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# Opposite effect of the Cu photodeposition on the •OH generation and diclofenac degradation depends on the exposed anatase facet

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### ABSTRACT

Exposition of a specific crystal facets and modification with co-catalysts are often used in order to improve photocatalytic activity of a material. However, considering complexity of these interactions, it is still challenging to fully understand and predict activity of a specific system. Here, we report combination of the {0 0 1} and {1 0 1} crystal facets of anatase TiO<sub>2</sub> with Cu<sub>x</sub>O co-catalyst, applied for diclofenac degradation, •OH generation and 4-nitrophenol reduction. Interestingly, we observed opposite effect of Cu for •OH generation and DCF degradation, with clear enhancement of the activity observed only for the {0 0 1} facets. EPR measurements further proved that for the most active material in this series, Cu exist as an atomically-scattered Cu<sup>2+</sup> species at the surface, achieved as the result of high surface development and high adsorption energy (from DFT). Based on the absorbance measurements and XPS analysis, these species are shown to be further reduced to Cu<sub>2</sub>O during degradation process. Compared to this, all other systems ({1 0 1} and all results for nitrophenol) showed reduction of the activity due to the Cu<sub>x</sub>O presence. In this regard, Cu<sub>x</sub>O cannot be seen as an universal reduction co-catalyst for TiO<sub>2</sub>, with final effect being clearly dependent on exposed facet and reaction. This is despite TiO<sub>2</sub>/Cu<sub>x</sub>O was found to form analogical, well-known Z-scheme junction in case of both surfaces. Ultimately, we showed that specifically {0 0 1} with Cu<sub>x</sub>O make highly synergistic combination for ROS generation.

### 1. Introduction

Photocatalytic degradation of organic pollutants is a promising strategy for enrironemntal remediation, which ideally does not require additional energy input apart from the sunlight. During the process, photocatalyst is not consumed, and the process takes place with the participation of solar radiation. However, due to the specificity of the solar light spectrum and the fact that the most commonly used photocatalysts, with TiO<sub>2</sub> at the forefront, show maximum absorption in the UV range, photocatalyst design is crucial regarding efficient utilization of incident light.

In this regard, multiple approaches are presented in order to improve photocatalyst performance. For example, surface modification with noble metals such as gold (Au), platinum (Pt), or silver (Ag) are a popular way to improve photoactivity. In such systems, deposition of the metal nanoparticles on the photocatalyst surface create heterojunctions capable of increased light absorption, mostly due to the effect of surface plasmon resonance (SPR) [1]. Occurrence of the SPR on metal surface creates possibility for electron transfer into the conducting band of TiO<sub>2</sub>, therefore increasing its photoactivity and additionally extending time of electron-hole recombination as the result of the metal–semiconductor junction (Shottky junction) [2,3]. However, using noble metals as co-catalysts increase overall cost of the materials, leading to the situation where other metal species gather increasing attention as an alternative to Au/Ag/Pt. From these possibilities, modification with copper became a promising way to increase TiO<sub>2</sub> activity, especially under continuous irradiation in the visible and UVA range, due to the formation of the Z-scheme junction between the TiO<sub>2</sub> and Cu<sub>X</sub>O species that forms as the result of partial copper oxidation at the TiO<sub>2</sub> surface [4].

However, despite rather large interest in the  $TiO_2/Cu$  system in the photocatalysis, some aspects of their operation still remain unknown. Particularly, this connects with interactions between Cu and  $TiO_2$ 

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surface features, such as a crystal facet engineering, which lately became important topic in the photocatalyst design [5–7]. Due to the anisotropy of the electronic properties in bulk crystal, termination of its structure with different crystal facets can heavily affect final properties of the material. This includes e.g. surface work function, band positions, surface energy and density of surface undercoordinated sites, which all affect further localization of excited charge carriers and interactions with surrounding species [8,9]. In case of the anatase  $TiO_2$ , its {0 0 1} amd {1 0 1} terminations are known to possess fundamentally different electronic properties, due to the unique electronic structure of the (0 0 1) crystal plane [10]. As the consequence, electron localization is known to be hindered for the {0 0 1}, while its favored on the surface atoms of the {1 0 1}. In this regard, both these surfaces are intriguing materials for further modifications with co-catalyst. For example, theoretical predications of the {101} termination with different metal nanoclusters were made concerning H<sub>2</sub> generation [11]. For such application, some interest in the {0 0 1} facets could also be noted, especially after their modification with Pt [12–14]. However, since it is still challenging to predict how fundament properties (e.g. change in work function) will affect efficiency of the specific photocatalytic process, it is still needed to systematically increase our knowledge about application of crystal facets in their bare and modified form for a different photocatalytic reactions. Concerning modification of different TiO<sub>2</sub> crystal facets with Cu, existing results report some of the possible applications. For example, Wei et al. studied Cu modification of anatase particles with either {1 0 1} or the combination of {0 0 1} with {1 0 1} facets exposed in the reaction of acetic acid decomposition [15]. However, more complex organic compounds were not investigated, as well as information about generation of reactive species in such system is generally missing. Also, nanoparticles that would exposed majority of the {0 0 1} were not investigated.

In this regard, we have especially focused on this problem and compared interactions between photodeposited Cu and  $\{0 \ 0 \ 1\}$  or  $\{1 \ 0 \ 1\}$  crystal facets of anatase TiO<sub>2</sub>. Modified and bare materials were studied for the degradation of diclofenac, which is one of the most frequently used non-steroidal anti-inflammatory drug (NSAID) [16,17]. Due to its popularity and ineffective treatment processes in wastewater treatment plants (3 to 60 % removal, depending on the method), it is commonly found in surface waters [18], inducing negative impact on aquatic organisms such as algae, crustaceans, and fishes [19–23]. Furthermore, generation of  $\bullet$ OH radicals by the prepared materials was studied in detail and discussed with respect to the electronic structure of the exposed anatase facet, followed by the 4-nitrophenol reduction.

### 2. Experimental

Detailed experimental description can be found in SI, Section 1.

### 2.1. Preparation of the $TiO_2$

Anatase nanoparticles with either {0 0 1} or {1 0 1} facets exposed in majority were prepared solvothermally based on the previous reports [24–26]. Briefly, for the {0 0 1} exposition, titanium n-butoxide (TBT) was mixed with the ethanol and concentrated HF solution and reacted at 180 °C for 10 h (please, be cautious that HF solutions are extremely dangerous and appropriate precautions must be used during work e.g. neoprene gloves and calcium gluconate gel are recommended for hand protection and neutralization after possible skin exposure). For the dominant exposition of the {1 0 1} facets, commercial P25 was hydrothermally treated with KOH solution to obtain  $K_2Ti_6O_{13}$  precursor. Obtained titanate was H-exchanged using HNO<sub>3</sub>, washed with water and dried. Finally,  $H_XK_2$ -xTi<sub>6</sub>O<sub>13</sub> was transformed into octahedral TiO<sub>2</sub> by reacting at 210 °C for 18 h with ammonia buffer. Final products were washed, neutralized with NaOH (in case of {0 0 1}), dried and hand-grounded.

