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Polyacrylamide exotemplate-assisted synthesis of hierarchically porous nanostructured $TiO₂$ macrobeads for efficient photodegradation of organic dyes and microbes†

Muhammad Ahmad Mudassir[,](http://orcid.org/0000-0002-6624-007X) D^{abc} Syed Zajif Hussain, D^a Mishal Khan, a e Syeda Tasmia As[ma,](http://orcid.org/0000-0001-5498-1236)^{ad} Zafar Iqbal[,](http://orcid.org/0000-0001-5142-5824)^a Zille Huma,^a Najeeb Ullah,^f Haifei Zhang, D^c Taria Mahmood Ansari^{*b} and Irshad Hussain **D**^{*af}

Nano/microscale TiO₂ materials and their composites have reached the pinnacle of their photocatalytic performances to destroy persistent organic pollutants and waterborne microorganisms, but their practical applications are limited by the drawbacks associated with their stability, leaching, processing and separation. To overcome these shortcomings, we have prepared hierarchically porous nanostructured TiO₂ macrobeads via an exotemplating or nanocasting strategy by infiltrating the TiO₂ sol into the emulsion-templated porous polyacrylamide scaffold followed by its gelation, drying and calcination. The nanoscale $TiO₂$ building units tailor the shape of the porous polymeric network after calcination thereby retaining the macroscale morphology of polymer beads after template removal. A novel combination of the hierarchical macroporosity, orderly crystalline anatase nature, nanoscale feature and good surface area revealed by the relevant characterization tools makes these TiO₂ scaffolds particularly effective for superior degradation of methylene blue with the enhanced rate constant and efficient disinfection of E. coli and S. aureus under UV light. The macrosize and mechanical stability of these purely TiO₂ beaded architectures have several potential advantages over conventional TiO₂ nanocomposites and slurry systems to address the inherent bottlenecks of secondary contamination, difficult operation and energy-intensive post-recovery processes that are indeed deemed to be the barriers to develop practically useful water treatment technologies. PAPER
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a Department of Chemistry & Chemical Engineering, SBA School of Science & Engineering (SBASSE), Lahore University of Management Sciences (LUMS), Lahore-54792, Pakistan. E-mail: ihussain@lums.edu.pk

b Institute of Chemical Sciences, Bahauddin Zakariya University (BZU), Multan-60800, Pakistan. E-mail: drtariq2000@gmail.com

Department of Chemistry, University of Liverpool, Oxford Street, Liverpool-L69 3BX, UK

d Institute of Industrial Biotechnology (IIB), GC University, Lahore-54000, Pakistan e Preston Institute of Nano Science & Technology (PINSAT), Islamabad, Pakistan f US-Pakistan Centre for Advanced Studies in Energy (USPCAS-E), University of Engineering & Technology (UET), Peshawar, Pakistan

† Electronic supplementary information (ESI) available: UV-vis diffuse reflectance spectra and plot of Kubelka–Munk function versus photon energy for bandgap measurement of PAM-TiO₂ NC and porous NS TiO₂ macrobeads sample; SEM images of PAM and PAM-TiO₂ NC beads; pore and particle size distributions of PAM, porous PAM-TiO₂ NC and porous NS TiO₂ macrobeads; removal efficiency (mg g^{-1}) of porous NS TiO₂ macrobeads without and with H₂O₂; time-wise bactericidal efficiencies (%) of 0.2 \rm{mg} \rm{mL}^{-1} dosages of PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads against *S. aureus* and *E. coli* under UV light and in the dark; fluorescence spectra of the irradiated TA and $TA-TiO₂$ samples to probe \cdot OH formation; SEM image of a single porous NS TiO₂ macrobead and the digital photo of porous NS $TiO₂$ macrobeads soaked in MB solution; a comparison for the removal of MB by different TiO₂-based materials. See DOI: 10.1039/c8ra06197a

Introduction

 $TiO₂$ -based materials have got vital importance in multidisciplinary research areas due to their large surface area, greater surface activity, long-term stability, low-cost, non-toxicity, earth abundance, unique physico-chemical properties, and so forth. Based on their excellent optical, electrical, thermal, mechanical and chemical properties, titania nanomaterials have extensively been used in electronics, photonics, energy production, biomedical engineering, deactivation of bioparticulates, gas sensing, air treatment, and photocatalytic degradation of inorganic as well as organic pollutants.¹–⁵

