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Single atom Cu(I) promoted mesoporous titanias for photocatalytic Methyl Orange

depollution and H₂ production

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Abstract

Tailoring the physicochemical properties and hence reactivity of semiconductor photocatalysts in a predictable fashion,

remains a challenge to their industrial application. Here we demonstrate the striking promotional effect of incorporating

single Cu(I) atoms, on aqueous phase photocatalytic dye degradation and H₂ production over surfactant-templated

mesoporous TiO₂. X-ray absorption spectroscopy reveals that ultra-low concentrations of copper (0.02-0.1 wt%)

introduced into the mesoporous TiO₂ surface create isolated Cu (I) species which suppress charge recombination, and

confer a six-fold photocatalytic promotion of Methyl Orange degradation and four-fold enhancement of H₂ evolution. The

impact of mesopore structure and photophysical properties on photocatalytic activity is also quantified for the first time:

calcination increases mesopore size and nanocrystalline order, and induces an anatase-rutile phase transition that is

accompanied by a decrease in the optical band gap, increased charge carrier lifetime, and a concomitant significant

activity enhancement.

Keywords: photocatalysis; titania; mesoporous; copper; XAS

Introduction

Heterogeneous, semiconductor photocatalysts have received significant attention due to their potential application in

diverse photoinduced reactions, notably solar fuels[1, 2] and hydrogen generation,[3] and wastewater and air

depollution.[4] Titania has been widely used for water depollution[5] due to its abundance, low cost, thermochemical

stability, low toxicity and suitable optical properties.[6] In this context, commercial P25 titania, comprising anatase and

rutile crystallites (which appear to exhibit synergistic photocatalysis due to heterojunction formation) is typically the first

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port of call for academic and industrial applications.[7, 8] However, P25 suffers from a low surface area (50 m².g⁻¹), wide band gap (restricting it to UV light absorption) and rapid recombination of photoexcited charge carriers.[9, 10] Consequently significant research efforts have focused on developing high area, thermally stable titanias with enhanced photophysical properties.

Antonelli et al[11] synthesised the first high internal surface area (200 m².g¹), mesoporous TiO₂ with uniform (3.2 nm) pores via a modified sol-gel synthesis using titanium alkoxides and phosphate surfactants. High area (150-240 m².g¹) and highly crystalline mesoporous titania microspheres with average pore diameters ~4.5 nm were more recently synthesised via evaporation induced self-assembly and acetic acid mediated sol-gel chemistries.[12] However, mesoporous TiO₂ catalysts with such mesopores are only amenable to substrates with small hydrodynamic diameters, and may suffer mass transport limitations for bulky molecules (such as azo dyes). Zhou et al[13] developed a lengthy, multistep route to achieve larger (10 nm) well-ordered mesopores in anatase TiO₂, and investigated the thermal stability and photoactivity of this material for 2,4-dichlorophenol degradation. The highly crystalline anatase mesoporous TiO₂ exhibited better photocatalytic activity than P25, nonetheless, charge carrier separation was limited in this single phase material. In this latter example, it is surprising to note that calcination of disordered mesoporous titanias failed to induce any systematic variation in the anatase:rutile phase composition, or photoactivity for imazapyr or phenol decomposition.[14] Despite efforts to optimise the synthesis of mesoporous TiO₂ photocatalysts,[11-17] there remains no general route to systematically tune their phase or mesostructure.

Titania photoactivity is predominantly influenced by structural and electronic properties, [18, 19] which may be tuned via crystal structure engineering, [20] The commonly studied metastable anatase and stable rutile phases both exhibit wide band gaps (3.2 eV and 3.0 eV respectively)[21], with anatase-rich mixtures with rutile proposed optimal for MO degradation due to charge separation across their interface. [22, 23] Density functional calculations have also highlighted the importance of crystal facets for anatase, with surface energies and photoactivity of decreasing from (001) 0.90 J m⁻² > (100) 0.53 J m⁻² > (101) 0.44 J m⁻².[24, 25] Surface defects such as oxygen vacancies (V_0) and Ti³⁺ sites also enhance TiO₂ photoactivity, [26] alongside particle size effects which dictate an optimum for 25-40 nm particles due to competition between light harvesting, scattering, bulk / surface recombination and geometric surface area.[27] Heterojunction formation[28] and doping[29] are promising approaches to overcome the limitations of pure titanias through modifying the quantum efficiency, band gap/energies, charge carrier mobility, surface structure, and adsorption properties.[23, 28-33] Functionalising by earth abundant metals such as copper is especially appealing with a wide range of promoter formats employed, including Cu metal, [34] Cu_xO clusters, [35] nano-Cu₂O, [28] Cu doped sol-gels, [36, 37] Cu(OH)₂ clusters, [38] and Cu₂O shells [39] to extend light absorption and improve charge carrier mobility, and in turn quantum efficiencies for photocatalytic pollutant degradation [28, 34-36] and solar fuels production. [38, 39] Cu-doped titania has also been exploited for thermo- and photocatalytic applications, including butene hydrogenation, [40] CO₂ reduction, [37] H₂ production,[41] and dye degradation.[42] In most cases, copper is introduced at moderate→high loadings spanning 0.5-5 wt% Cu, resulting in the formation of Cu₂O₂(37, 42) CuO₂(41) and/or Cu metal nanoparticles of 1-10 nm.[40]

However, it has proven difficult to unequivocally identify the copper oxidation state (due to XPS sensitivity and energy resolution limitations) or location of copper species (discrete particles or within the titania framework) for low loadings and highly dispersed particles.

