SIMS instrument used. For detailed explanation of the results in Table 2 please consult the SI (sections EDX, AES, XPS and ToF-SIMS). LoD= limit of detection

***At this low level F is detectable, but not reliably quantifiable, see also SI

The calcination process has also a strong impact on the electronic properties of the pristine F⁻ containing nanoparticles. Indeed, the electronic spectrum of TiO₂ n-sh (Figure 3A) shows a very broad electronic absorption spread over the 600-2500 nm range, due to trapped excess electrons resulting from the substitution of O²⁻ ions by F⁻ ions.³²⁻³⁴ Most part of this absorption is still present in the spectrum of TiO₂ n-sh_NaOH. Moreover, in the MIR spectral range (Figure 3B) these samples show a spectral profile resulting from the superimposition of a continuous increase in absorbance, from 2500 up to 10000 nm (spectral cutoff due to bulk Ti–O vibrations), and of a series of minor localized signals, vibrational in nature. The latter will be commented on in the following, while the first one is typical of free conduction band and shallow-trapped electrons.^{10, 35-36} After calcination, the spectral features in the overall 600-10000 nm (panels A+B of Figure 3) due to electronic defects associated with the presence of F⁻ disappear, confirming that fluorine has been completely removed.

We performed also a parallel XPS analysis to investigate the evolution of the valence states. In the Ti2p XPS spectra reported in Figure S10 we can identify a specific component of the Ti⁴⁺2p_{3/2} at 459.9 eV ascribed to the influence of the F⁻ anions.³⁷⁻³⁸ The intensity ratio between this Ti⁴⁺ component and the main Ti⁴⁺2p_{3/2} signal decreased from 0.3 for TiO₂ n-sh to *ca.* 0.1 for the other samples (see SI for the details), in agreement with the evolution of the electronic properties highlighted by UV-Vis-NIR-MIR spectroscopy and with the progressive decrease of F⁻ content (see Table 2). A contribution of Ti³⁺ to the Ti2p signal can be also present, especially for the TiO₂ n-sh sample (see SI).³⁹

Finally, for the bare TiO₂ nanoparticles we calculated the band gap using a Tauc plot (Figure S11) to evaluate the impact of the different exposed facets. The results are 3.10 ± 0.07 eV for TiO₂ bipy and 3.26 ± 0.06 eV for TiO₂ n-sh_873K. These data seem to be in contradiction with experimental results on micrometric crystals⁴⁰ and with DFT calculations which reported a smaller band gap for the {001} facets. However, it should be considered that the particle thickness for TiO₂ n-sh_873K is below 10 nm, while it is about 40 nm for TiO₂ bipy (see Table 1), therefore the higher band gap for nano-sheet particles could be also due to quantum confinement effects which have been reported to play a role for anatase nanoparticles up to 15 nm in diameter.⁴¹ Anyhow, these small differences in band gap should

not impact our tests of the photocatalytic activity which are performed under pure UV irradiation at 3.4 eV (see section 3.5).