Pertaining to its tremendous role in environmental applications, TiO₂ has previously been utilized in different sizes from the nanometer to the micrometer range. 6 In the case of TiO₂ nanoparticles (NPs) prepared via different routes, the targeted goals have partially been achieved just by tuning their size, morphology, nitrogen doping and surface chemistry. However, high surface energy and small size of $TiO₂$ NPs undermined their handling and separation during and after their use for the wastewater remediation.^{7,8}

To circumvent the post-separation complications, $TiO₂ NPs$ have been immobilized onto the rigid supports comprised of glass,⁹ quartz,⁸ stainless steel,7 silica,¹⁰ alumina,¹¹ polymer,¹² carbon,¹³ clay and various zeolites¹⁴ by using electrophoretic deposition,⁸ dip-coating¹⁵ and electrospinning¹⁶ techniques, and so forth. However, the photocatalytic activity of $TiO₂$ was observed to be affected by the adsorptive nature of the support,¹⁷ and the possible reduction in the available surface-tovolume ratio.¹⁸

Apart from using the substrate-supported $TiO₂$ NPs, the mesoporous TiO₂ nanocrystal clusters,¹⁹ fiber-like²⁰ and necklace-structured²¹ TiO₂–SnO₂ hybrid nanofibers, Au/TiO₂ nanoforests, and $NaYF_4:Yb,Tm@TiO_2$ core–shell nanomicrospheres have also been designed for their specific morphologydependent recovery.^{22,23} Together with their one-dimensional morphology, these hybrid nanofibers have been doped with Fe to add magnetic character for making their separation easier.²⁴ Furthermore, the efficiency of such magnetic photocatalysts was boosted by introducing Ag NPs, that is generally helpful in excitation of electrons, visible light photocatalytic activity and separation of electron–hole pairs.25,26

The direct contact of magnetic iron oxide with $TiO₂$ under particular conditions may cause photodissolution,²⁷ promote transfer of e^{-}/h^{+} from TiO₂ to iron oxide particles, and lead to Fabry–Perot oscillations.²⁸ In the same way, the direct interaction of chemically reactive Ag with $TiO₂$ may oxidize Ag into AgO at the TiO₂-Ag junction. In order to avoid these problems, the iron oxide and Ag cores were coated by the non-toxic and hydrophilic inorganic or organic materials including $SiO₂$ and carbon.²⁸ For instance, the $(\gamma Fe_2O_3@SiO_2)_n@TiO_2$,²⁹ Ag/SiO₂-
TiO_c composites ³⁹ and triple shelled Ag@Ee O @SiO @TiO $TiO₂$ composites,³⁰ and triple-shelled Ag@Fe₃O₄@SiO₂@TiO₂ nanospheres²⁸ have been designed and used for the photodegradation of methylene blue. But achieving strong binding of $TiO₂$ onto the $SiO₂$ was a challenge that, somehow, was addressed in $Fe₃O₄/SiO₂/TiO₂$ composites by polyacrylic acid treatment.³¹

In the microscale size regime, hollow $TiO₂$ sphere-in-sphere³² structures with urchin-like morphology have been used for the photodegradation of phenols. An efficient photocatalysis of rhodamine B has also been achieved by the multi-shelled $TiO₂$ hollow spheres.³³ Likewise, the nacre-like nanolignocellulosepoly(vinyl alcohol)–TiO₂ composite microstructures have been used for photodegradation of methyl orange.³⁴ Whereas, the hierarchical flower-like $Co_{3-x}Fe_{x}O_{4}$ hollow microspheres and the hierarchically porous Fe₃O₄@SiO₂@TiO₂@Pt core-shell microspheres presented improved photocatalytic degradation of methylene blue and rhodamine B together with their rapid magnet-assisted separation.^{35,36} Moreover, a polymer-templatedirected TiO₂ and the mixed TiO₂/metal oxide (Al, Ga and In) microspheres have been developed for the efficient photocatalytic decomposition of 2-chlorophenol.³⁷

The isolation of the photocatalysts has become more feasible with the help of newly developed $TiO₂$ spheres (larger in size than 1000 µm) which are also more effective in photoconversion of $CO₂$ to $CH₄$.³⁸ From this perspective, we have prepared hierarchically porous nanostructured (NS) $TiO₂$ macrobeads by applying a well-established nanocasting technique by using porous polyacrylamide (PAM) beads as the exotemplates.^{39,40} These $TiO₂$ beads are highly UV-active and very efficient for the degradation of methylene blue and inactivation of E. coli and S. aureus bacteria. Notably, these can be handled and separated conveniently without requiring any external magnetic field and cumbersome separation processes, and thus alleviates the chances of secondary contamination caused by the photocatalysts themselves during scaled-up water treatment.