Considering the former application, water pollution and its associated impact on human health accounts for >840,000 fatalities annually worldwide.[43] Around 80 % of contaminants in wastewater arise from the unregulated discharge of toxic, recalcitrant organic compounds by the textile, paper, petrochemical, food, energy and mineral processing sectors into aquatic ecosystems, and their depollution represents a significant challenge particularly for emerging countries.[44] Recalcitrant organic compounds, which include organic Azo dyes such as Methyl Orange, cannot be treated by conventional biological and/or physicochemical processing (e.g. microorganisms, flocculation or chlorination);[45, 46] the concentration of such Azo-dyes can reach 500 ppm in in textile effluent.[47] Advanced oxidation processes, including Fenton oxidation, [48] photocatalytic oxidation, [49, 50] photo-Fenton oxidation, [51] catalytic wet air oxidation, [52] electrocatalytic oxidation, [53] electro-Fenton oxidation[54] are promising solutions for the oxidative removal of persistent organic compounds from wastewater. However, while Fenton-type AOPs exhibit high removal efficiencies (100 mg/L) for diverse ROCs including cyanotoxins, they share a common high demand for H₂O₂ (~2.1 kg per kg of biological/chemical oxygen demand).[55] Furthermore, heterogeneous Fenton analogues are susceptible to transition metal leaching, [56] Hence, photocatalytic solutions to wastewater depollution are highly sought after.[4] In regard of the second application above, energy security and climate change represent key global challenges arising from historic reliance on fossil fuels. [57] Artificial photosynthesis offers the possibility of clean energy through water photolysis and renewable chemicals through CO₂ utilisation as a sustainable feedstock, commonly termed solar fuels and chemicals.[58] Hydrogen production through photocatalytic water splitting over semiconductor nanomaterials represents one of the most promising routes for the conversion and storage of solar energy in a form amenable for transportation.[59]

Here, we demonstrate the first synthetic route to high area, mesoporous TiO₂ with systematically tunable textural and photophysical properties, enabling facile control over their photocatalytic activity for both aqueous phase Methyl Orange degradation (in the absence of dye sensitisation) and hydrogen production. Post-functionalisation of mesoporous titania surfaces by ultra-low (0.1 wt%) concentrations of copper confers a six-fold enhancement in the rate of Methyl Orange photodegradation, and a four-fold increase in H₂ evolution, which Cu K-edge XAS reveal are associated with the genesis of atomically-dispersed Cu(I) species, and concomitant improvements in charge carrier lifetime. This work represents the first report that single Cu(I) atoms can promote titania photocatalysis, and quantitative correlation of the effect of titania mesoporosity on photocatalytic activity. The unexpected synergy between isolated Cu(I) atoms and mesoporous titania offers a new approach to promoted photocatalysts utilising earth abundant elements.

Experimental

Materials

Titanium (IV) n-butoxide (ACROS Organics, 99.0 %), Pluronic F-127 (Sigma-Aldrich), nitric acid (Fisher, 70 %), ethanol (Fisher, analytical reagent grade), copper (II) nitrate trihydrate (ACROS Organics, 99 %), Methyl Orange (Sigma-Aldrich, Reag. Ph. Eur.), anatase (ACROS Organics, 98.0 %), rutile (Sigma-Aldrich, 99.99 %) and P25 (Sigma-Aldrich, 99.5 %) were used as received.

Catalyst preparation

Mesoporous titanias (meso-TiO₂-XC, X=calcination temperature) were synthesised through a surfactant templating route employing the non–ionic triblock copolymer Pluronic F-127 (12,600 g mol⁻¹, 7.7 g) dissolved in ethanol (120 ml), followed by HNO₃ (6.3 ml) and titanium (IV) n-butoxide (16 ml) addition under rapid stirring. The resulting solution was stirred at room temperature for 6 h, and then transferred to an oven for solvent evaporation at 50 °C for 18 h. The gel thus obtained was calcined in a muffle furnace at a range of temperatures between 300-700 °C (ramp rate of 1 °C.min⁻¹ in each case) for 4 h, and finally ground to a fine powder. Copper doped materials were prepared by subsequent wet impregnation of a Cu(II) nitrate trihydrate solution of variable concentration on the 500 °C calcined mesoporous TiO₂ (meso-TiO₂-500C) to achieve Cu loadings spanning 0.02 to 12.33 wt%. The aqueous slurry was stirred in the dark at 50 °C for 17 h until a dry powder was obtained, and the resulting solid subsequently re-calcined at 400 °C (ramp rate of 10 °C.min⁻¹) under flowing oxygen (10 ml.min⁻¹) for 2 h, and finally ground to a fine powder.

Characterisation

Nitrogen porosimetry was performed on Quantachrome NOVA 4000e and 4200e porosimeters with analysis by NovaWin v11.03 software. Samples were degassed at 120 °C in vacuo for 16 h before measurement. BET (Brunauer-Emmett-Teller) surface areas were calculated over the relative pressure range 0.05-0.35. Mesopore size distributions were estimated applying the BJH (Barrett-Joyner-Halenda) method to the desorption isotherm for relative pressures >0.35. Powder X-ray diffraction (XRD) patterns were recorded on a Bruker D8 ADVANCE diffractometer (40 kV, 40 mA) employing a Cu K_{α} (1.54 Å) source fitted with a LynxEYE high speed strip detector and calibrated against a corundum standard. Wide-angle patterns were recorded for 2θ =10-80° with 0.02° steps and low-angle patterns recorded for 2θ =0.3-8° in 0.01° steps. The proportion of crystalline anatase and rutile was estimated from the Spurr–Myers equation:

% $Anatase = \frac{1}{1+1.265\frac{I_R}{I_A}}x100$ %; % $Rutile = \frac{1}{1+0.8\frac{I_A}{I_R}}x100$ % where I_R and I_A are the respective peak height of the rutile