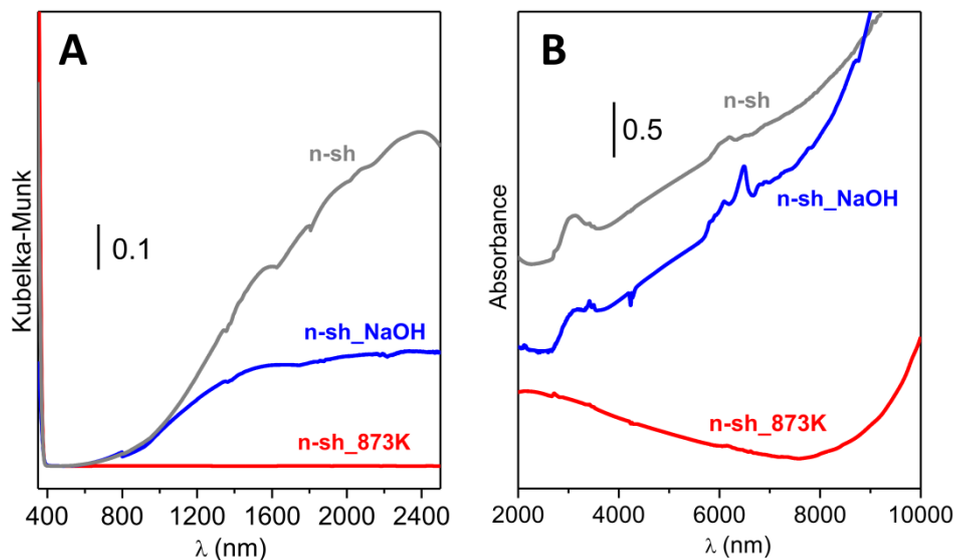


Figure 3. Diffuse reflectance spectra, in the 380-2500 nm range and in Kubelka-Munk units (A), and transmission NIR-MIR spectra, in the 2000-10000 nm range and in absorbance units (B) of TiO₂ n-sh, TiO₂ n-sh_NaOH and TiO₂ n-sh_873K. On the X-axis of panel B, where spectra in the MIR range are shown, wavelength values are reported, instead than conventional wavenumber values, for the sake of consistency with spectra in panel A (being wavelength values commonly used for electronic spectra in the UV-Vis-NIR ranges). Note on the MIR spectrum of TiO₂ n-sh_873K: the apparent decrease in absorbance from 8000 to 2000 nm is the typical effect of decreased scattering of the samples at higher wavelengths, observed when measuring the spectrum in transmission, instead of in diffuse reflectance.

3.3 Surface hydration and hydroxylation

The interaction between water and fluorinated (TiO₂ n-sh) and F⁻ free (TiO₂ n-sh_NaOH and TiO₂ n-sh_873K) was investigated by measuring the amount of H₂O molecules that can be reversibly adsorbed at room temperature (r.t) (Figure 4A). This amount depends on the adsorptive properties of the “first hydroxylation/hydration layers”, constituted by surface hydroxy groups and strongly adsorbed water which are not desorbed by outgassing at r.t. (*vide infra*). Pristine TiO₂ n-sh adsorbs up to ~8 H₂O molecules/nm² when in contact with H₂O at 15 mbar. The removal of F⁻ by the two different methods impacts in two opposite directions the surface hydrophilicity: for TiO₂ n-sh_NaOH, the substitution of surface F⁻ by exchange with OH⁻ results in an increase to ~12 H₂O molecules/nm², whereas for calcined TiO₂ n-sh_873K the water adsorption capacity becomes limited to only ~5 H₂O

molecules/nm². This finding is in agreement with DFT calculations which suggested that in the terrace region of the (1×4) reconstructed (001) surface, which should be obtained after calcination, water is physisorbed or not adsorbed at all.¹⁶ It is worth mentioning that TiO₂ bipy, overwhelmingly exposing {101} surfaces, adsorbs up to ~10 H₂O molecules/nm². For TiO₂ n-sh_873K, where these facets account on an average for 40% of the surface, the actual amount of H₂O molecules adsorbed on fluorine free calcined {001} terminations might be of the order of ~2 H₂O molecules/nm². A similar consideration holds for TiO₂ n-sh_NaOH, which exposes on average 20% of {101} facets, but in this case the F⁻ free unreconstructed {001} surface exhibits an affinity towards water similar or slightly higher than the {101}. No definitive conclusions can be derived for the pristine TiO₂ n-sh since F⁻ can modify also the properties of {101} surfaces.¹⁴