Experimental

Materials

Titanium(IV) isopropoxide (TTIP, 97%) acrylamide (AM, \geq 99%), bis-acrylamide (BAM, 99%), ammonium peroxydisulfate (AP, \geq 98.0%), 1,2-bis(dimethylamino)ethane (TEMED, 99%), poly(vinyl alcohol) (PVA, 98–99%), polyethylene glycol tert-octylphenylether (triton X-405, 70%), terephthalic acid (TA, 98%), sodium hydroxide (NaOH, 97%) and methylene blue (MB, \geq 82%) were purchased from Sigma-Aldrich. Potassium chloride (KCl, 100%) and hydrogen peroxide (H_2O_2) were obtained from Fischer. Glacial acetic acid (AA, 100%) was purchased from Labscan. Ethanol (analytical grade), n-hexane (95%), acetone (99.50%), 2-propanol (\geq 99.5%) and LB (Luria–Bertani) nutrient broth were procured from Merck. The cooking oil (sunflower) was of the commercial grade purchased from a local market in Lahore. The Escherichia coli [ACCN, KJ880039] and Staphylococcus aureus [ACCN, KY635411] strains, and deionized water purified by Milli-Q Plus system (Millipore, Bedford, MA, USA) was used for the whole research study. Paper

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Methods

Synthesis of PAM beads. The crosslinked PAM beads were synthesized by double $(O/W/O)$ emulsion polymerization^{41,42} with some modifications. The clear solution of AM (0.96 g) , BAM (0.24 g) and PVA (0.06) was prepared in deionized water (3 mL) and evenly mixed with a triton X-405 (650 μ L) solution. An oil phase (6 mL) containing TEMED (0.016 g) was added to the aqueous monomer solution and mixed well under stirring at 160 rpm with an overhead paddle stirrer to get a stable oil-inwater (O/W) emulsion. The oil-in-water emulsion polymerization was initiated by the addition of aqueous APS solution (0.03 g in 600 μ L), transferred dropwise into the hot (80–90 °C) oil phase, and baked for about 3 h to complete oil-in-water-in-oil (O/W/O) emulsion polymerization. The PAM beads were filtered out of the oil, washed with n -hexane (24 h) followed by acetone (thrice for 1 h) and isopropanol (overnight), and dried in oven at 85 °C overnight.

Preparation of PAM-TiO₂ NC beads. The sol-gel process was used to prepare PAM-TiO₂ nanocomposite (NC) beads by impregnation and gelation of $TiO₂$ sol into the macroporous PAM beads.⁴³ To prepare a TiO₂ sol, TTIP (0.3 mL) was hydrolysed with slow mixing of AA (5.11 mL) followed by dropwise addition of water (6.2 mL in 1 h) under vigorous stirring which continued further for 3 h. The resultant suspension was sonicated for about 30 min to get a clear solution. The PAM beads were soaked into the freshly prepared $TiO₂$ sol and shaken gently for about 3 d, separated by a Buchner funnel, kept undisturbed in the dark for about 24 h to complete the nucleation process, placed in an oven at 70 \degree C for 12 h, and subsequently vacuum-dried at 100 °C. The same procedure of soaking, gelation and drying was again repeated twice to achieve higher mechanical stability by the maximum loading of $TiO₂$.