(110) and anatase (101) reflections.[60, 61] Temperature-programmed oxidation (TPD) was conducted using a Mettler Toledo TGA-DSC 2 thermal analyser from 45-800 °C (ramp rate of 10 °C.min⁻¹) under flowing N_2/O_2 (40:20 v/v, total flow of 60 ml.min⁻¹). Diffuse reflectance UV-vis spectroscopy (DRUVS) was performed on a Thermo Scientific

Evolution 220 spectrophotometer equipped with an ISA-220 integrating sphere. Optical band gaps were determined from Tauc plots for an indirect band gap material, $F(R) = (\frac{(1-R)^2}{2R} \times hv)^{1/2}$ using the Kubelka-Munk formalism, where R corresponds to reflectance (%R/100), with the band gap obtained from the intersection of a tangent to the function with the energy axis. High-resolution scanning transmission electron microscopy (HR-STEM) images were recorded on a JEOL 2100F FEG microscope operating at 200 keV and equipped with a spherical aberration probe corrector (CEOS GmbH). Samples were prepared by dispersion in methanol and drop-casting onto a copper grid coated with a holey carbon support film (Agar Scientific). Images were analysed using ImageJ v1.46r software. X-ray absorption spectroscopy (XAS) measurements were undertaken at the XAFS beamline of the Elettra synchrotron. Ex-situ fluorescence and transmission Cu K-edge (8987.96 eV) spectra were acquired on powder samples diluted with BN and packed in Teflon washers. Spectra were normalised, background subtracted, and fitted to Cu₂O, CuO, Cu(NO₃) and Cu foil standards using the Athena and Artemis components of the IFEFFIT software suite version 1.2.11d respectively. Time-resolved photoluminescence spectra of aqueous suspensions (1 mg.mL⁻¹) of samples were recorded on a F-4500FL spectrometer at an excitation wavelength of 380 nm.

Photocatalytic activity

Photocatalytic Methyl Orange (MO) degradation tests of meso-TiO2 and Cu/meso-TiO2 materials were performed in a sealed 150 ml quartz photoreactor (27 mm i.d.) maintained at an ambient of temperature 20°C through a recirculating cooling-jacket connected to a Huber Minichiller. 50 mg of catalyst was added to 50 ml of a 20 ppm MO aqueous solution stirred at 625 rpm in the dark for 1 h to establish an adsorption-desorption equilibrium prior to illumination. The reactor was subsequently irradiated (26 mW.cm⁻² reactor internal flux) under stirring by a 200 W Newport Oriel Instruments 66002 Hg Xe arc lamp employing a water filter to remove infrared wavelengths. Reaction aliquots were periodically withdrawn, centrifuged to remove catalyst, and analysed in cuvettes by UV-Vis absorption spectroscopy employing a Thermo Scientific Evolution 220 spectrophotometer. Initial degradation rates were calculated spectrophotometrically from the 464 nm MO absorption band, employing multi-point calibration curves and corrected for background rates of dye photolysis in the absence of catalyst. Degradation products were determined on periodically withdrawn and filtered aliquots using an Agilent 1200 series HPLC equipped with refractive index and UV diode array detectors, and a Zorbox Eclipse column and 10 % acetonitrile/90 % water mobile phase. For hydrogen production via water splitting, 50 mg of sample was dispersed in 50 ml of 0.5 M Na₂SO₃ aqueous solution (sulphite as a hole scavenger), and ultrasonicated for 10 min to obtain a uniform dispersion. Subsequently, the reaction mixture was purged with He (15 ml/min) for 1 h to remove air from the system. Aliquots of the reaction gas mixture was periodically withdrawn in both reactions to measure gas products with a 1 ml air-tight gas syringe for analysis on a Shimadzu Tracera GC-2010 Plus gas chromatography fitted with a Carboxen1010 (30 m × 0.53 mm × 0.1 μm) column with a He carrier gas and Barrier Ionization Detector. Values reported for both reactions are the mean of 3 repeats. Mass-normalised photocatalytic activities are reported in all cases since this is the only means to enable quantitative benchmarking of different catalysts: photocatalytic reactions, in

common with all types of catalytic reactions, must always be performed in regimes wherein the rate of reactant conversion is directly proportional to the catalyst mass; if this is not the case, then the activity observed may reflect e.g. mass transport kinetics or self-absorption, rather than intrinsic reaction kinetics of the catalyst. The masses of different photocatalysts should not be arbitrarily optimised (for example focusing solely on maximising product yield), since this prohibits fair comparison of their intrinsic performance.

Results and discussion

Thermal evolution of mesoporous TiO₂

A series of meso-TiO₂ materials was synthesised by a one-pot surfactant templating method and thermal processing at temperatures spanning 300 to 700 °C, and characterised by a range of structural and photophysical methods. Wide angle XRD diffractograms for all meso-TiO₂ materials (**Fig. 1**) exhibited reflections indexed to (101), (004), (200), (105,211), (204), (116,220) and (215) planes of anatase crystallites (PDF no. 21-1272).[62] Higher temperature calcination at 500-700 °C also induced additional reflections characteristic of a crystalline rutile phase (PDF no. 21-1276). The relative rutile:anatase peak intensity increased with calcination temperature (from 3 % to 15 % between 500-700 °C, **Table 1**) consistent with the greater thermodynamic stability of rutile. Crystallites sizes were estimated from the Scherrer equation and evidenced sintering of both anatase (from $5\rightarrow31$ nm) and rutile (from $29\rightarrow46$ nm) particles with increasing calcination temperature. Unit cell parameters of the anatase phase estimated from the d₂₀₀ reflection of a=b=3.78 Å were in accordance with the literature (a=b=3.78 Å).[62]

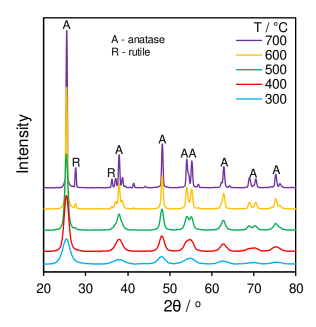


Fig. 1. Powder X-ray patterns of meso-TiO₂ as a function of calcination temperature.