### 2.2. Cu photodeposition

The photodeposition process was carried out in a quartz reactor using  $CuSO_4$  as the Cu source and 50 cm<sup>3</sup> of 50/50 v/v ethanol–water mixture as environment. The suspension was stirred and purged for 30 min with nitrogen, closed tightly and 1 h with a 300 W Xe lamp. Modified material was washed multiple times and dried.

### 2.3. Structural characterization

Crystal structure of the prepared samples was analyzed using powder X-ray diffraction (XRD) with the Rigaku MiniFlex diffractometer and the Cu Ka radiation source. The refinement of the pattern performed with the available.cif file of the anatase structure (COD ID 9015929). The absorption properties of the powders were analyzed using ThermoFisher Scientific Evolution 2000 spectrometer (0.5 nm resolution) with the BaSO<sub>4</sub> as a standard for the diffuse reflectance spectroscopy measurements (DR-UV/vis). For the measurements of the suspensions we have followed procedure suggested by Cabrera et al. The measurements were performed between the 300–800 nm, with the 10 nm step. Surface areas were determined from the BET adsorption isotherm of N<sub>2</sub> at 77 K, after degassing of the sample under N<sub>2</sub> flow at 200 °C for 2 h. X-ray photoelectron spectroscopy (XPS) measurements were performed using a PHI VersaProbeII Scanning XPS system with monochromatic Al Ka (1486.6 eV) X-rays All XPS spectra were charge referenced to the unfunctionalized, saturated carbon (C-C) C1s peak at 285.0 eV. Transmission electron microscope images were collected using TEM TECNAI TF 20 X-TWIN microscope by FEI, operating at 200 kV, which is equipped with field emission gun. Photoluminescence spectra were recorded using Shimadzu RF-6000 spectrofluorometer with 300 nm and the  $\lambda$  < 400 nm cut-off filter on the emission side.

### 2.4. Local volume rate of photon absorption (LVRPA) modeling

The local volume rate of photon absorption (LVRPA) distribution within the photocatalytic rector was estimated with the six-flux model (SFM [27,28]), using procedures described by Acosta-Herazo et al. [29]. Detailed procedure is described in SI.

### 2.5. Photocatalytic studies and analytical procedures

Evaluation of photocatalytic activity was carried out in a cylindrical reactor with a water jacket cooling, airflow and a quartz glass as the UV–vis entry. We used a 300 W xenon lamp (LOT Oriel, Darmstadt, Germany) with both IR and  $\lambda < 310$  nm radiation filters as the light source. The UVA light intensity maximum on the reactor surface was adjusted as 24 mW•cm<sup>-2</sup>. DCF solution with a concentration of 20 mg•dm<sup>-3</sup> or 500 µmol•dm<sup>-3</sup> coumarin solution were used for the photocatalytic studies. Analysis of the DCF and its by-products was performed with HPLC detection (MS and DAD), while formation of 7-hydroxycoumarin was determined via its fluorescence. Estimation of errors associated with the adapted procedures was based on the repeated processes for the selected cases. The highest observed error was than adapted for the remaining experiments. Detailed results are shown in SI, section 5.

### 2.6. Computational details

Molecular properties of the proposed diclofenac by-products were simulated using Orca 5.0.4 software [30] with the B3LYP functional, aug-cc-PVDZ basis set and CPCM model of solvation. All structures were geometry-optimized using BFGS algorithm to the standard convergence criteria Absorption spectra were simulated using time-dependent density functional theory (TD-DFT). The energy difference between diclofenac and its byproducts was finally calculated using H<sub>2</sub>O, O<sub>2</sub> and H<sub>2</sub> energies.

Interaction between TiO<sub>2</sub> surfaces and Cu atoms were additionally investigated using Quantum Espresso 7.2 software package [31] with PAW pseudopotentials. Energy cut-offs for wavefunction and charge density were 40 Ry and 400 Ry respectively. We used 3x3x1 and 3x2x1slab models with a single k-point for {0 0 1} and {1 0 1}surfaces respectively however, to reduce computational cost their thickness was reduced approximately by half. Single Cu atom was introduced on each surface to study its adsorption and resulting DOS distribution.

### 3. Results and discussion

## 3.1. Structure of the unmodified $TiO_2$ with dominant exposition of {0 0 1} and {1 0 1} facets

Both prepared TiO<sub>2</sub> samples were found to be single-phase anatase powders, without other signals observed in the XRD patterns, as presented in Fig. 1a, including Rietveld refinement of the anatase crystal structure ( $\chi^2 = 2.322$  for the {0 0 1} exposition and 2.332 for the {1 0 1} one). Despite this, both patterns show significant differences in the overall character. In the case of the {0 0 1} exposition, selective broadening of the signals with high values of the l index is clearly observed, e.g. for the (0 0 4), (1 0 5) and (1 1 6) planes, which agrees with the expected 2-dimensional character of the obtained particles [25]. Although it is known that precise determination of the crystallite size from XRD patterns is a challenging problem [32], it can be estimated that their corresponding sizes along the [0 0 1] direction are approximately 16 nm for the {0 0 1} and 400 nm for the {1 0 1}, which gives over 20 times difference between both materials. Furthermore, highly anisotropic growth of the {0 0 1} exposing sample resulted in a slight increase of the unit cell volume, compared to the {1 0 1} sample, as well as relatively lower crystallinity of the structure, which can be noticed due to lower intensity of the overall pattern. Both these observations are in logical agreement with the small crystallite size along the l direction for the {0 0 1} exposing sample, since it is known that perfect bulk crystal structure is commonly not conserved near the surface. In this regard, XRD characteristics confirm expected morphological differences for the  $\{0 \ 0 \ 1\}$  exposing sample.

Following crystal structure analysis, both materials were

investigated spectroscopically to check their absorbance properties and chemical states at the surface. Fig. 1b presents the results of the DR-UV/ vis measurements with bandgap determination using the Tauc method in the inset. As observed, both materials show absorbance spectra characteristic for anatase TiO<sub>2</sub>, with the main electron transition between the valence and conduction band occurring at approximately 400 nm. Interestingly, compared to the  $\{1 \ 0 \ 1\}$ , the bandgap of the  $\{0 \ 0 \ 1\}$ exposing sample is slightly reduced by  $\sim 0.1$  eV with the final value of 3.16 eV. This effect is contradictory to the common observations that the bandgap of different semiconductors is increasing for smaller nanostructures [33,34]. In this regard, it is noteworthy that previous reports have concluded that for the isotropic anatase nanoparticles, analogical bandgap variation with size is almost negligible [35]. On the other hand, DFT-based simulation of the (0 0 1) model has previously suggested the existence of the additional electronic states on this specific surface, which could result in the bandgap narrowing [36]. Analogical increased activity in the visible light region was also suggested for the {0 0 1} facets in the photooxidation of formic acid [37]. In this regard, smaller bandgap is expected to result from the specific structure of the  $\{0 \ 0 \ 1\}$ facets, rather than size effects or impurities. In the case of the  $\{1 \ 0 \ 1\}$ sample, the determined bandgap is in a range of commonly reported values at room temperature.

