Quasi-homogenous photocatalysis of quantum-sized Fe-doped TiO₂ in optically transparent aqueous dispersions

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In this study, the preparation of anatase TiO₂ nanocrystals via a facile non-aqueous sol–gel route and their characterization are reported. The 3–4 nm particles are readily dispersable in aqueous media and show excellent photoreactivity in terms of rhodamine B degradation. The catalytic performance can be further increased considerably by doping with iron and UV-light irradiation as a pre-treatment. The effect of surface ligands (blocked adsorption sites, surface defects etc.) on the photoreactivity was thoroughly probed using thermogravimetric analysis combined with mass spectrometry. Photoelectrochemical characterization of thin-film electrodes made from the same TiO₂ nanocrystals showed the opposite trend to the catalytic experiments, that is, a strong decrease in photocurrent and quantum efficiency upon doping due to introduction of shallow defect states.

The wide-bandgap semiconductor titanium dioxide, TiO₂, is the most widely investigated photocatalyst for the decomposition of organic contaminants and water splitting¹. At the same time, TiO₂ is one of the most promising transition-metal oxides for industrial catalytic applications due to its unique optical and electronic properties, high chemical stability, negligible toxicity and relatively low costs³. Overall, TiO₂ is still heavily researched by materials scientists across the world and is often utilized as a benchmark or reference material⁴. The photooxidation of organic species by TiO₂ would be beneficial for industrial purposes, such as the control of ground water contamination⁵ or reduction of air pollution⁶. The major advantage of using TiO₂ in water purification compared to conventional technologies, such as activated carbon adsorption, is the complete mineralization to CO₂ and H₂O combined with low processing costs⁶,⁷. However, the degradation rate of many organic pollutants (e.g., dioxins) when using common photocatalysts is too low for practical applications⁸. For increasing the activity of catalysts in heterogeneous photocatalysis, a high dispersability in the reaction medium is desirable to avoid light scattering processes at the surface by suspended particles⁹–¹¹. Note that the activation of bulk particles requires an isotropic medium that provides a sufficient light penetration depth¹²,¹³.

The relatively large bandgap of anatase TiO₂ (E_g ≈ 3.2 eV), corresponding to wavelengths shorter than ~390 nm, allows < 4% of solar energy to be converted into electrical energy. The most effective way of achieving an absorption red shift in the photocatalyst, thereby enhancing the generation of photoexcited charge carriers, has been found in the doping of TiO₂ with various cations¹⁶. Metal-ion dopants substitute the Ti⁴⁺ site in the crystal lattice, creating discrete energy levels within the electronic band structure¹⁷. Iron has been demonstrated to be one of the most efficient dopants for increasing the photoreactivity¹⁸. Iron-doping induces energy levels between 0.2 and 0.4 eV above the valence-band edge, as shown for quantum-sized TiO₂ particles¹⁹. Furthermore, large improvements in photoreactivity have been observed through nanostructuring²⁰–²² and substitutional/interstitial anion doping with nitrogen²³ and sulfur²⁴,²⁵, shifting the optical absorption spectrum into the visible range²⁶.

Since Niederberger et al. applied the general concept of non-aqueous sol–gel chemistry to the preparation of TiO₂ nanoparticles in 2002²⁷, several authors reported about modified low-temperature synthesis routes of anatase TiO₂ for photocatalytic applications²⁸–³⁰. For example, Zhang et al. reported about improved photo-reactivity of 15 nm TiO₂ particles and proved their long-term stability as a dispersion in water³¹. However, to
the best of our knowledge, there are no studies available on the preparation and photocatalytic performance of (non-aqueous) sol–gel derived Fe-doped TiO₂ nanocrystals having a size below 5 nm. It is worth mentioning that apart from photocatalysis, the surface composition of such particles can be tailored toward the fabrication of well-defined mesoporous or macroporous thin-film electrodes for electrochemical applications.