Textural properties were evaluated by N_2 porosimetry (**Fig. 2a**), and revealed Type IV isotherms for all samples, characteristic of mesoporous materials with H2 type hysteresis loops indicating either restricted pore entrance (ink bottle shape) or an irregular channel structure of a mesopore framework.[63] The decrease in the hysteresis loop size and concomitant shift towards higher relative pressure with increasing calcination temperature evidence a progressive loss of mesoporosity, although hysteresis can still be clearly observed for the 700 °C calcined meso-TiO₂ sample. Sintering and a fall in mesopore volume with calcination temperature, were accompanied by a decrease in the BET surface area from 313 m^2 . g^{-1} for the 300 °C sample to only 19 m^2 . g^{-1} for meso-TiO₂-700C. TPO shows that the 300 °C calcined meso-TiO₂ had a high area despite retention of significant carbon from the F127 template within the mesopores (whose complete combustion required calcination \geq 400 °C, **Fig. S1**). Samples calcined between 300-500 °C exhibited narrow BJH mesopore size distributions (**Fig. 2b**), with higher temperatures expanding and broadening mesopores. Structural changes \geq 600 °C coincide with loss of both a well-defined d_{10} low angle XRD reflection, and hexagonal close-packed mesopore channels observed by HRTEM (**Fig. 3**). collectively the data indicate that nanocrystallite sintering induces the gradual collapse of ordered mesopores and coincident formation of large intraparticle voids. Mesopore diameters and anatase crystallite diameters of the 400 °C calcined meso-TiO₂ from HRTEM were in good agreement with values from porosimetry and wide angle XRD.

Table 1. Physicochemical properties of meso-TiO₂

Calcination temperature	BET area ^a	Mesopore size ^b	Mesopore volume ^b	TiO ₂ phase ^c	Crystallite size ^c /	Band gap ^d	Mass loss ^e
/ °C	$/ m^2.g^{-1}$	/ nm	/ g.cm ⁻³		iiii	/ eV	1%
300	313	4.6	0.41	Anatase	5	-	20 (5)
400	194	5.4	0.34	Anatase	7	3.07	6 (3)
500	126	6.5	0.28	97 % Anatase	11 (29)	3.02	3 (2)
				/3 % Rutile			
600	51	9.7	0.15	96 %Anatase	20 (35)	2.97	1 (1)
				/4 % Rutile	20 (33)		
700	19	16.0	0.11	85 % Anatase	21 (46)	2.94	0
				/15 % Rutile	31 (46)		

^aN₂ BET, ^bBJH, ^cXRD (rutile values in parentheses), ^dDRUVS, ^cTPO (mass loss between 150-350 ^cC in parentheses).

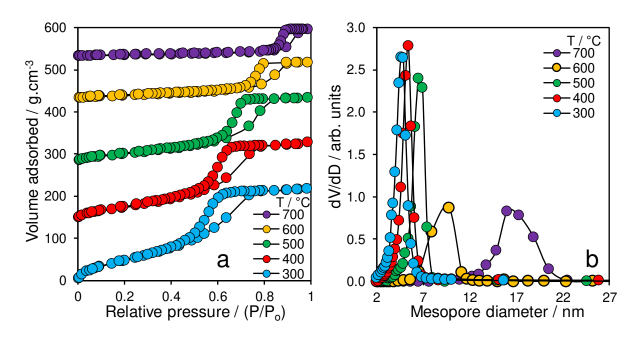


Fig. 2. (a) N_2 adsorption-desorption isotherms, and (b) corresponding BJH pore size distributions of meso-TiO₂ as a function of calcination temperature. Note that isotherms are offset for clarity.

DRUV spectra (**Fig. S2**) and associated Tauc plots (considering all materials as indirect band gap semiconductors due to the dominance of anatase) shown in **Fig. 4** demonstrate a strong trend between calcination temperature and optical properties of the meso-TiO₂ family. Besides the 300 °C sample, whose dark brown colour (due to ~15 wt% residual carbon) conferred strong UV-Vis absorption, meso-TiO₂ samples exhibited distinct absorption band edges between 350-415 nm which shifted to visible wavelengths with increasing calcination temperature. Corresponding optical band gaps (**Table 1**) decreased from 3.07 eV (400 °C) to 2.94 eV (700 °C) consistent with the coincident anatase→rutile phase transition.

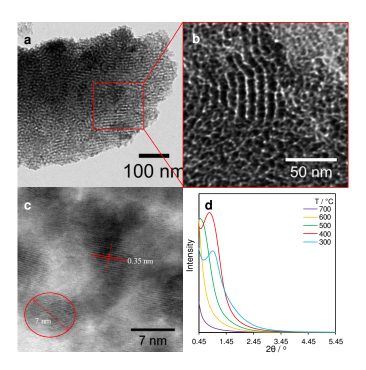


Fig. 3. (a-c) HRTEM images of 400 °C calcined meso-TiO₂, and (d) low angle XRD patterns of meso-TiO₂ as a function of calcination temperature.

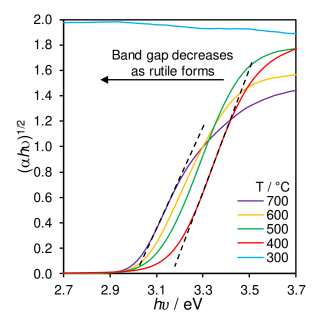


Fig. 4. Tauc plots of meso- TiO_2 as a function of calcination temperature.