The amount of H₂O per nm² reversibly adsorbed on TiO₂ bipy in contact with H₂O at 15 mbar might correspond to the formation of 2 molecular layers, adsorbed on the first hydration layer, assuming an average density of ~5.3 molecules/nm² for each of the three layers, on the basis of molecular dynamic calculations of H₂O on a TiO₂ anatase (101) surface,⁴²⁻⁴³ (details on the calculations in the SI). For H₂O on (001) TiO₂ anatase, a radial distribution function was found only for an unreconstructed hydroxy-free surface,⁴² resulting in an average density of ~9.3 molecules/nm² per each of the first three layers (see Figure S12). This result cannot be compared with experimental results on neither TiO₂ n-sh_NaOH nor TiO₂ n-sh_873K, where {001} surfaces are expected to be hydroxylated or (1×4) reconstructed, respectively. Nevertheless, an additional insight on the state of water on TiO₂ n-sh_873K in comparison with TiO₂ bipy was obtained by NIR spectroscopy (Figure 4B). The $\nu_{\text{asym}} + \delta(\text{H}_2\text{O})$ band observed for TiO₂ bipy in equilibrium with 15 mbar H₂O (*i.e.* ~3 molecular layers of adsorbed water) is composed by a main absorption at ~5180 cm⁻¹, and a shoulder at ~5300 cm⁻¹, this latter due to the H-bonding free OH pointing towards the exterior of the upmost water layer.⁴⁴⁻⁴⁵ This component accounts for ~3.7% of the overall integrated intensity, but this value increases to ~8.5% for H₂O on TiO₂ n-sh_873K. In agreement with the microgravimetric data, this feature should be related to

a decrease of the amount of water molecules underneath the upmost layer, i.e. a decreased number of H₂O layers on TiO₂ n-sh_873K.

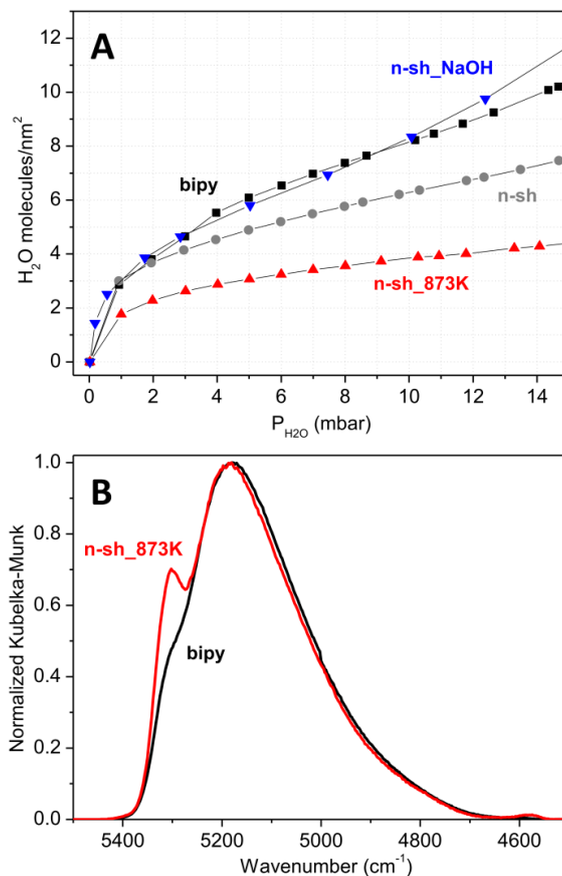


Figure 4. (A) Water adsorption isotherms at 298 K obtained from microgravimetric measurements. The samples have been previously outgassed at r.t. (B) Normalized NIR spectra of the TiO₂ bipy (black) and TiO₂ n-sh_873 K (red) samples in contact with H₂O at 15 mbar.

Information on the relative amount of water molecules left irreversibly adsorbed at r.t. on the various samples and on their hydroxylation state was provided by MIR spectroscopy. As a link with microgravimetric data (providing information on H₂O reversibly adsorbed at r.t.), also spectra of samples in contact with H₂O at 15 mbar were collected (Figure 5A). Relevant spectral features are: i) a weak, sharp band at ~ 3700 cm⁻¹ due to the $\nu(\text{OH})$ of H-bonding free H₂O molecules at the surface of the upmost water layer (see above), ii) an intense and broad absorption spread over the 3600-2800 cm⁻¹ range, due to the $\nu(\text{OH})$ of H-bonded water molecules and surface hydroxy groups, iii) the $\delta(\text{H}_2\text{O})$