In this study, the photocatalytic activity of undoped and Fe³⁺-doped TiO₂ nanocrystals synthesized via a facile benzyl alcohol-based sol–gel route was examined regarding the bleaching of rhodamine B (RhB). The 3–4 nm particles exhibited a high dispersability in aqueous media (up to 10 wt-%), forming colloidal dispersions that were stable over days. In addition to nanostructuring and doping, emphasis is placed on the correlation between ultraviolet (UV)-light irradiation as a kind of pre-activation and photoreactivity. It is demonstrated that the TiO₂ nanocrystals show significantly improved photodegradation kinetics after irradiation due to the removal/stripping of organic surface ligands.

Results

Considering the parameters discussed in the Introduction section, photocatalyst materials need to be tailored in terms of particle size, bulk and surface composition, degree of crystallinity and dispersability. The photocatalysts employed in this work were prepared by a non-hydrolytic benzyl alcohol-based synthesis route, allowing the formation of both undoped and doped TiO₂ nanocrystals, the latter by incorporation of Fe³⁺ ions during the condensation reactions. In recent years, Fe(NO₃)₃·9H₂O for cation doping has been shown to be a suitable precursor due to the relatively weak adsorption of NO₃⁻ on the particle surface. To study the effect that the iron doping level has on the photoreactivity, TiO₂ nanocrystals of different composition were prepared and tested. The amount of incorporated dopants was determined by energy-dispersive X-ray spectroscopy (EDS) and found to be ~ 1.0, 1.5 or 4.3 mol.% (5% error). Overall, EDS indicated the presence of Ti, O, Cl and Fe (see Supplementary Figure S1). In this context, we also note that the actual Fe³⁺ doping level deviated strongly from the amount of precursor used in the synthesis (e.g., 1.5 versus 7.5 mol.%).

X-ray diffraction (XRD) patterns of the undoped and Fe³⁺-doped TiO₂ nanocrystals (see Fig. 1 and Supplementary Figure S2) were found to match the anatase phase of TiO₂ (JCPDS card no. 21-1272). The pronounced reflections confirm the crystallinity of the TiO₂ samples. This result is noteworthy, as sol–gel derived materials often require some kind of post-treatment for achieving a reasonable degree of crystallinity. Furthermore, no α-Fe₂O₃ (note that hematite is the thermodynamically most stable iron-oxide phase) reflections were observed, supporting the assumption that Fe³⁺ (0.64 Å, octahedral coordination, high-spin state) substitutes the Ti⁴⁺ (0.605 Å) site in the lattice. The average crystallite size was determined by applying the Scherrer equation to the full width at half-maximum (FWHM) intensity of the 101 reflection at 2θ = 25.2°. It was found to be 3.5 nm for both the undoped and Fe³⁺-doped TiO₂ materials (i.e., the doping level has no notable effect on the crystallite size). For such small-size particles, high photocatalytic activities have been observed, as the electrons and holes after optical excitation can readily react with the adsorbate on the surface of the adsorbent if the dopant acts as a mediator of interfacial charge transfer. Taken together, XRD provides clear evidence of the crystalline nature of the TiO₂ nanoparticles, which is required for proper charge-carrier migration to the surface (interface). Amorphous photocatalysts usually show lower photocatalytic activity due to the presence of abundant defect sites, acting as recombination centers.

To probe the dispersability of the TiO₂ nanocrystals in water, dynamic light scattering (DLS) measurements were performed (see Supplementary Figure S3). DLS indicates a narrow particle size distribution with an average hydrodynamic diameter of (12.5 ± 2.6) nm. This result suggests that the nanoparticles tend to form aggregates in polar solvents, which is reasonable from a surface-energy perspective and is in agreement with electron microscopy imaging data (see Supplementary Figure S4). In addition, selected-area electron diffraction (SEAD) confirmed the phase purity of the anatase TiO₂ nanoparticles.