Photocatalytic Methyl Orange depollution

Photocatalytic activity of meso-TiO₂ materials was subsequently evaluated for Methyl Orange (MO) degradation under UV-Vis irradiation (**Fig. 5**). It is important to recognise that Azo-dyes (exemplified by MO) are integral to the textile industry, and major sources of waste water contamination,[47] and hence their removal from aquatic environments is critical to improve human health. In the present work, dye degradation was investigated to target the removal of a specific Azo-dye commonly discharged to waste water in developing countries, and not simply to assay the general photocatalytic performance of our materials (Rochkind and co-workers published a recent critical review on this distinction [64]).

Prior to illumination, samples were stirred in the dark for 1 h to equilibrate MO adsorption: around 15 % of the initial dissolved MO adsorbed over the 300 °C calcined meso-TiO₂, with only ~3 % adsorbing over the other materials, and reaction data presented are corrected to discount this contribution. Negligible MO photolysis was observed over 5 h in the absence of titania. All meso-TiO₂ samples were active for MO photocatalytic degradation under UV-vis irradiation, with mass normalised initial rates increasing monotonically with calcination temperature (**Fig. 5b**). A plot of ln [MO] versus time reveals a linear dependence (**Fig. S3**) with a common rate constant k_{avg}=-0.0115 min⁻¹, characteristic of first order rate kinetics.[65, 66] Four distinct mechanisms are proposed for MO catalytic degradation over titania; a direct hole oxidation process[67] (**Equation 1**), hydroxyl radical oxidation[68, 69] (**Equations 2-4**), superoxide radical formation[70] (**Equations 5-9**) and ultra-fast direct electron injection into adsorbed MO[68] (**Equation 10**):

$$h_{VB}^{+} + dye \rightarrow dye^{\bullet +} \rightarrow dye$$
 oxidation products (1)

$$h_{VB}^{+} + H_2O \rightarrow H^{+} + OH^{-}$$
 (2)

$$h_{VB}^{+} + OH^{-} \rightarrow OH^{-}$$
 (3)

$$OH' + dye \rightarrow dye$$
 oxidation products (4)

$$e_{CB}^{-} + O_2 \rightarrow O_2^{-}$$
 (5)

$$O_2^{-} + H_2O \to HO_2^{-} + OH^{-}$$
 (6)

$$HO_2^{-} + H_2O \rightarrow H_2O_2 + OH^{-}$$
 (7)

$$H_2O_2 \rightarrow 2OH$$
 (8)

$$OH' + dye \rightarrow oxidation of the dye$$
 (9)

$$e_{CB}^{-} + dye \rightarrow degradation of the dye$$
 (10)

The pH of the reaction mixture rose slightly from 5.2 to 5.7 after 4 h reaction, consistent with hydroxyl radical generation during irradiation.

Since calcination simultaneously alters the structural and optical properties of mesoporous titanias, it is difficult to unequivocally identify the origin of the rate enhancement, with Fig. 5c evidencing a direct relationship between mesopore diameter and MO degradation, and concomitant inverse relationship with band gap. The former may be readily understood in terms of improved mass transport of the bulky dye into the mesopore network through Knudsen diffusion, for which the diffusion coefficient is proportional to pore diameter. However, the relatively weak dependence of this correlation suggests that MO (with a molecular diameter of 1.3 nm) can readily access even the smallest 5 nm mesopores.[71, 72] The far stronger rate dependence on band gap may simply reflect an exponential increase in (UV) photon absorption, or improved photoexcited charge transport and/or suppressed charge carrier recombination. Table 2 (Fig. S4) shows time-resolved photoluminescence data for meso-TiO₂ and P25 (whose anatase:rutile composition is closest to the 700 °C calcined sample). The average radiative lifetime ($\tau_{average}$) is fitted to arise from two separate components, designated τ_1 and τ_2 . $\tau_{average}$ increases with calcination temperature as a result of the increase in τ_2 (attributed to radiative relaxation processes originating from the indirect formation of self-trapped excitons) and therefore the overall increase in the average radiative lifetime is likely due to improved charge separation across a type-II, staggered anatase/rutile heterojunction interface.[73, 74] High-resolution TEM is required to directly visualise such a heterojunction, [75] however this is extremely challenging for our mesoporous titanias, which only contain <15 % rutile randomly dispersed in a non-uniform matrix. It is important to note that activities were unchanged under pure UV irradiation (<400 nm), indicating a negligible role for dye sensitisation in the photocatalytic mechanism (MO adsorbs strongly in the visible region, **Fig. S5**).[64, 76] Calcination >700 °C induced additional crystallite sintering (60-140 nm) and collapse of the mesopore framework, accompanied by an almost complete loss in photoactivity. HPLC analysis identified trace succinic and malic acid as reactively-formed intermediates during the first 2 h reaction, with fumaric acid the sole organic decomposition product after 4 h (**Figure S6**), equating to ~80 % mineralisation over the meso-TiO₂-500C photocatalyst.

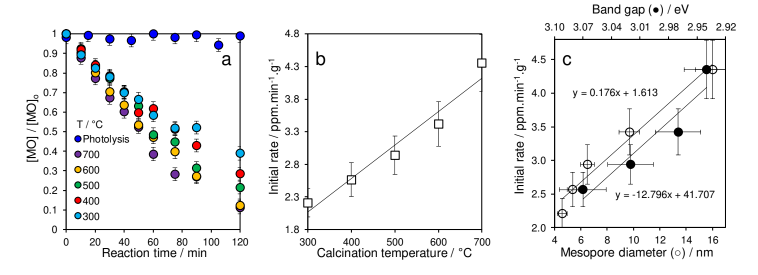


Fig. 5. (a) Reaction profiles for MO degradation, and corresponding initial rates of MO degradation as a function of (b) calcination temperature, and (c) mesopore diameter/band gap of meso-TiO₂.