signal of adsorbed H₂O molecules, with maximum at ~1635 cm⁻¹.⁴⁶⁻⁴⁷ In addition, also the $\delta(\text{H}_2\text{O})$ of water left adsorbed irreversibly by outgassing at r.t. is reported, for the sake of comparison (dashed lines). For each sample, the integral of the area between the full and dotted lines is proportional to the amount of reversibly adsorbed H₂O molecules. The relative values obtained (1.0/bipy, 0.7/n-sh, 0.9/n-sh_NaOH, 0.3/n-sh_873K) are in good agreement with those provided by microgravimetry (1.0/bipy, 0.7/n-sh, 1.1/n-sh_NaOH, 0.4/n-sh_873K). Thus, also the relative integrated intensity of the $\delta(\text{H}_2\text{O})$ signals obtained for samples outgassed at r.t. (Figure 5B) should correspond to the relative amount of H₂O molecules left adsorbed in this condition, that was not possible to determine by microgravimetry. With respect to the pristine TiO₂ n-sh, the removal of fluorides by NaOH washing resulted in an increased capability of retaining water in the first hydration layer, becoming similar to that of TiO₂ bipy. Moreover, the upshift in position of the $\delta(\text{H}_2\text{O})$ band indicates that a change in the interaction of water molecules with surface sites occurred, their behavior as H-bonding donors gaining in relative extent with respect to the interaction through the oxygen atom.⁴⁸⁻⁴⁹ Noteworthy, the amount of water molecules retained by the surface of TiO₂ n-sh calcined at 873 K appears rather low. This well agrees with DFT calculations which suggested that, on the reconstructed (001) surface in the terrace region, water is physisorbed or not adsorbed at all.¹⁶ Concerning the 3800-2700 cm⁻¹ $\nu(\text{OH})$ spectral profile of samples outgassed at r.t., TiO₂ bipy exhibits the well-known pattern made of a series of sharp bands at $\nu \geq 3600$ cm⁻¹ due to isolated hydroxy groups, a band at 3470 cm⁻¹ assigned to water molecules strongly coordinated with Ti⁴⁺ sites on {101}, overlapped to the onset of a broad absorption due to an extended network of hydrogen bonding between OH groups and water molecules.⁴⁶⁻⁴⁷ This absorption becomes the dominant component of the $\nu(\text{OH})$ profile of TiO₂ n-sh, at the expenses of signals due to isolated hydroxy groups. These signals are totally absent in the spectrum of TiO₂ n-sh_NaOH, indicating that the treatment in basic solution affected also the hydroxylation of {101} surfaces. Conversely, bands due to isolated OH are the main feature of the $\nu(\text{OH})$ pattern of TiO₂ n-sh_873K, characterized by an overall weak intensity.