Because the specific surface area plays an important role in catalytic applications, N₂-physisorption measurements were conducted on the TiO₂ nanocrystals. The adsorption/desorption isotherms (see Supplementary Figure S5) confirm the high surface area and porosity of the nanocrystals. The specific surface area calculated using the BET method is 150 m²/g, indicating a good accessibility of the active sites. The pore size distribution shows a bimodal pore structure with pores in the mesopore range (2-50 nm) and a small contribution from macropores (50-200 nm), which is beneficial for the fast diffusion of reactants and products.
Figure S5) show a combination of H2- and H4-type hysteresis. In general, H2 hysteresis is attributed to condensation of adsorbate in porous materials that show pore blocking/percolation effects. In contrast, H4 hysteresis can be primarily assigned to cavitation-induced evaporation (in mesoporous and microporous materials). Both can be somewhat expected, as the aggregation of TiO2 nanocrystals inevitably leads to interparticle void formation. Brunauer–Emmett–Teller (BET) analysis provided a specific surface area of $A_{\text{BET}} \approx 260 \text{ m}^2/\text{g}$ for both the undoped and Fe3+-doped TiO2 nanocrystals. This result agrees well with theoretical calculations assuming a cubic close-packed arrangement of 3.5 nm spherical particles.

Figure 2 shows optical absorption spectra for the undoped and Fe3+-doped TiO2 nanocrystals. The data indicate that the doped material absorbs light much stronger at wavelengths shorter than 400 nm than the undoped counterpart does. The 1.5 mol.% Fe3+-doped TiO2 nanocrystals exhibited the largest optical absorption properties (see Supplementary Figure S6). The intrinsic band-edge absorption due to interband transition from the O-2p to Ti-3d orbitals of undoped titania can be found in the range between 375 and 340 nm. The stronger absorption of the doped material (red shift) between 450 and 355 nm results from the excitation of electrons from the Fe-3d states to the TiO2 conduction band. For TiO2 as an indirect bandgap semiconductor, the square root of the absorption coefficient is expected to vary linearly with photon energy near the bandgap energy. The optical bandgaps were determined from the respective Tauc plots (see Supplementary Figure S6) to be about 3.4 eV (365 nm) and 3.3 eV (376 nm) for the undoped and Fe3+-doped TiO2, respectively. The fact that $E_{\text{gap}}$ is larger by 0.1–0.2 eV compared to that of bulk anatase TiO2 can be ascribed to nanoconfinement effects, in agreement with the bandgap shift of 0.16 eV for 3.8 nm TiO2 particles observed by Anpo et al. The theoretical calculation model for bandgap shifts of quantum-sized particles was first described by Brus et al. in 1984. The nanoconfinement effect, in general, can be explained by Heisenberg’s uncertainty principle: As the particle size decreases, the ground state energy of confined electrons must increase to satisfy the uncertainty principle ($\Delta x \cdot \Delta p \leq \hbar$).

The photocatalytic activity of the undoped, (1.0, 1.5 and 4.3 mol.%) Fe3+-doped and anatase/rutile P-25 (Degussa) TiO2 was studied by monitoring RhB [as a representative environmental (organic) pollutant] degradation under UV-light irradiation [< 254 nm, see UV–visible (UV–vis) spectra in Fig. 3A,B]. The 1.5 mol.% Fe3+-doped TiO2 was found to be the most active photocatalyst among the materials tested in this work (see Supplementary Figure S7) and therefore chosen for further structural/optical investigations together with the undoped TiO2 sample. The results in Fig. 3C show that the photobleaching in water follows pseudo-first-order reaction kinetics. The degradation rate, $k$, for the 1.5 mol.% Fe3+-doped TiO2 was about two times larger ($k = 9.0 \cdot 10^{-2} \text{s}^{-1}$) compared to that of undoped TiO2 ($k = 4.3 \cdot 10^{-2} \text{s}^{-1}$) and the P-25 reference material ($k = 4.6 \cdot 10^{-2} \text{s}^{-1}$). This result demonstrates that the photocatalytic activity of TiO2 nanocrystals can be substantially improved via doping with iron. Interestingly, the experimental data for the as-prepared TiO2 nanocrystals did not show the same linear (kinetics) trend observed for P-25. This finding can probably be explained by the presence of organic surface ligands from the synthesis. The ligands may act as trap states for charge carriers at the particle surface and can sterically hinder the adsorption of RhB. The first data point ($t = 0$ min, black lines in Fig. 3A,B) was measured in the absence of photocatalyst material. For the P-25/RhB suspension (Fig. 3A), a sudden increase in absorption at $\lambda \leq 500$ nm after $t = 0$ min was observed, which is due to scattering of incident light, especially in the visible range, by the TiO2 particles.