Table 2. Time-resolved photoluminescence spectroscopy fitted parameters^a for meso-TiO₂ and reference materials

Sample	τ_1	B ₁ (%)	τ_2	B ₂ (%)	$ au_{average}$	χ^2
Sample	/ ns		/ ns		/ ns	
400 ^{b,c}	2.2	595 (83%)	8.5	121 (17%)	5.0	1.10
500 ^{b,c}	1.4	2435 (98%)	22.6	35 (2%)	5.4	1.10
700 ^{b,c}	2.2	381 (83%)	17.0	76 (17%)	11.2	1.10
P25 ^{b-d}	1.0	3190 (83%)	2.6	610 (83%)	1.5	1.17
$0.02 \text{ wt}\% \text{ Cu/meso-TiO}_2\text{-}500\text{C}^{\text{b,d}}$	5.6	2328	0	-	5.6	1.19
$CuO^{b,e}$	3.5	1019	0	-	3.5	1.15

^aParameters τ_1 and τ_2 are the calculated lifetimes of the two contributors to the radiative lifetime. B_1 and B_2 express the relative contribution of each component to the weighted average lifetime, $\tau_{average}$. Goodness of fit between model and recorded data express by χ_2 being close to unity. ^b380 nm excitation, ^c470 nm emission, ^d420 nm emission, ^e530 nm emission.

Photocatalytic H_2 production

Calcination of meso-TiO₂ exerted a similar effect on photocatalytic hydrogen production from water splitting under UV-Vis irradiation to that observed for MO degradation (**Figure 6**); H_2 productivity increased almost linearly with calcination temperature up to 700 °C. However, in this instance mesopore expansion cannot be responsible for the (dramatic) 25-fold rate-enhancement observed, since chemical mass transport of water will never be rate-limiting in aqueous solution. Rather, the increase in H_2 productivity with calcination temperature, which mirrors the concomitant decrease in band gap, must arise from the phase transition from anatase—rutile (quantified in **Table 1**). As discussed above for MO degradation, this phase transition is expected to result in a type-II staggered semiconductor heterojunction, thereby increasing charge carrier lifetimes and hence activity for any photocatalytic process. Furthermore, it is important to recall that the energetic position of the conduction band minimum (CBM) should be more negative than the reduction potential of water (-0.41 V vs SHE at pH 7) to produce H_2 ($2H^++2e^-\rightarrow H_2$).[77] Recent hybrid quantum chemical calculations and high-resolution XPS on titania nanostructures[74] and electrochemical Mott-Schottky analysis[78] reveals that the rutile CBM lies 0.2-0.4 eV more negative than that of anatase (typically \sim -0.4-0.65 eV vs SHE[79]). Rutile formation on meso-TiO₂ calcination will thus also provide a stronger thermodynamic driving force for H_2 evolution.

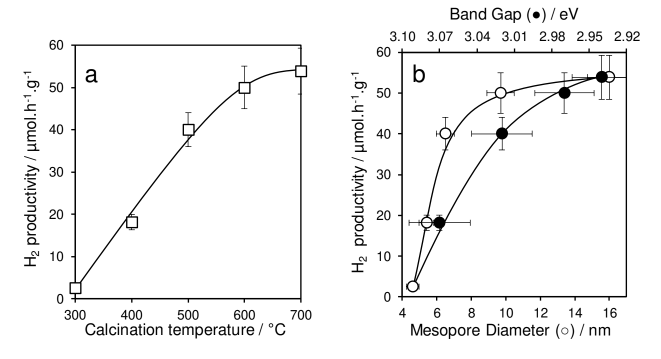


Fig. 6. H₂ production as a function of (a) calcination temperature, and (b) mesopore diameter/band gap of meso-TiO₂.

Cu functionalised mesoporous TiO₂

The impact of Cu deposition on the structural and photocatalytic properties of the preceding meso-TiO₂-500C was subsequently investigated; this support offering the optimal trade-off between textural properties and rutile content (**Table 1**), the latter desirable for heterojunction formation and enhanced photoactivity. Porosimetry evidenced minimal change in textural properties of the titania following post-synthesis functionalisation by copper at loadings between 0.02-12.33 wt% (**Table S1** and **Fig. S7a**): surface areas, mean mesopore diameters and mesopore volumes remained close to the parent meso-TiO₂ values demonstrating retention of the mesopore framework.

Wide angle XRD of the Cu/meso-TiO₂ materials (**Fig. S7b**) showed only pure anatase reflections for copper loadings between 0.02 and 8.34 wt%, with additional weak peaks at 35.7, 38.6 ° associated with (-111)/(002), (111)/(200) reflections characteristic of tenerite CuO crystallites (JCPDS card no. 45-0937) with a volume averaged diameter of 34 nm. Neither the anatase unit cell (a=b=3.78 Å) or crystallite size (11 nm) were changed following copper addition (**Table S1**), indicating that the latter was located predominantly extra-framework.

DRUVS spectra (**Fig. 7a**) and corresponding Tauc plots (**Fig. 7b**) demonstrate a systematic evolution in optical properties of the parent meso- TiO_2 with Cu functionalisation: the band gap absorption edge progressively increased from 350 \rightarrow 430 nm with increasing Cu loading; this was accompanied by increased absorption between 400-600 nm possibly due to interface states arising from charge transfer between TiO_2 and Cu(I)/Cu(II) species, and the emergence of a new absorption band spanning 580-900 nm associated with d-d transitions in CuO[80] emerges for loadings \geq 0.3 wt%. These

changes are reflected in a systematic decrease in the band gap from 3.00 eV for the parent titania to only 1.10 eV for the 12.33 wt% Cu sample (**Table S1**), consistent with the formation of bulk CuO[81] in accordance with XRD.