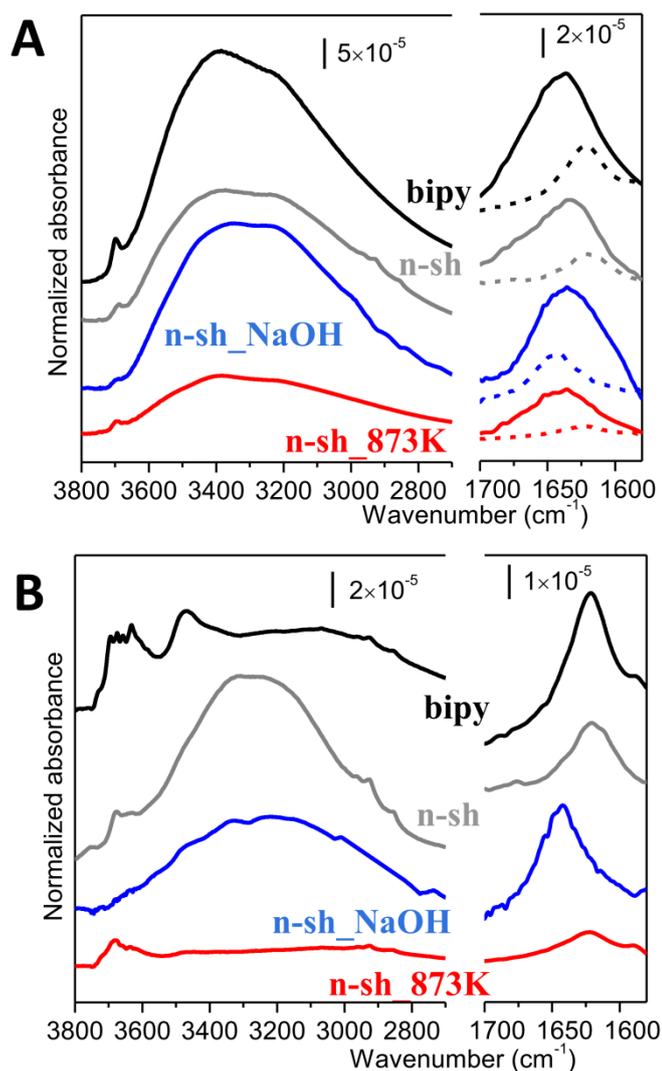


Figure 5. FT-IR spectra of the samples in contact with H₂O at 15 mbar (A) and after outgassing for 1 hour at r.t. (B). To facilitate the comparison, the spectra after outgassing for 1 hour are also reported in part (A) as dashed curves in the H₂O bending region. The spectral features of surface hydroxy groups and water molecules were extracted from original spectra as shown in Figure S14. The spectra here reported have also been normalized by dividing the absorbance by the specific surface area (m²/g) and the “optical thickness” (g/m²) of the pelletized samples.

3.4 Surface Lewis acidity

The Lewis acidity of surface Ti⁴⁺ sites was probed by IR spectroscopy of adsorbed CO.⁵⁰⁻⁵¹ This method requires the pre-outgassing at high temperature and, therefore, was applied only to TiO₂ n-sh_873K, that already experienced a thermal treatment, and to TiO₂ bipy, for the sake of comparison. Surprisingly, the main feature of the spectrum of CO adsorbed at ~100 K on TiO₂ n-sh_873K is the peak at ~2180 cm⁻¹ due to probe molecules on {101} surfaces,^{20, 52} similar in

position to the dominant peak of CO on TiO₂ bipy, whereas signals associated to {001} surfaces are not noticeable (Figure 6A). Details on the assignment of the other minor components are reported in Table S1 in the Supporting Information. By further decreasing the adsorption temperature down to ~60 K (Figure 6B) the dominant spectral feature of CO on TiO₂ n-sh_873K becomes a new band at ~2155 cm⁻¹, thus falling in a frequency range expected for CO adsorbed on Ti⁴⁺ sites on (1x4) reconstructed (001) surface as predicted by DFT calculations.²⁰ This band is fully reversible upon CO outgassing at 60 K, whereas the peak at ~2180 cm⁻¹ remains almost unaffected (appearing reversible upon outgassing in the experiment at 100 K). Moreover, the maximum moves from ~2155 to ~2160 cm⁻¹, because of the fading away of adsorbate-adsorbate interactions, i.e. approaching the singleton ν(CO).⁵³ The very limited upshift of the singleton signal with respect to the stretching mode of CO in gas phase (2143 cm⁻¹) indicates that Ti⁴⁺ on such surfaces are rather weak as Lewis acid sites. This agrees with the weakness of their interaction with CO, actually scarce at a temperature as low as ~100 K.

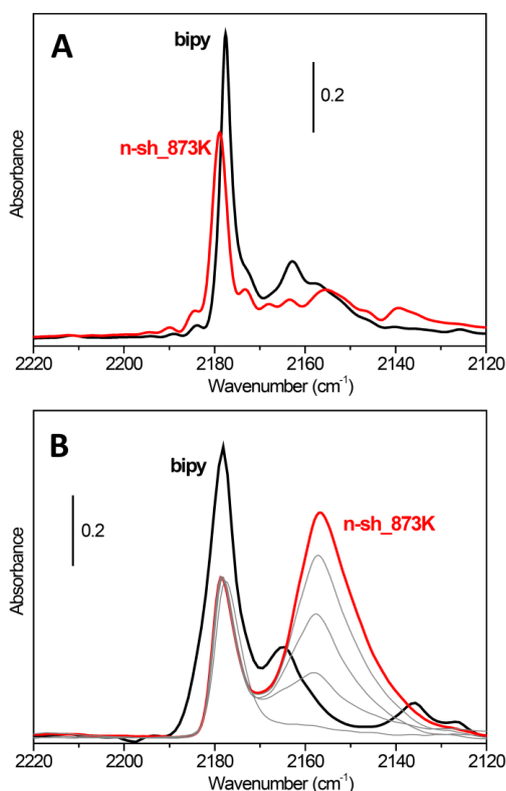


Figure 6. IR spectra of CO adsorbed on TiO₂ bipy (black) and TiO₂ n-sh_873 K (red) at 100 K (A) and 60 K (B). In both cases, the sample was put in equilibrium with CO at 15 mbar at 100 K and then, for (B), the temperature was decreased to 60 K. The grey spectra in part (B) refer to progressive CO outgassing at 60 K for sample TiO₂ n-sh_873 K.