Finally, acquired high-resolution XPS spectra in the Ti 2p, O 1 s and C 1 s regions after the fitting procedure are shown in Fig. 1c. As can be seen, in both cases, Ti exists solely as the Ti<sup>4+</sup> species, in accordance with the stoichiometric TiO2 structure. Particularly, no reduction of the surface can be noticed. This is in agreement with the amount of the lattice oxygen observed in the O 1 s spectra for both samples (O-Me at approx. 530 eV), which give O/Ti ratios equal 1.99 for the {001} exposing sample and 2.13 for the {101} one, as well as the absorbance results which have not indicated any absorption increase in the NIR region [38]. Another signals observed in the O 1 s spectrum show only moderate differences between both materials and generally follow the characteristics of the C1 s signal (that is, higher O1 s signal ascribed to the mixed O-C and -OH species in case of the {101} sample correlates with the higher O-C signal in the C1 s spectrum as well). Finally, in the case of the {001} sample, the signal corresponding to the adsorbed fluorine (~684 eV) was noticed with the final Ti/F ratio of



Fig. 1. Rietveld-refined XRD patterns of the prepared  $TiO_2$  samples (a), together with the corresponding absorbance spectra (b) and deconvoluted high-resolution XPS signals for the observed elements (c). Inset of (b) presents bandgap determination using Tauc method for the indirect transition.

approximately 4. This is fluorine residue left after the preparation, which is commonly observed in similar materials [39,40]. Besides this, other differences result most likely from the adsorbed impurities and the oxide surface itself in both cases is almost identical before the modification with Cu.

Fig. 2 shows TEM/HRTEM images of the prepared samples. As presented in Fig. 2a, in the case of the HF-assisted synthesis, the 2-dimensional morphology of the nanoparticles is clearly observed. The HRTEM analysis of the selected plates (Fig. 2b) revealed their singlecrystalline nature, as confirmed by the discrete diffraction pattern obtained from the fast Fourier transform (FFT). Moreover, despite a slightly twisted orientation of the particles, these images confirmed expected orientation of the crystal structure, with symmetrical patterns observed in both directions towards the edges of the plates, in accordance with the fourfold symmetry of the anatase unit cell along its *c* axis (visualization of the approximated orientation of a crystal structure is presented in Fig. 2b). Moreover, in the case of the {1 0 1} exposing nanoparticles, their octahedral morphology can be easily observed (Fig. 2c), in accordance with the 8 equivalent (1 0 1) planes exposed. Analogically, HR-TEM observations confirmed their single-crystalline nature, with the FFT diffraction pattern corresponding to the [1 0 0] zone axis of the anatase crystal structure when analyzing isolated octahedron from its side (Fig. 2d). Based on these results, the successful preparation of both materials was clearly confirmed. Finally, structural characterization was completed with the control measurements of the specific surface areas of both materials using the BET method. These analyses revealed the high difference in the surface development, being either 122 or 13  $m^2 \cdot g^{-1}$  for the {0 0 1} and {1 0 1}, respectively. This difference results from the adapted preparation procedures, needed to stabilize specific facet, where condensation of TBT (strictly bottom-up process) allows to achieve high surface development. On the other hand, transformation of the solid precursor used for the preparation of the {1 0 1}, P25 and K-titanate, allows only for a limited control over surface development.

## 3.2. LVRPA analysis and diclofenac degradation over unmodified samples

Following the large difference in the specific surface area of both materials, optical properties of their suspensions were investigated, in order to further compare their activity under similar illumination of the reaction system. Fig. 3a and b show trends of the average measured absorbance between 300–380 nm, including either the light transmitted directly through the suspension (a) or both light transmitted directly and after scattering on the particles (b). Based on the obtained data, spectral average extinction ( $\beta$ ) and absorption ( $\kappa$ ) coefficients were determined as the slope of these trends, with correction on the optical path in both cases (L). The difference between these values results from the light scattering on the particles. As presented, the absorption coefficient of

both materials is fairly similar, however, particles with high  $\{0 \ 0 \ 1\}$ exposition show visibly higher  $\beta$  values, which indicate larger scattering in this system. This difference is reasonable, since it is well known that light scattering is a complex process, affected by many factors such as particle shape, size and possible agglomeration [41]. In this regard, the platelet shape of the {0 0 1}-exposing particles and different surface potential of the (0 0 1) plane might be a source of increased scattering. Following this observation, the apparent optical thickness ( $\tau_{app}$ ) of the utilized photoreactor was calculated for different loadings of both TiO2 structures, as presented in Fig. 3c. Ultimately, the TiO<sub>2</sub> concentrations of  $0.4 \text{ g} \cdot \text{dm}^{-3}$  and  $0.5 \text{ g} \cdot \text{dm}^{-3}$  were selected respectively for the {0 0 1} and {1 0 1} exposing structures, which corresponds to the  $\tau_{app}$  of approximately 7. Ideally, the  $\tau_{app}=1$  value corresponds to the minimum photocatalyst loading required for total absorption of the incident light and therefore should result in the most uniform irradiation of the whole reactor space. However, a very low number of dispersed particles also reduces the available surface sites for the reaction to occur. Therefore, maximum activities are commonly observed for higher  $\tau_{\text{app}}$  values, which represents the compromise between a large volume of the reactor being effectively irradiated and a large number of active sites being "free" for reaction. Noteworthy, different values of  $\tau_{app}$ , such as a 3–4, are often suggested as optimal for different TiO<sub>2</sub> materials [28,29], although, it should be noted that photocatalytic processes are known to be sensitive to the technical details of the experimental setup and in this regard, we have previously observed that reduction of the TiO<sub>2</sub> concentration below  $\sim 0.5 \text{ g} \cdot \text{dm}^{-3}$  significantly decrease final activity in our case [9]. Therefore,  $\tau_{app} = 7$  was adapted for further studies. As presented in Fig. 3d, both materials show almost identical course of the LVRPA curve under such conditions. After applying detailed distribution of the UVA light that reaches the reactor border (Fig. 3e), this results in approximately 60 % of the reactor cross-section being effectively irradiated in both cases (Fig. 3f with limit adapted as the 1 % of maximum LVRPA value).