Designing hierarchically porous NS $TiO₂$ macrobeads. Porous TiO₂ beads were obtained by calcining PAM-TiO₂ NC in air at 520 °C for 5 h with temperature ramp rate of 1 °C min⁻¹ $(Fig. 1).^{40,44,45}$

Characterization

Attenuated total reflectance-infrared spectrometer (Alpha, Bruker) was used to record FTIR spectra while taking 24 scans at a resolution of 4 $\rm cm^{-1}.$ Thermal analyser (TA Instruments, Q600-SDT) was used to perform TGA analysis under an inert (N_2) gas) atmosphere (flow rate, 50 mL min $^{-1})$ in temperature range of 25–1000 °C (ramp rate, 10 °C min^{-1}). X-ray diffractometer (D2 Phaser, Bruker) with CuK α radiations ($\lambda = 1.54$ Å, 30 kV, 10 mA) was used to determine XRD pattern in the range of $20-80^\circ$ (scan rate, 1 $^{\circ}$ min $^{-1}$). The surface area analyser (3Flex, Micromeritics ASAP 2420) was used to measure the Brunauer–Emmett–Teller specific surface area by using the N_2 sorption isotherms at 77 K. Field emission electron microscope (FEI, Nova NanoSEM) was used to obtain electron micrographs of the samples mounted on aluminium stubs with conductive carbon adhesive stickers. ImageJ software was used to measure the average particle size of $TiO₂$ nanobuilding units (NBUs) and the pore size distribution of beads. Zeta potential measurements were performed by the Zetasizer Nano ZSP (Malvern Instrument Ltd., UK). The absorption and diffuse reflectance spectra of porous NS $TiO₂$ beads samples were taken with the ultraviolet-visible (UV-vis) double beam spectrophotometer (JASCO, V-770) equipped with a solid sample accessory (Integrating Sphere, ILN-925). The experiments for the degradation of methylene blue and inactivation of E. coli and S. aureus were carried out in the dark, and under simulated sunlight (100 W xenon lamp) and UV-A light (UV irradiation chamber equipped with UV-A lamps) conditions. The catalytic performance of porous NS $TiO₂$ macrobeads was quantitatively analysed by ultraviolet-visible light spectrophotometer (Shimadzu UV-1800). The fluorescence emission spectra were recorded on a multilabel plate reader PSC Advances

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Fig. 1 Schematic illustration of exotemplate-directed preparation of hierarchically porous $NS TiO₂$ macrobeads.

PAM bead

shaking

nucleation
aggregation

gelation
drying

Optical properties evaluation

The optical absorption properties were analysed by the absorption spectra of the PAM, PAM-TiO₂ NC and NS porous $TiO₂$ macrobeads samples to screen the absorption edges for optimizing the hydroxyl radicals' generation in order to achieve efficient degradation of methylene blue and bacteria. Furthermore, the diffuse reflectance measurements were used to estimate the indirect bandgap of porous NS $TiO₂$ beads sample by extrapolating the tangent line in the plot of Kubelka–Munk function $[F(R)hv]^{0.5}$ versus photon energy (E_g) .

Photocatalytic performance of porous NS $TiO₂$ macrobeads

Degradation of methylene blue. In a typical experiment, each of PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads were separately placed in 10 mL of 25 ppm MB solution at pH 10 and kept in the dark for 1 h to attain adsorption–desorption equilibrium followed by turning on light (simulated solar and UV-A) further for 1 h under the identical condition of continuous shaking (250 rpm) at room temperature in the absence as well as presence of $H₂O₂$ (0.03 M). The absorbance of MB was monitored at λ_{max} of 661 nm as a function of time (0, 12, 24, 36, 48, and 60 min) by the UV-visible spectrophotometer to determine its concentration using the calibration curve. All experiments were performed in triplicate and the error bars used to denote the standard deviation. The degradation efficiencies were demonstrated in both mg g^{-1} (using 3 mg of TiO₂ beads) and % (using 5 mg each of PAM, PAM-TiO₂ NC and porous TiO₂ beads). The reaction rate constant was deduced by the slope of the straight line through zero from the plot of $ln(C_0/C)$ versus illumination time using the following pseudo-first-order kinetics, *i.e.*, $ln(C_0/C) = kt$, where the ln is the logarithm of natural base, C_0 and C are initial and final concentrations of MB, t is the illumination time and k represents the pseudo-firstorder reaction rate constant.