3.5 Photocatalytic activity

To investigate the impact of the differences in surface features between bare TiO₂ nanoparticles mainly terminated by {101} or (1x4) reconstructed {001} surfaces, TiO₂ bipy and TiO₂ n-sh_873K were tested in the phenol photodegradation reaction. The mechanism of phenol degradation has been widely studied demonstrating that for pure TiO₂ under UV irradiation ·OH radicals are the primary active species.²³ Recent studies highlighted the crucial role of both hydroxyls and adsorbed water in stabilizing the surface hole centers.⁵⁴ Moreover, also the amount of water adsorbed at the photocatalyst surface was found to strongly influence the efficiency of the phenol degradation process.³⁶ Thus, hydroxylation and surface affinity towards H₂O molecules are expected to have both a relevant influence on this process. For the sake of completeness, tests were carried out also for pristine TiO₂ n-sh and TiO₂ n-sh_NaOH, although the photocatalytic behavior of these nanoparticles should be affected also by the electronic states due to the presence of fluorides. Finally, we performed a test to further check that the thermal treatment at 873 K does not change the functional properties of TiO₂ bipy: in agreement with the morphological stability, already discussed in section 3.1, the photocatalytic activity does not significantly change between TiO₂ bipy and TiO₂ bipy_873K (see Figure S15).

Figure 7 shows the phenol photodegradation curves (panel A) and the related initial degradation rates (panel B), normalized by weight and specific surface area of the samples of nanoparticles. The effectiveness of TiO₂ n-sh_873 (red curve) in the phenol photodegradation is very limited, whereas TiO₂ bipy (black curve) appears significantly more active (*ca.* -6% and -23% of C₆H₅OH, respectively, after 180 min irradiation). Noteworthy, TiO₂ bipy has an initial degradation rate *ca.* 4.5 times higher than TiO₂ n-sh_873 K. Thus, the collection of data on surface features of these two types of nanoparticles allows to state that low hydrophilicity, poor hydroxylation, and weak Lewis acidity of Ti sites exhibited by reconstructed {001} surfaces affect negatively the photocatalytic performance of bare TiO₂ nano-sheet particles in the tested reaction with respect to the TiO₂ bipy ones, mainly exposing {101} facets.

As far as the other two forms of titania nano-sheet particles, TiO₂ n-sh and TiO₂ n-sh_NaOH, are concerned, they act as better photocatalysts in phenol degradation than TiO₂ n-sh_873K, in terms of both C₆H₅OH conversion along the duration of the test and initial rate. Interestingly, TiO₂ n-sh_NaOH nanoparticles, with surfaces freed from F⁻, richer in hydroxy groups and more hydrophilic than pristine TiO₂ n-sh ones, are the most active, approaching the performance of TiO₂ bipy.