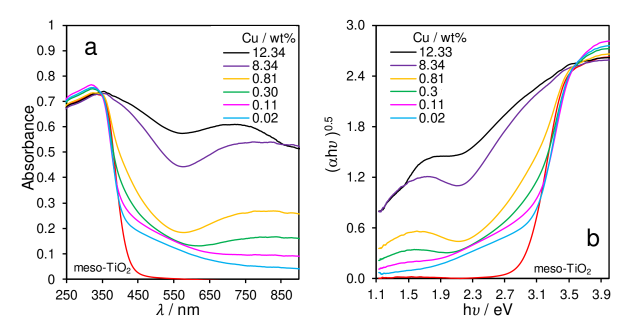


Fig. 7. DRUV spectra and corresponding Tauc plots of Cu/meso-TiO₂-500C as a function of Cu loading.

Copper co-catalysts may promote TiO₂ by various means, including trapping of charge carriers at the semiconductor surface and/or electron injection into the titania conduction band by metallic nanoparticles,[82-84] heterojunction formation between discrete copper oxide and titania nanoparticles,[85, 86] or framework substitution of copper cations accompanied by band energy modification and/or improved charge transport.[87, 88]

Identification of the copper chemical environment within the Cu/meso-TiO₂ series is thus critical to interpreting their photocatalytic performance. However, the extremely low Cu loading within some samples, and similar binding energies of Cu 2p metal and (I) oxide photoemission features, prohibits their quantification and discrimination by laboratory X-ray sources.[89] Hence the local chemical environment and oxidation state of Cu species was probed by Cu K-edge XAS. Background subtracted, normalised XANES spectra of the 0.02-12.33 wt% Cu/meso-TiO₂ series and Cu₂O, CuO and nitrate references are shown in **Fig. 8a-b**. The copper local environment evolves from that reminiscent of the Cu(II) nitrate trihydrate complex to that of bulk CuO with increasing Cu loading, with spectra exhibiting three distinct features: a weak pre-edge feature around 8977 eV, a strong rising-edge feature around 8986 eV, and an intense white line around 8996; these are attributed to dipole-forbidden $1s\rightarrow 3d$ (not observed for Cu(I) species), $1s\rightarrow 4p_z$ (or 4px, ypz for Cu(I) species) and $1s\rightarrow 4p$ (continuum) transitions respectively. Least squares spectral fitting to Cu, Cu₂O, CuO and Cu(NO₃)₂ references suggested only Cu(II) were present in all Cu/meso-TiO₂ samples (**Fig. S8**) with loadings ≥ 0.81 wt% a good fit to CuO, and those < 0.3 wt% an excellent fit to a mononuclear copper species. However, **Fig. 8b** reveals that the pre-edge feature characteristic of Cu(II) only emerges for Cu loadings ≥ 0.3 wt%, and hence copper atoms in the ultra-dilute Cu/meso-TiO₂ materials appear electronically similar to Cu(I) but structurally similar to those in mononuclear

complexes. Fitting of the corresponding EXAFS spectra of the 0.1 wt% Cu/meso-TiO₂ sample (**Fig. S9**) revealed only Cu-O scatterers around 1.94 Å, with no Cu-Cu interatomic distances (**Table S2**), consistent with isolated Cu(I) species either within the titania framework or decorating the semiconductor surface. The Cu-O nearest neighbour coordination number of 2.5 (versus 4 for both CuO and Cu(NO₃)₂) for 0.1 wt% Cu/meso-TiO₂ is consistent with either anion vacancy formation upon framework substitution of Ti(IV) with Cu(I), or atomically-dispersed, undercoordinated Cu(I) atoms at the titania surface. Since copper was introduced by post-modification of mesoporous titania, employing a comparatively low temperature processing step, the latter scenario is more feasible. The constant band gap, and anatase lattice parameter and crystal size, following the addition of ultra-low Cu concentrations to the parent meso-TiO₂ supports the hypothesis that single copper atoms are dispersed over titania. Time-resolved photoluminescence of the 0.02 wt% Cu/meso-TiO₂ highlight a significant increase in charge carrier relaxation time τ_1 relative to the parent 500 °C calcined titania upon copper functionalisation (**Table 2**), possibly due to isolated Cu(I) species acting as hole traps.

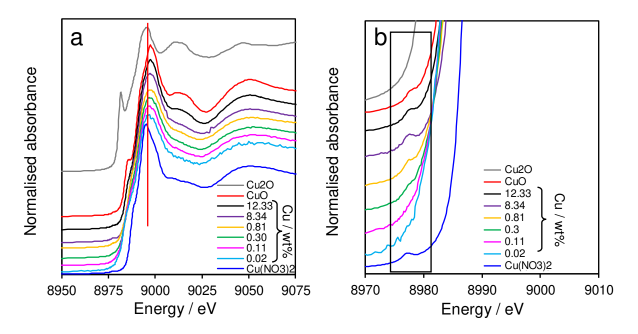


Fig. 8. (a-b) Cu K-edge XANES of Cu/meso-TiO₂-500C as a function of Cu loading.