Following LVRPA analysis, the photodegradation of diclofenac (DCF) was studied under the selected photocatalyst concentrations and compared to the spontaneous photolysis (Xe lamp as the light source with an additional 310 nm filter to cut-off high energy UV radiation). As presented in Fig. 4a, photodegradation of DCF under such conditions was quite efficient, with approximately 85 % degradation observed after 60 min of irradiation. As presented in Fig. 4c, photolysis of DCF results in the formation of multiple by-products characteristic of this process, which are marked as P1-P4. Further combination of MS analysis (spectra in SI, Fig. 1 section 2), DAD spectra and molecular simulations revealed that these structures are mostly carbazole derivatives, formed due to the initial dehydrohalogenation of the DCF molecule (product P1) with subsequent hydroxylation and oxygenation of the benzene rings (P2-P4). As presented in Fig. 4d, the accumulation of these products over time shows different kinetics, supporting step-by-step transformation mechanism of these compounds. During this process, product P4 might



**Fig. 2.** Exemplary TEM images of the prepared  $TiO_2$  samples, including nanoparticles with dominant {0 0 1} exposition (a), selected HRTEM image of the individual nanoplate (b), nanoparticles with dominant {1 0 1} exposition (c) and selected HRTEM images of the individual octahedron (d). The HRTEM images are accompanied by the FFT transformation, that corresponds to the diffraction pattern, as well as visualization of the approximated orientation of the anatase crystal structure for the {0 0 1} exposition. Scalebars are 200 nm for the panels (a) and (c); and 20 nm for the (b) and (d).

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Fig. 3. Determination of spectral average extinction  $\beta$  (a) and absorption  $\kappa$  coefficients (b) of the suspensions in the light range of 300–380 nm, as the slope of the linear absorbance vs. concentration trend. The calculated apparent optical thickness  $\tau_{app}$  of the suspension as the function of the photocatalyst concentration (c) and detailed LVRPA curve for the selected  $\tau_{app}$  conditions at the reactor center (d). The UVA flux distribution at the border of reactor used for the photocatalytic experiments (e) and comparison of the LVRPA distribution in the full reactor cross-section for both TiO<sub>2</sub> materials, colored between 100–1 % of the maximum LVRPA value presented in panel d (f).

be seen as particularly interesting, due to the clear appearance of the unique  $\sim 420$  nm absorption band. Excessive simulation of the possible structures suggests that such visible-light absorption is strictly connected with the transformation of the amine moiety (–NH-) to the imine one (=N-) within the DCF structure (Fig. 4g), which must also involve substitution of the benzene rings with keto-oxygen. Ultimately, we propose a detailed P4 structure, as presented in Fig. 4e. Noteworthy, the transformation between the P3 and P4 closely resembles this between the hydroquinone and benzoquinone, described in detail for the phenol degradation.

Compared to the simple photolysis, the introduction of the TiO<sub>2</sub> slightly improves overall DCF removal, however both mechanism and the kinetics of this process differ significantly. In the case of the {0 0 1} exposing structure, the adsorption during in-dark stabilization of the system achieved nearly 90 %, of initial DCF concentration, which was followed by selective formation of only one degradation by-product in the suspension (P5), as presented in Fig. 4a and 4d. Both MS data and absorption spectra clearly identify the P5 as hydroxylated DCF, without additional changes to the parent molecule. On the other hand, the {1 0 1} exposing sample shows negligible adsorption ability with a clearly improved DCF degradation after light introduction. Interestingly, in this case, both P1 and P2 products were also observed, independently of the preferred P5 formation. Ultimately, MS analysis has indicated the formation of the additional product P6 in this case, with the observed m/z (+) = 310.0. The possible structure of this product is presented in

Fig. 4e, however it should be noted that, similarly to the P4, such structure should exhibit absorption at approx. 520 nm, which was not observed in the DAD spectra, possibly due to its low concentration. Noteworthy, as shown in Fig. 4f, DFT simulations clearly suggest that hydroxylated products (P5 and P3) are particularly stable, which might be additionally responsible for the preferred P5 formation in the TiO<sub>2</sub> presence. Moreover, further transformation of P5 to P6 is clearly endothermic, which also supports its possibly low concentration and absence in the DAD spectra. Optimized models of the simulated molecules are presented in Fig. S2. Finally, we will note that TOC measurements indicated increased mineralization efficiency of the TiO2-assisted degradation than simple photolysis (approx. 40 % for the {1 0 1}, 23 % for the  $\{0\ 0\ 1\}$  and  $1\ \%$  for photolysis). This shows that despite efficient dehydrohalogenation, UVA alone is not effective for the diclofenac mineralization which was also observed for other compounds [42]. Detailed TOC results are presented in SI (Fig. S3).

Ultimately, the effect of both  $\text{TiO}_2$  presence and its exposed facets were clearly observed in the DCF degradation kinetics and mechanism. Particularly, significant adsorption and selective formation of P5 were observed in the case of the {0 0 1} facets. Noteworthy, both these observations are closely related. Firstly, high energy of the clean {0 0 1} facets [43] should stimulate DCF adsorption, creating a complex between undercoordinated 5c-Ti atoms and  $-\text{COO}^-$  group [44–46] (please note that during the degradation in water, DCF is in its deprotonated salt form, but became protonated again during the chromatographic



**Fig. 4.** Observed changes in the diclofenac concentration (a) due to the UVA photolysis and TiO<sub>2</sub>-assisted photocatalytic degradation, determined I-order kinetics of the process (b), exemplary DAD chromatograms of the photolysis/photocatalysis process (c), observed accumulation of the selected products over time (d), suggested mechanism of the initial DCF transformation in both processes (e), energy diagram of both initial degradation paths (f) and simulated absorbance spectra of the selected products (g).

analyses due to the acidic separation conditions). Therefore, since almost all DCF became adsorbed before the irradiation, formation of the P1-P4 products became practically eliminated. However, the high adsorption energy might also stabilize the DCF/DCF-OH at the surface, which is not released back to the environment after excitation, leading to the inhibition of the degradation rate for longer irradiation times. On the other hand, the low energy {1 0 1} facets do not show significant indark adsorption, leading to the situation where all transformations and adsorption start to occur only after introduction of light. Therefore, mixed photolysis of the free molecules and photocatalytic degradation of photoadsorbed molecules are occurring simultaneously, creating both dehydrohalogenated products P1 and P2, as well as hydroxylated P5.

### 3.3. Structure and properties of the Cu modified samples

Following the above results, clear effects of the TiO<sub>2</sub> presence and its surface structure on the DCF degradation were observed, including the particular role of the in-dark adsorption and UV-induced dehydrohalogenation process. To investigate further possible control over degradation efficiency and its detailed pathway, both TiO<sub>2</sub> materials were surface-modified with Cu clusters/nanoparticles. Copper species are well-known to easily deposit on anatase surfaces under illumination [47], effectively alternating their photocatalytic activity. Due to the low cost of copper, this approach presents a promising and affordable method for TiO<sub>2</sub> modification. However, the detailed effect of the Cu species with respect to the exposed facet has been less investigated compared to isotropic particles, and the impact of the resulting interactions on the degradation process of complex compounds like DCF remains relatively unknown. In this regard, we have studied thoroughly these aspects.