The photographs in Fig. 4 clearly show that the P-25 particles cannot be dispersed well in water, forming a suspension and thereby scattering the incident light to some degree. In contrast, the sol–gel derived TiO2
nanocrystals produced an optically transparent (colloidal) solution under the very same conditions. Therefore, it was not necessary to centrifuge the particles for photocatalytic measurements.

The degradation rate has been reported to be proportional to the incident light intensity below 20 mW/cm² and exhibit square root dependence above ~ 25 mW/cm²[55,56]. This suggests that the photoreactivity is strongly affected by the penetration depth of UV light. The latter was examined for the TiO₂ nanocrystals dispersed in 80 mL aqueous RhB solution (25 μmol/L, see Supplementary Figure S8). The absorbance of incident photons by the RhB/TiO₂ dispersion revealed an exponential decay from 300 to 370 nm, meaning that the absorption of high-energy photons is significantly increased. On the other hand, the physical penetration depth decreased from 24 to 2.5 cm between 370 and 300 nm. Hence, to make use of high-energy photons (in an effective manner) for charge-carrier generation in the TiO₂ photocatalyst, the penetration depth of light and the reactor...
mirradiation removed remaining chlorine ligands (Supplementary Figure S10) further supports the presence of organic surface contaminants. In addition, the UV-light bulk anatase TiO₂, electrons may be trapped in the respective states energetically located within the electronic conduction-band structure and holes in the Fe³⁺ states (located within the TiO₂ bandgap) after photoexcitation. However, for quantum-sized particles, they are at least several tenths of meV below the conduction band due to nanoconfinement-induced bandgap widening. Therefore, recombination of electron–hole pairs is somewhat inhibited, resulting in extended diffusion lengths. The trap-mediated charge transfer is well known to occur through detrapping mechanisms and tunneling processes. In a 4 nm TiO₂ particle (~ 1000 atoms, ~ 2 nm exciton diameter), the majority of dopant atoms are located close to or at the surface. Consequently, trapped electrons and holes may lead immediately to the formation of highly reactive oxygen species, O₂•−, HOO• and/or 'OH. Such mediated charge-transfer processes (besides the superior light absorption in the visible range due to introduction of Fe-3d states, see Fig. 2) help to explain the substantially increased photoreactivity observed for the 1.5 mol.% Fe²⁺-doped TiO₂ as compared to the undoped nanocrystals. The highest photodegradation rate was demonstrated for a doping level of 1.5 mol.% (see Supplementary Figure S7), in good agreement with the study reported by Ranjit et al. ¹⁻°³××°¹⁻°⁻°×°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°⁻°°°resco
The radicals formed through redox reactions with O₂ and H₂O initiate the photobleaching of RhB by attacking the aromatic chromophore ring structure and inducing de-ethylation and oxidative degradation.

Apart from photocatalysis, TiO₂ nanomaterials have been widely investigated in photoelectrochemical applications, e.g., in dye-sensitized solar cells (DSSC) or for solar hydrogen generation. The latter describes the process of splitting water into hydrogen and oxygen in a simple electrochemical experiment (using a semiconductor electrode and the energy of light irradiation) and was first reported by Fujishima and Honda in 1972.

Figure 7A shows the I-U characteristics of nanocrystal-based TiO₂ thin-film photoelectrodes on FTO-coated glass substrates. The measurements were carried out at a sweep rate of 2 mV/s. The undoped and 1.5 mol.% Fe³⁺-doped TiO₂ nanocrystals showed an open cell voltage, $U_{oc}$, of about −0.95 and −0.8 V (no steady-state value) and a steep increase in photocurrent, $j_{ph}$, with increasing applied voltage, $U_{appl}$. $j_{ph}$ increased only slightly between −0.5 and +0.8 V (0.54 and 0.02 mA/cm² short-circuit current densities for the undoped and Fe³⁺-doped TiO₂, respectively). For $U_{appl} > 0.8$ V, $j_{ph}$ strongly increased again. However, this result has no photoelectrochemical origin, as the same behavior was observed without illumination. The maximum conversion efficiency was determined to be 0.18% for the undoped and 0.007% for the Fe³⁺-doped TiO₂ (at $U_{appl} = 0$ V).