The impact of copper on the photocatalytic performance of mesoporous titania for MO photodegradation under UV-Vis irradiation is shown in **Fig. 9**. Mass normalised initial rates of dye degradation were inversely proportional to copper loading (and bulk CuO content determined by XANES fitting), but directly proportional to the concentration of isolated Cu(I) species (from XANES). We attribute the latter relationship to the creation of oxygen vacancies upon introducing Cu(I) species into the titania framework; these oxygen vacancies likely capture photoexcited electrons to form F-centres,[90] thereby suppressing charge carrier recombination, promoting either direct hole oxidation or indirect hydroxyl oxidation mechanisms (**Equations 1** or **2-4**). Dye degradation via a hydroxyl radical mechanism (**Equation 2**) requires photoexcited holes in the valence band with energies >2.6 eV. This threshold is exceeded by the valence band maxima of all Cu/meso-TiO₂ catalysts (determined as ~2.72-2.95 eV from valence band spectra in **Fig. S10**), and hence MO degradation may occur via oxidation by hydroxyl radicals. For loadings above 0.81 wt% Cu, photoactivity fell to that

of the parent 500 °C calcined meso-TiO₂ support, indicating negligible benefits from CuO despite its impact on optical properties. The 0.02 wt% Cu/meso-TiO₂ catalyst also exhibited excellent stability, with negligible loss in photodegradation activity over three consecutive recycles, consistent with preservation of Cu(I) isolated sites from XANES analysis (**Fig. S11**). CuO displays poor performance as a promoter, probably due to its smaller valence band maximum relative to the meso-TiO₂-500C anatase parent (~2.66 eV versus 2.95 eV [91]) which offers a lower thermodynamic driving force for catalytic oxidation, and the large size of CuO nanoparticles (34 nm for 12.33 wt% Cu) and corresponding poor interfacial contact and inefficient heterojunction formation with titania. It is interesting to note that only the 12.33 wt% Cu/meso-TiO₂, for which crystalline CuO was observed, was active under visible light irradiation (>400 nm, 8 mW cm⁻²). This confirms that Cu(I) species did not significantly modify the band gap of the titania support, and that the poor observed initial rate of only 0.244 ppm.min⁻¹.g⁻¹ arose solely from the visible absorbing CuO phase.

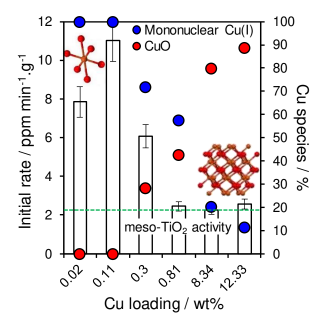


Fig. 9. Correlation between initial rate of photocatalytic MO degradation and Cu species (from XANES) in Cu/meso-TiO₂-500C. Dotted line indicates activity of parent meso-TiO₂-500C.

In contrast to MO degradation, photocatalytic H₂ production over in Cu/meso-TiO₂-500C exhibited a volcano dependence on copper loading (**Fig. 10**), reaching a maximum for 0.81 wt% Cu, a material comprising ~60 % mononuclear Cu(I) species and 40 % CuO. The lack of promotion at low Cu loadings, wherein only Cu(I) species exist, is understandable in terms of the preceding hypothesis that Cu(I) doping introduces F-centers into the meso-TiO₂ framework (which trap photoexcited electrons required for H₂ production). Similarly, the baseline activity of the 12.33 wt% Cu/meso-TiO₂-500C which is dominated by crystalline CuO is also explicable, since the CBM of CuO lies ~0.96 eV, significantly below that necessary to reduce protons,[91] and therefore cannot contribute directly to H₂ production as a semiconductor photocatalyst. The origin of the synergy between Cu(I) and CuO responsible for the 3-4-fold rate-enhancement observed for the 0.81 wt% Cu/meso-TiO₂-500C remains unclear, but highlights the impact of subtle variations in the composition of doped titania on corresponding photophysical/catalytic properties.

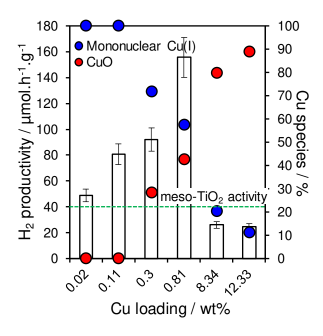


Fig. 10. Correlation between H₂ productivity and Cu species (from XANES) in Cu/meso-TiO₂-500C. Dotted line indicates activity of parent meso-TiO₂-500C.

Conclusions

A facile route is demonstrated to high surface area, mesoporous TiO₂ in which the pore diameter, anatase:rutile phase composition and corresponding photophysical properties may be readily tuned through thermal processing of a surfactanttemplated sol-gel, and whose performance may be dramatically enhanced through the incorporation of single atom Cu (I) species. Increasing calcination temperature from 300 \rightarrow 700 °C induces progressive expansion of uniform mesopores from 5→16 nm respectively, and a concomitant decrease in surface area, accompanied by the transformation of smaller $(11\rightarrow31 \text{ nm})$ anatase to larger $(30\rightarrow46 \text{ nm})$ rutile nanocrystallites) and slight band gap narrowing. These structural and electronic changes strongly influence photocatalytic activity for Methyl Orange degradation under UV-Vis irradiation. Initial rates of photodegradation exhibit a weak, direct correlation with mesopore diameter and strong inverse (direct) correlation with band gap (rutile concentration), imparting a two-fold rate enhancement through high temperature calcination. Photocatalytic H₂ production is even more sensitive to the anatase→rutile phase transition, which confers a dramatic 25-fold increase in water splitting, likely due to a combination of heterojunction formation and the higher conduction band minimum of rutile titania. Post-functionalisation of mesoporous TiO₂ by wet-impregnation with ultralow (≤ 0.1 wt%) concentrations of copper affords atomically-dispersed Cu(I) species, likely extra-framework decorating the surface of titania crystallites. Despite their high dilution, the incorporation of Cu(I) species results in a six-fold rate enhancement for Methyl Orange photodegradation relative to the unmodified (500 °C) mesoporous TiO₂, possibly due to the introduction of oxygen vacancies which may act as electron traps to increase charge carrier lifetimes. A synergy between Cu(I) and CuO over 0.81 wt% Cu/meso-TiO₂-500C increased photocatalytic H₂ production four-fold relative to the mesoporous anatase scaffold. The importance of tailoring both textural and photophysical properties of heterogeneous

photocatalysts to optimise chemical mass transport, and the energetics and lifetime of charge carriers, emerge as key design features.

Acknowledgements

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