Ultimately, both TiO<sub>2</sub> materials were modified with nominal Cu amounts of 0.5 %, 1 %, 3 % and 5 % (atomic), which were photodeposited under the Xe lamp irradiation from the CuSO<sub>4</sub> solution in water/ethanol mixture. In every case, the irradiated solution became pale-purple during the deposition, as presented in Fig. 5a, indicating the formation of the reduced Cu<sup>0</sup> species, with characteristic absorption maximum at approximately 580 nm due to the localized surface plasmon resonance effect (LSPR) [48]. However, after drying, the optical properties of the prepared materials underwent further changes in the span of the next day with a clear shift of the absorption maximum to the higher wavelengths, as presented in Fig. 5b, especially for the lower Cu loadings. Similar maxima were reported previously for the TiO2/Cu materials and resulted from the oxidation of Cu to the Cu<sub>X</sub>O species at the surface [49,50]. Noteworthy, the new maxima at 783 nm and 688 nm are noticeably shifted between both surfaces, indicating the formation of a slightly different band structure in each case, resulting reasonably from the different electronic properties of both surfaces [23,36]. Interestingly, for the Cu loading of 3 % and 5 %, the overall increase of the absorbance in the broad visible light spectrum is clearly observed. This connects with the appearance of metallic copper in the



**Fig. 5.** Exemplary images of the sample color change during the photodeposition and after storing for a few days, with additional absorbance spectrum measured strictly after initial drying of the sample (a); the absorbance spectra of all modified samples after drying and storing (b), corresponding XRD patterns focused on the Cu presence (c), optimized models of the Cu adsorption on both facets with corresponding adsorption energy (d) and calculated DOS distribution for both systems (e).

XRD patterns, as presented in Fig. 5c. The significantly lower signal of the (1 1 1) Cu plane in case of the 5 % modification of the {0 0 1} facets agrees with the high surface development and surface energy of this structure [51], which might improve the dispersion of the Cu species during the deposition. To get better insight into Cu adsorption on both facets, DFT simulations were performed for the periodic slab models of both surfaces with single Cu atom adsorbed. As presented in Fig. 5c, the geometry of Cu adsorption in both cases involve bonding with the 2 of undercoordinated O atoms at the surface. The resulting adsorption energy is exothermic with the high preference of Cu bonding at the  $\{0 \ 0 \ 1\}$ over the  $\{1 \ 0 \ 1\}$  surface (-4.85 eV vs -2.08 eV respectively). Therefore, better dispersion of the Cu species on the {0 0 1} might be expected both due to the high surface area development and high adsorption energy of Cu. Finally, simulated DOS patterns, presented in Fig. 5e, revealed very analogical structure with occupied Cu states adopting energy levels with the bandgap. However, it can be noted that the bandgap of  $\{0 \ 0 \ 1\}$  is visibly reduced, comparing to the  $\{1 \ 0 \ 1\}$ , which is a known to result from the unique electronic structure of the  $(0\ 0\ 1)$  plane. As the result, Cu states are closer to the CB edge, which might be responsible for the maximum shift observed in the absorbance spectra.

Detailed analysis of the surface presence of Cu and its chemical states were further investigated using XPS spectroscopy. As presented in Fig. 6a, the Cu  $2p_{3/2}$  signal can be observed in each case around 932.7 eV, however for the lowest concentrations of 0.5 % at. it is approximately at the detection limit. Further increase of the Cu loading clearly enhances its intensity especially for the {1 0 1} samples, which connects well with the smaller surface area of this material, leading to a visibly higher-than-designed Cu concentrations at the surface (surface Cu/Ti are between 0.9 % and 21 % for this series vs. 0.4 % and 7.5 % for the {0 0 1}). No clear difference in the oxidation state was noticed between the

samples, with a single line fitted for the overall signal corresponding to the  $\mbox{Cu}^0$  or  $\mbox{Cu}^+$  oxidation [52].

Outside of the Cu presence, XPS studies have revealed rather similar characteristics between the samples, with detailed results presented in SI (Figs. S4 and S5). However, an increased contribution of the O1 s signal at ~531.4 eV can be noted for the higher concentrations of Cu (Fig. S5). This signal might origin from the increased presence of oxygen vacancies in the TiO<sub>2</sub> structure, which might affect its photocatalytic activity. In this regard, XPS studies were followed with EPR measurements in order to investigate the formation of possible structural defects. As presented in Fig. 6b, both unmodified materials were found to be relatively defect-free. Specifically, the only signal at g = 2.0034 was noted for the {101}, which could be ascribed to the hole trapped at the oxygen site [53]. However, concentration of this species is very low, being a few orders of magnitude lower than introduced Cu. Starting from this point, copper introduction results in the appearance of the clear EPR signal around g = 2.1205, which is characteristic for the Cu<sup>2+</sup> ions [54]. Moreover, in the case of the {0 0 1} with 1 % Cu, the hyperfine structure of the Cu<sup>2+</sup> was clearly observed with 4 characteristic signals around g = 2.3895 [55,56]. This proves that in this case, Cu exist as the atomically-dispersed species on the surface. For other samples, analogical spectra were not observed and the g = 2.389 signals start to overlap due to the increased dipole-dipole interactions, connected with the reduced distance between neighboring Cu<sup>2+</sup> centers. In this regard, EPR measurements additionally confirmed better dispersion of the Cu species on the  $\{0 \ 0 \ 1\}$  samples. Interestingly, since only  $Cu^{2+}$  can be observed during the EPR measurements, the obtained results stands in some contrast to the XPS, where reduced species were observed. In this regard, we estimated possible concentration of the Cu<sup>2+</sup> ions from the EPR measurements and compared it to the total amount of introduced



**Fig. 6.** Deconvolution of the Cu  $2p_{3/2}$  XPS signals observed for all modified samples (a), together with the EPR measurements for the bare, 1 % and 5 % modified samples in both series (b) and Cu<sup>2+</sup> concentration estimated from the EPR measurements, compared to the total amount of introduced Cu (c).

copper. As presented in Fig. 6c, nearly all of the introduced copper might exist as the Cu<sup>2+</sup> in the case of the {0 0 1} 1 % sample. For other samples however, the amount of Cu<sup>2+</sup> is not proportional to the overall amount of copper. Based on combined results so far, we conclude that Cu became reduced during the deposition process, and further appearance of the

 $Cu^{2+}$  resulted from the subsequent oxidation in air. Moreover, in the case of the {0 0 1} facets, high surface area and adsorption energy promote atomic dispersion of the Cu species, which facilities their effective oxidation to  $Cu^{2+}$  observed in the EPR spectra. However, both increasing Cu content and changing the material to {1 0 1}, with lower



Fig. 7. HR-TEM images of the 0.5 and 5 % Cu-modified samples. Yellow lines highlight exemplary surface irregularities, resulting from the (possible) Cu presence. Scale bars are 10 nm. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

surface area and weaker interactions with Cu, leads to aggregation of Cu species and stabilize  $Cu^0/Cu^+$  in the final material. Ultimately, we will note that Cu presence has negligible effect on the appearance of other possible EPR signals, although 5 % Cu on {1 0 1} revealed some additional bands for g = 2.0061 and g = 2.0093 (Fig. S7). Therefore, we conclude that ~531.4 eV XPS signal cannot be ascribed in this case to structural defects of TiO<sub>2</sub> and its increase with Cu content results most likely from the post-deposition oxidation of Cu species, leading to different adsorbed O species and mixed oxidation state.