The quantum efficiency or incident photon-to-current efficiency (IPCE) is also commonly used to evaluate the performance of photoelectrodes. Assuming ideal monochromatic light and complete Faradaic conversion, the IPCE can be calculated from the photocurrent response (Fig. 7B,C). As is evident, both samples showed notable IPCEs for wavelengths shorter than 380 nm, with maximum efficiencies of about 23 and 1.8% (within the uncertainty indicated by the error bars) for the undoped and Fe³⁺-doped TiO₂, respectively. Measurements at lower wavelengths were not feasible for experimental reasons. The large error for small wavelengths is due to the UV fraction in the light-source spectrum (relatively large uncertainty of the intensity measurement).

Although the 1.5 mol.% Fe³⁺-doped TiO₂ showed superior performance in the photobleaching of RhB, in this kind of photoelectrochemical experiments the doping reduced the efficiency by more than an order of magnitude. In contrast to photocatalysis with nanoparticles (3–4 nm) dispersed in solution, efficient electron transport over extended distances (several hundred nanometres) is required in a photoelectrode. As known from literature and as described above, Fe dopants introduce shallow trapping sites, thereby lowering the recombination rates of photoexcited charge carriers. This in turn leads to improved photocatalytic activity as long as the carriers are trapped near the particle surface or diffusion pathways are short. Zhang et al. showed that the optimal iron doping level decreases (from 0.2 to 0.05 at%) with increasing particles size. In a photoelectrochemical experiment, only efficient “long-distance” charge transport, from the position of the excited site to the electrode back
contact, results in high photocurrents and therefore high performance. Egerton et al. studied sol–gel derived TiO₂ photoelectrodes with varying iron levels and observed a characteristic decrease in photocurrent with increasing content from 0 to 2.2%68. This is in agreement with the findings of this work.

Discussion
In summary, we have applied a non-hydrolytic sol–gel route for the preparation of anatase TiO₂ nanoparticles that can be readily re-dispersed in aqueous media. The particles had a size of 3–4 nm in diameter, leading to a high specific surface area of ~ 260 m²/g. Because of their excellent dispersability, the TiO₂ nanoparticles were investigated in the photocatalytic degradation of RhB in water. In general, the dispersability provided an ideal experimental setup, with even deeply dispersed particles in the photoreactor being addressable for high overall reactivity. The photocatalytic performance was further improved by a factor of two through appropriate iron doping and pre-irradiation using UV light. TGA-MS measurements showed that the latter leads to surface ligand removal/stripping, increasing the number of active adsorption sites, among others, and therefore the rate of mineralization. However, photoelectrochemical studies revealed that iron doping adversely affects the photoresponse of TiO₂ thin-film electrodes because of differences in the migration distances of charge carriers (bulk recombination determines the photocurrents of TiO₂ photoelectrodes).

Materials/methods
Materials. 
Titanium tetrachloride (99.9%), anhydrous benzyl alcohol (99.8%), iron nitrate (FeNO₃·9H₂O) and rhodamine B (RhB, dye > 90%) were purchased from Sigma-Aldrich. Ethanol (absolute grade) and methanol were purchased from VWR and propanediol (99%) from Acros Organics.

Synthesis of TiO₂ nanocrystals. 
The preparation of anatase TiO₂ nanoparticles was adopted from Niederberger et al.27 and Brezesinski et al.34 1 mL (9.12 mmol) TiCl₄ was added dropwise to 5 mL absolute ethanol in a water-free and loosely sealed glass vial. The resulting transparent (pale yellow) solution was combined with 0.25 mL propanediol and mixed with 20 mL benzyl alcohol under vigorous stirring. Finally, the required amount of FeNO₃·9H₂O was dissolved into the solution, followed by treatment in an ultrasonication bath to achieve a homogenous and transparent sol. The sol was filtered using a millipore filter (0.2 μm) and then heated at 110 °C for 3 h under constant stirring. To isolate the nanoparticles, the opaque suspension was precipitated in 250 mL diethyl ether, centrifuged at 6000 rpm for 10 min and subsequently dried at ambient conditions. The undoped and Fe-doped TiO₂ appeared as white and yellow powder, respectively. Transparent aqueous dispersions were obtained by re-dispersion of nanoparticles in deionized water.