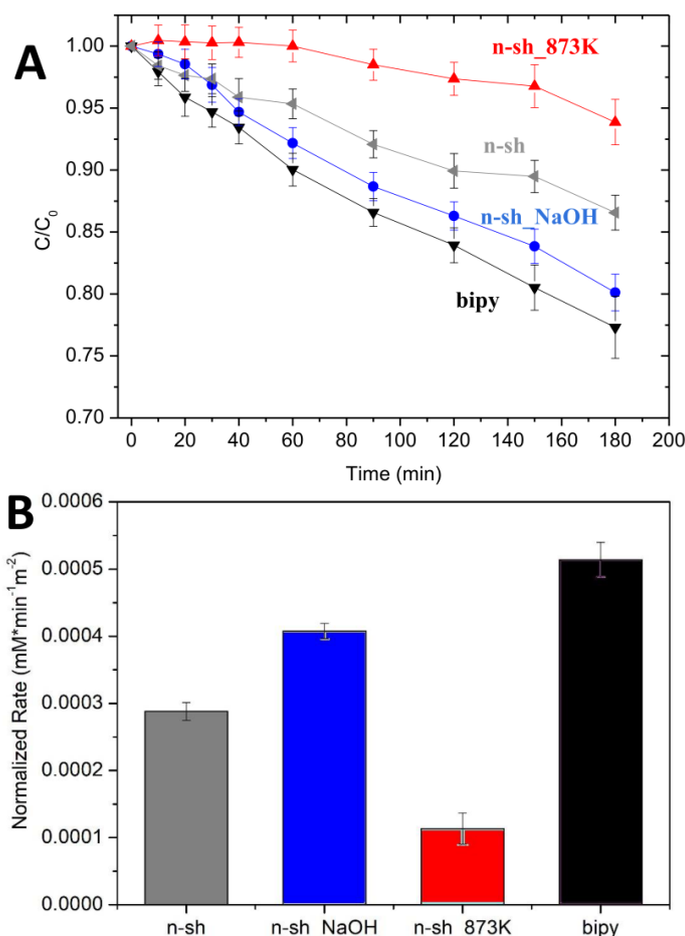


Figure 7. Phenol photodegradation curves (A) and related initial degradation rates, normalized by weight and specific surface area, (B) obtained using the various types of TiO₂ NPs.

4. Conclusions

The comprehensive set of data presented in this work leads to a main conclusion: TiO₂ nano-sheets exposing {001} facets as prevalent surfaces, prepared by capping with fluorides, can appear as particles similar in shape, but they actually show significant differences in surface/interfacial features, namely hydroxylation and hydrophilicity, in dependence on the treatment carried out (if any) for the

removal of F⁻ ions, which remain in the bulk and on the surface. Thus, other than obvious differences in electronic states due to these anions, also these aspects should be considered when evaluating functional properties involving surface/interfacial phenomena, as photocatalysis. Interestingly, a proper treatment in a basic solution of TiO₂ nano-sheets templated with fluorides allows to attain a complete exchange of surface F⁻ with OH⁻, resulting in nanoparticles retaining the platy shape, and combining a high surface hydroxylation with the electronic states due to fluorides still retained in their bulk, as highlighted by XPS and UV-Vis spectroscopy. TiO₂ nano-sheets that underwent this treatment show an increase of 40% in normalized initial rate of phenol photodegradation with respect to pristine TiO₂ nano-sheets fluorinated both in the bulk and at the surface, which are also less hydroxylated and hydrophilic. Conversely, complete F⁻ elimination by calcination at 873 K results in a decrease of 60% in normalized initial rate with respect to pristine nanoparticles. The disappearance of electronic states due to F⁻ is accompanied by a significant decrease in surface hydroxylation and hydration, likely resulting from a (1x4) reconstruction of {001} facets.

Calcined TiO₂ nano-sheets, being free from both bulk and surface F⁻, appear as the most suitable of the three types of platy-shaped TiO₂ nanoparticles, considered in this work, to be compared in a straightforward way with non-doped TiO₂ nanoparticles with different morphology. In this respect, (1x4) reconstructed {001} facets, mainly exposed by calcined TiO₂ nano-sheets, show significant lower hydrophilicity, poorer hydroxylation, and weaker Lewis acidity of Ti sites than {101} facets, mainly exposed by bipyramidal TiO₂ nanoparticles. These differences in surface features are accompanied by a normalized initial rate of phenol photodegradation 4.5 times lower for F⁻-free calcined TiO₂ nano-sheets.

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Supporting Information. Details of the model adopted to calculate the expected specific surface area and percentage of exposed {001} surfaces; HR-TEM images of TiO₂ n-sh_NaOH; SEM/EDX, XPS, AES and ToF-SIMS data; Tauc plots; complete FT-IR spectra of the samples after outgassing for 1 hour at r.t.; phenol photodegradation curves for TiO₂ bipy and TiO₂ bipy_873K samples are available free of charge on the ACS Publications website.

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Anatase TiO_2 nanoparticles