Ultimately, the morphology of the modified materials was investigated in detail for the 0.5 % and 5 % modified samples using TEM observations. As presented in Fig. 7b, in the case of the 5 % modified {1 0 1} particles, the formation of the few-nanometers thick coating of the Cu species is clearly observed, in contrast to the 0.5 % case (Fig. 7a). Noteworthy, no clearly-defined Cu particles were observed in case of other materials, although some surface irregularities, possibly connected with the Cu presence, can be noticed in the range of  $\leq$  1 nm. Comparing between the {1 0 1} and {0 0 1}, TEM analysis further confirmed better dispersion of the Cu species on the {0 0 1} facets, especially for the higher Cu concentration. For the lower Cu concentrations, no clear formation of any bigger Cu entities was observed, in accordance with the low XPS signals and possible formation of single-atom Cu<sup>2+</sup> species.

### 3.4. Degradation of diclofenac over modified samples and $\bullet OH$ generation

Following modification with Cu, obtained materials were tested for diclofenac degradation and compared with the previous observations for unmodified TiO<sub>2</sub>. Interestingly, as presented in Fig. 8a, the deposition of Cu had the opposite effect on the activity of both materials, clearly reducing the activity of the  $\{1 \ 0 \ 1\}$  facets and improving it for the  $\{0 \ 1\}$ 

0 1}. Furthermore, in the latter case, deposition of the Cu clearly influences the DCF adsorption, as presented in Fig. 8b. This is in accordance with the capping of the adsorption sites with Cu species, which should affect adsorption energy on the (001) plane. Moreover, as shown in Fig. 8c, modification with Cu also alternated relative concentration of the identified by-products, especially P1 and P5, which were assigned to spontaneous dehydrohalogenation (P1) and hydroxylation due to the reaction with  $TiO_2$  (P5). As observed, especially in the case of  $\{1 \ 0 \ 1\}$ samples, the relative presence of P5 is generally proportional to the degradation rate and also decreases with Cu presence. On the other hand, a slight increase of the P5 formation with Cu is observed for the {0 01}, in line with the positive deposition effect on the activity. Focusing on P1, modification with Cu increases its presence for both facets and in the case of the  $\{0 \ 0 \ 1\}$ , its formation is inversely proportional to the observed adsorption. Based on these results, it is concluded that Cu presence decreases activity of the  $\{1 \ 0 \ 1\}$ , which might include a lower ability to generate •OH since reaction with hydroxyl radicals often leads to the formation of hydroxylated products. On the other hand, photocatalytic activity of the  $\{0 \ 0 \ 1\}$  seems to be improved, however high adsorption capacity makes it a bit ambiguous to confirm. In this regard, to better decouple the effect of Cu modification on the adsorption ability and overall photocatalytic activity, additional experiments were performed using coumarin as a probe to detect generated •OH in the system via monitoring of the 7-hydroxycoumarin formation (7–OHC) [57,58]. As presented in Fig. 8d, the observed effect was analogical to the DCF degradation, however the actual maximum of the •OH generation was for the 1 % modified {0 0 1} sample, which was followed by the removal of coumarin itself (Fig. 8e). This suggests that the determined k for DCF degradation for this sample was underestimated due to the high adsorption and the optimal Cu loading is somewhere around 1 % Cu/Ti. Interestingly, for the {1 0 1}, the effect of Cu presence is again negative and starting from the concentration of 1 % Cu the formation of



**Fig. 8.** Effect of Cu loading on the kinetics of diclofenac degradation (a) and in-dark adsorption on the  $TiO_2$  surfaces (b), together with the •OH generation rate (c) determined from the formation of 7-hydroxycoumarin (7–OHC) and corresponding rate of the parent coumarin removal from the solution (d). During the •OH generation, additional {1 0 1} material modified with 0.05% Cu was included to achieve a similar  $Cu^{2+}/m^2$  ratio as for the 0.5% {0 0 1} case.

7-hydroxycoumarin was practically stopped, although some removal of the coumarin was still occurring. This results probably from the dense distribution of Cu<sub>x</sub>O species on this sample, as clearly evidenced for the 5 % Cu modified sample during TEM observations, which prevents withdrawal of h<sup>+</sup> from the TiO<sub>2</sub> structure. Moreover, since the problem of optimal surface modification connects with the large difference in the determined surface areas between both TiO<sub>2</sub> materials, we have performed additional modification of the  $\{1 \ 0 \ 1\}$  sample with 0.05 % of Cu, in order to achieve a comparable amount of  $Cu^{2+}/m^2$  between {1 0 1} and the 0.5 % modified {001}. However, as shown in Fig. 8d and 8e, the effect was still negative and overall within the trend. Finally, activity in the visible light ( $\lambda \ge 420$  nm) was also tested, focusing on the {0 0 1} with 1 % Cu as the best-performing sample. As shown in Fig. 8f, DCF degradation was initiated under such conditions, which was connected with the appearance of P5 as the only by-product observed in DAD spectra. Furthermore, control experiments confirmed that under these conditions, spontaneous photolysis of DCF is not occurring. However, the observed photocatalytic degradation rate was relatively low, which is a known problem of the TiO<sub>2</sub>/Cu<sub>x</sub>O systems [59]. Detailed results of the observed kinetics during photocatalytic reactions are presented in Figs. S8 and S9 in SI. Noteworthy, the Cu effect on TOC removal during DCF degradation followed overall trends, achieving the highest removal of 47 % for the 1 % Cu on the  $\{0 0 1\}$ , while for the 1 % Cu on the  $\{1 0 1\}$ , final TOC removal was reduced to 22 % (from 40 %).

Compared to commercial TiO<sub>2</sub>, we included analogical modification of anatase ST01 and tested it in both reactions. As presented in Fig. 8, ST01 show high ability to degrade diclofenac, probably connected with its exceptionally high surface development ( $\sim$ 300 m<sup>2</sup>•g<sup>-1</sup>). Starting from this point, the introduction of Cu decreases its activity linearly, analogically to the {1 0 1}. Coumarin-based probing of •OH further showed that its generation by ST01 is very low and shows very little change upon Cu modification. This revealed that degradation by ST01 is not dominated by •OH generation and separation of electrons between Cu and ST01 could not increase its activity.