Figure 7. (A) I–U characteristics of the undoped and 1.5 mol.% Fe³⁺-doped TiO₂ nanocrystals deposited onto FTO-coated glass substrates under irradiation (~ 360 mW/cm²). Quantum efficiency of undoped (B) and doped (C) TiO₂ at Uₐₕₗ = 0 V. The connecting lines are for eye guidance.
Synthesis of TiO₂ thin films. Thin films were prepared by the dip-coating method on quartz glass (Lithosil®, Schott) or FTO-coated glass substrates. For the dip-coating solution, 500 mg (as-prepared) nanoparticles were dispersed in 4 g methanol and 0.3 g double distilled water. Optimal coating conditions included a relative humidity of 70% and constant withdrawal rates from 3 to 13 mm/s. The as-made films were dried in air at 100 °C for 1 h and then at 300 °C for 12 h. To remove the organic constituents, the samples were heated to 550 °C within 50 min and kept at this temperature for 5 min.

Characterization. The dopant concentration was probed via energy-dispersive X-ray spectroscopy (EDS, Link Pentafet, 7426, Oxford Instruments) by irradiating a (10×10) μm² sample area at 10 kV. X-ray diffraction (XRD) measurements were performed on an XPert Pro diffractometer from Panalytical Instruments (Cu Kα radiation) at an acceleration voltage of 40 kV and emission current of 30 mA. XRD data were collected in θ–2θ geometry in the range of 20°–80° in step scan mode (0.008° step size). N₂-physisorption experiments were carried out at 77 K using a Quantachrome Autosorb instrument. Prior to the measurements, the samples were degassed in a vacuum at 120 °C. For thermogravimetric analysis (TGA), a QMG421 mass spectrometer system (Balzers) combined with a Netzsch STA409PC thermal analyzer were used. Specifically, 10 mg vacuum-dried (24 h) TiO₂ powder was heated to 800 °C at a rate of 5 °C/min in an oxygen/argon (20:80) atmosphere. The optical properties of nanoparticles were determined by measuring the light absorption spectra of diluted dip-coating solutions (~6 mg/mL) using a UVIKON XS spectrophotometer equipped with 3 mL cuvettes.

Photocatalytic experiments. The photocatalytic activity was measured by monitoring the photobleaching of RhB. In a quartz reactor, 5 mg TiO₂ nanoparticles were dispersed into 35 or 70 mL deionized water under moderate stirring. Subsequently, 5 or 10 mL aqueous solution of 200 μmol/L RhB was added to the dispersion, yielding a 25 μmol/L RhB solution. The transparent dispersion was aged for 10 min to attain adsorption/desorption equilibrium of dye on the photocatalyst surface, as reported in detail elsewhere²⁹. In case of the pre-irradiation treatment, TiO₂ nanoparticle dispersions were illuminated with a UV lamp (8 W, λ < 254 nm, Benda NU-8 KL) for 120 min to decompose the organic (and inorganic) surface ligands. The distance between the light source and the sample was about 6 cm. The quantitative change in absorbance at 554 nm was measured using an UVIKON XS spectrophotometer.

Photoelectrochemical experiments. Photoelectrochemical measurements were carried out in a rectangular PMMA reactor [(30×50×100) mm³] equipped with a quartz window. A two-electrode system was used with a Pt foil counter-electrode (300 mm²) and the TiO₂ electrode on FTO-coated glass substrate (19.6 mm²) in an aqueous 0.1 M NaOH electrolyte. For bias supply and current measurements, a Zahner IM-6 electrochemical workstation was used. A 150 W Xe Arc lamp (LOT-Oriel) served as the light source. A thermopile and powermeter were used for light-intensity measurements. The incident photon-to-charge carrier conversion efficiency was examined by irradiating the sample with monochromatic light (LOT-Oriel monochromator MSH101, 2.5 mm width of entrance and exit slit). The photocurrent at each wavelength was recorded until steady-state condition was reached (at constant electrode potential of 0 V).

Data availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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