Concerning other studies (Table 1), prepared materials allow to degrade DCF rather effectively, with >90 % of degradation achieved at least within 60 min or faster. For example, the study by Castañeda-Juárez et al. reported only 47 % of degradation over TiO<sub>2</sub>-Cu supported on clinoptilolite zeolite after 480 min of irradiation [60]. On the other

hand, high degradation rates were reported in systems with high energy radiation, including UVC, as such Hg lamps. Since DCF was found to be prone to UV photolysis, it is likely that this process is also enhanced with high energy radiation included with such light sources.

Concerning stability of the obtained photocatalysts, the bestperforming material ( $\{0 \ 0 \ 1\} + 1 \ \% \ Cu$ ) was resynthesized and tested in the 3 consecutive cycles of •OH generation. During these experiments, photocatalyst concentration was increased to  $1.5 \text{ g} \cdot \text{dm}^{-3}$  and the recovered material was washed and dried between each cycle. As presented in Fig. 9a, the material underwent visible activity drop after the first cycle and became stable afterwards. This is further connected with the visible colour change from pale-green to pale-yellow (after the first cycle), with corresponding absorbance spectra shown in Fig. 9b. The new band shows origin approximately at.  $\sim 2.15$  eV, which is a value characteristic of Cu<sub>2</sub>O. Transformation of a mixed-valence Cu species deposited on TiO<sub>2</sub> to the Cu<sub>2</sub>O after the analogical degradation process was recently reported by Wang et al., who also observed a similar activity decrease after the first reaction [59]. In this regard, it is likely that Cu<sup>2+</sup> species that are present pre-degradation became reduced to Cu<sup>+</sup> during the reaction. Noteworthy, XPS analysis revealed a slight increase of the reduced copper after the process in accordance with this explanation, as presented in Fig. 9e (please note that due to the separate preparation of  $\{0 \ 0 \ 1\} + 1 \ \%$  Cu for these experiments, detailed Cu dispersion and oxidation state might be slightly different than before, affecting XPS results). No other significant changes were noticed in the XPS spectra. XRD and SEM experiments revealed no visible changes in both materials (Fig. 9c and d).

Following observed activity trends, it is clear that the effect of Cu deposition is strictly opposite for both {0 0 1} and {1 0 1} surfaces, at least when concerning •OH generation and degradation of organic compounds. Interestingly, this was not observed before, despite a rather large interest in the TiO<sub>2</sub>/Cu systems. In this regard, a possible explanation is suggested. Recently, reported studies agree that deposition of the Cu<sub>x</sub>O species on the TiO<sub>2</sub> forms a Z-scheme junction, which should be especially beneficial for the overall activity under UV excitation [59,70]. In such heterostructure, recombination of charge carriers occurs between the TiO<sub>2</sub> conduction band (e<sup>-</sup>) and valence band of Cu<sub>x</sub>O (h<sup>+</sup>) when irradiated with the UV light. As a result, spatial separation of both carriers is achieved while preserving the high redox ability of the

Comparison of the DCF photocatalytic degradation reported in the literatu	ıre.
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Photocatalyst	Photocatalyst concentration	DCF conc.	Irradiation source and intensity	Degradation rate/removal	Ref.
TiO <sub>2</sub> -Pd 1 %	1000 mg•dm <sup>-3</sup>	50 mg•dm <sup>-3</sup>	4x Philips CLEO tubes 15 W,	$k = 0.048 \text{ min}^{-1}, \text{ pH} = 5$	[61]
TiO <sub>2</sub> -Ag 1 %	5	0	$\lambda_{\text{Max}} = 365 \text{ nm},$	$k = 0.040 \text{ min}^{-1}, \text{ pH} = 5$	
- 0			9mW		
WO <sub>3</sub>	250 mg•dm <sup>-3</sup>	$0.5 \text{ mg} \cdot \text{dm}^{-3}$	1500 W Xe lamp,	100 % after 120 min.	[62]
			550 W∙m <sup>-2</sup>		
Supported TiO <sub>2</sub> -Cu	10 000 mg•dm <sup>-3</sup>	20 mg•dm <sup>-3</sup>	Solar radiation,	47 % after 480 min.	[60]
			UV light = $383.40 \pm 0.39 \text{ Wh} \cdot \text{m}^{-2}$		
TiO <sub>2</sub>	50 mg•dm <sup>−3</sup>	10 mg•dm <sup>-3</sup>	Radium Ralutec, 9 W/78,	85 % after 240 min.	[63]
			350-400 nm		
			3.37 µeinstein∙s <sup>−1</sup>		
SrFeO <sub>3</sub>	300 mg•dm <sup>-3</sup>	50 mg∙dm <sup>-3</sup>	Hg vapor lamp 6 W	$k=0.090\ min^{-1}$	[64]
Graphite/C-doped TiO2	1666 mg•dm <sup>-3</sup>	10 mg∙dm <sup>-3</sup>	Short-arc Xe lamp	$k=0.061\ min^{-1}$	[65]
			$1.31 \bullet 10^{21} \text{ photons} \bullet \text{s}^{-1}$		
ZnO-Ce	1000 mg•dm <sup>-3</sup>	2 mg∙dm <sup>-3</sup>	Blacklight Philips TLK 05 40 W	$k=0.110\ min^{-1}$	[66]
			$\lambda_{Max} = 360 \text{ nm}$		
			$\sim 30 \text{ W} \cdot \text{m}^{-2}$		
TiO <sub>2</sub>	500 mg•dm <sup>-3</sup>	8 mg∙dm <sup>-3</sup>	2x Heraeus Hg vapor lamps (low and medium pressure)	$k = 0.903 \ min^{-1}$	[67]
			Flux at 366 nm 2.46 μeinstein•s <sup>-1</sup>		
ZnO-WO <sub>3</sub>	800 mg•dm <sup>-3</sup>	20 mg•dm <sup>-3</sup>	Metal halide lamp 400 W	$k=0.013\ min^{-1}$	[68]
Sr <sub>2.7</sub> Nd <sub>0.3</sub> Fe <sub>1.4</sub> Co <sub>0.6</sub> O <sub>7-δ</sub>	100 mg•dm <sup>-3</sup>	20 mg•dm <sup>-3</sup>	Xenon lamp 75 W	100 % after 150 min.	[69]
{1 0 1} TiO <sub>2</sub>	500 mg•dm <sup>-3</sup>	20 mg∙dm <sup>-3</sup>	Xenon lamp 300 W with IR and 310 nm filter.	$k=0.038\ min^{-1}$	This work
			UVA flux 24 mW•cm <sup>-2</sup>	90 % after 60 min	
{0 0 1} TiO <sub>2</sub> 1 % Cu	400 mg•dm <sup>-3</sup>			$k=0.024\ min^{-1}$	
				90 % after 30 min	
ST01 TiO <sub>2</sub>				$k=0.059\;min^{-1}$	
				100 % after 45 min	



**Fig. 9.** Stability tests of the  $\{0\ 0\ 1\} + 1\ \%$  Cu sample during •OH generation (a) with corresponding characteristics before and after the process: absorbance (b), XRD (c), SEM (d) and XPS (e). Please note that new sample was prepared for these experiments and some differences from previous results might be present.

Cu<sub>X</sub>O electrons and TiO<sub>2</sub> holes. Outside of the clear reduction of copper observed in the XPS spectra, we have additionally confirmed this separation with the photoluminescence spectra measurements, confirming that increased Cu loading reduces emission around ~450 nm, which should occur due to the electron-hole recombination within the TiO<sub>2</sub> structure. As shown in Fig. 10a, the effect is the same for both TiO<sub>2</sub> facets. In this regard, effective charge separation is occurring in both systems. In the case of {0 0 1} facets, this straightforwardly increases photocatalytic activity, including •OH generation, which should mostly result from the h<sup>+</sup> induced oxidation of water/hydroxyl groups present on the TiO<sub>2</sub> surface (please note that valence band potentials of Cu oxides are too low to effectively generate  $\bullet$ OH from H<sub>2</sub>O or -OH [71–77]). Noteworthy, due to the high surface energy, it is known that H<sub>2</sub>O can spontaneously dissociate on the {0 0 1} surface, forming multiple –OH, which are particularly good substrates for  $h^+$  oxidation due to the higher electron density [78]. On the contrary, the reduction ability of the {0 0 1} is generally worse than other TiO<sub>2</sub> surfaces, due to its unfavored electronic structure compared to the bulk [9,23]. In this regard, "shifting" the reduction reaction to Cu<sub>x</sub>O might be additionally beneficial for the overall activity. On the other hand, the reduction process is important for the {1 0 1} facets, which preferentially trap electrons on the surface sites [79]. Particularly, this connects with the increased hydroxylation of this surface due to the electron-induced water dissociation [80], which otherwise is partially limited. As a consequence, oxygen reduction contribute also to the •OH generation [81], and some excess electrons are important for the surface hydroxylation and h<sup>+</sup> trapping [82]. In this regard, the Z-scheme recombination of the TiO<sub>2</sub> electrons reasonably reduces the overall activity of the {1 0 1} facets, since oxygen reduction must be hindered at this surface. The scheme of these processes (charge separation and ROS generation in water/air) is presented in Fig. 10c, based on the known information about band position of TiO<sub>2</sub> and Cu<sub>2</sub>O [83-87]. Interestingly, we have also tested obtained materials in the reaction of 4-nitrophenol reduction to 4-aminophenol in methanol, to verify if the Cu effect on other reduction process would be analogical. However, as presented in Fig. 10b, the effect was negative in both cases and Cu clearly reduced amount of 4aminophenol evolved. This show that deposited Cu<sub>X</sub>O species cannot be seen as universal reduction co-catalyst and they are especially beneficial for the degradation process when combined with the  $\{0 \ 0 \ 1\}$ facets. Ultimately, these results suggest that the same Z-scheme junction might have different effect on the photocatalytic activity, depending on the electronic structure and reactions details occurring on the TiO<sub>2</sub> surface, which was clearly presented for the first time.

### 4. Conclusions

In the present work, detailed insight into the complex effects of  $Cu_XO$  loading at the {0 0 1} and {1 0 1} crystal facets of  $TiO_2$  and how it affects photodegradation of diclofenac (DCF), generation of •OH and 4-nitrophenol reduction were investigated. Overall results have presented the existence of two independent degradation pathways, either dehydrohalogenation or hydroxylation, both of them sensitive to different factors. Specifically, dehydrohalogenation occurs spontaneously in the solution under irradiation with suitable light (UVA range), leading to the



**Fig. 10.** Recorded photoluminescence spectra of the bare and Cu-modified materials after excitation with UV light (a), results of the 4-nitrphenol reduction (b) and proposed scheme of the possible reactions and charge separation during the diclofenac photooxidation in water (c). In case of the {1 0 1} facets, the scheme in (c) include dissociation of water due to the oxygen reduction on the unmodified surface, which was recently recognized as important for the {1 0 1} activity.

formation of carbazole derivatives. On the other hand, in the presence of TiO<sub>2</sub> photocatalyst, DCF hydroxylation is preferentially occurring. Selectivity of both these processes depends strongly on light energy, as well as on the DCF adsorption on TiO<sub>2</sub> surface. Noteworthy, high DCF adsorption was observed for the {0 0 1} facets, which led to almost selective formation of hydroxylated product. Following this, a very strong effect of CuxO in all systems were observed with quantitively different effects in the case of •OH generation and DCF degradation. From overall results, the synergy of  $Cu_XO$  cooperation with {001} facets in both these reactions was clearly observed, e.g. boosting their ability to generate •OH over 2-times. Combination of EPR, XRD and XPS studies revealed that depending on the Cu concentration and type of the surface, different amounts of  $Cu^{2+}/Cu^{+}/Cu^{0}$  can be observed in the system. For the most active material ({0 0 1} with 1 % Cu), copper exists mostly as the atomically-scattered Cu<sup>2+</sup> species, which are later transferred to the Cu<sub>2</sub>O during the degradation. Such atomic dispersion is a combined effect of high Cu adsorption energy and high surface development of the {0 0 1} exposing materials. Interestingly, we were not able to recreate analogical effect on the activity of the {1 0 1}, despite testing Cu concentration over 3 orders of magnitude (from 0.05 % to 5 %). Therefore, despite forming the same Z-scheme junction in both cases, the qualitative effect of Cu<sub>x</sub>O deposition was found to be different for both facets in these reactions. This is explained to result from the different roles of both  $e^{-}$  and  $h^{+}$  on the generation of reactive species on both {0 0 1} and {1 0 1}, where electrons are especially needed to stimulate the activity of the {1 0 1}, including generation of •OH. Concerning 4-nitrophenol reduction, Cu<sub>x</sub>O presence decreased its efficiency for both systems, proving that synergy between copper and {0 0 1} facets are unique for the ROS generation.

### CRediT authorship contribution statement

Szymon Dudziak: Writing - review & editing, Writing - original draft, Methodology, Investigation, Formal analysis, Conceptualization. Jakub Smoliński: Writing - original draft, Investigation, Formal analysis. Adam Ostrowski: Writing - original draft, Methodology, Investigation, Formal analysis. Aleksandra Szkudlarek: Writing - review & editing, Methodology, Investigation, Formal analysis. Mateusz Marzec: Writing - review & editing, Methodology, Investigation, Formal analysis. Krystian Sokołowski: Writing - review & editing, Methodology, Investigation, Formal analysis. Karol Sikora: Methodology, Investigation, Formal analysis. Anna Zielińska-Jurek: Supervision, Resources, Project administration, Methodology, Funding acquisition, Conceptualization.

### Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Anna Zielinska-Jurek reports financial support was provided by National Science Centre Poland. Szymon Dudziak reports financial support was provided by Foundation for Polish Science. If there are other authors, they declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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### Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.cej.2024.159146.

#### Data availability

Data will be made available on request.

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