Disinfection of E. coli and S. aureus. To perform the antibacterial activity tests of PAM, PAM-TiO₂ NC, and porous NS $TiO₂$ macrobeads, pure cultures of two representative Gramnegative and Gram-positive bacteria (E. coli and S. aureus, respectively) were cultivated in the fresh sterile LB (Luria–Bertani) broth media. The exponentially growing bacterial cells of both bacterial cultures were harvested by centrifugation at 5000 rpm for 5 min and the bacterial dead cell mass was washed off with normal saline (NS). The washed bacterial cells were suspended in NS, and serially diluted to adjust bacterial concentration at 10⁷ CFU mL⁻¹. The 10 mL of each bacterial cell suspension was then added into two separate sets of sterilized flasks containing PAM, PAM-TiO₂ NC and porous NS TiO₂ beads.

Together with their controls, one set of all flasks in the dark while the other set in a UV chamber, were incubated at 37 $^{\circ}$ C with a constant shaking speed of 150 rpm. After regular time intervals (10-60 min), 100 μ L aliquot of suspension mixture was withdrawn from each flask of both sets (dark and UV), plated

TiO₂ sol

 $AA + H₂O$

stirring

onto the freshly prepared sterile LB plates, and incubated at 37 °C for 24 h. The number of grown colonies in each LB plate was enumerated and compared with its corresponding control. To determine the effect of higher dosages, the same procedure was also repeated for 0.4 and 0.8 mg mL^{-1} of the PAM, PAM- $TiO₂ NC$ and porous $TiO₂$ beads in the dark as well as under UV light. All experiments were carried out in triplicate and their average values are presented.

Probing \cdot OH generation

A simple fluorescence-based approach involving chemical reaction of TA with \cdot OH to form fluorescent hydroxyterephthalic acid (TAOH) was used to justify the formation of \cdot OH during photocatalysis by porous $TiO₂$ beads.⁴⁶ To perform it, an aqueous solution (3.5 mL) containing 0.01 M NaOH, 0.1 M KCl, 0.03 mM $H₂O₂$ and 3 mM TA (without TiO₂), and another aqueous suspension (3.5 mL) containing 0.01 M NaOH, 0.1 M KCl, 0.3 mM $H₂O₂$, 3 mM TA and 1 mg of fine TiO₂ powder were prepared and placed under UV-A chamber under identical conditions of stirring for only 15 min. Subsequently, the fluorescence spectra for the UV-irradiated samples were recorded (excitation wavelength $=$ 325 nm, emission wavelength $=$ 426 nm) and compared.

Results and discussion

Synthesis

A bottom-up approach was used to hierarchically organize porous NS TiO₂ macrobeads from the sol-gel processing of easily condensable molecular precursor in the presence of porous PAM macrobeads prepared via an oil-in-water-in-oil polymerization. The slow hydrolysis of TTIP precursor produced the colloidal $TiO₂$ NPs. These nanometer-sized colloidal TiO₂ sol particles were then infiltrated into the pores and channels of the emulsion-templated porous PAM beads by a simple soaking method with constant mechanical shaking. The impregnated $TiO₂$ NPs tailored the shape of PAM pores and channels, progressively nucleated and grown during the process of gelation. The subsequent drying of the resultant material removed the trapped solvent, solidified the wet $TiO₂$ gel, and evolved a compact network of $TiO₂$ inside of PAM beads. As a result of calcination, PAM was removed and the remaining TiO2 retained the proxy structure of hierarchically porous macrosized (0.6–1 mm) beads (see Fig. S-7A†). These NS $TiO₂$ macrobeads show good mechanical stability while maintaining their integrity in water and MB aqueous solution even after the continuous shaking at 250 rpm (see Fig. S-7B \dagger). These specific properties are highly demanding and effective for the practical applications of such materials.

Characterization

The UV-visible absorption spectra were recorded to evaluate the photoactive nature of the PAM, PAM-TiO₂ NC and porous NS $TiO₂$ macrobeads (see Fig. 2(A)). The absence of any characteristic absorption band in the PAM beads sample spectrum shows its inactivity under UV-visible light. The appearance of broader absorption band in PAM-TiO₂ NC beads sample confirms

Fig. 2 (A) UV-vis absorption, and (B) FTIR spectra of (a) PAM, (b) PAM- $TiO₂$ NC and (c) porous NS $TiO₂$ macrobeads.

successful incorporation of Ti species into the PAM beads.⁴⁷ However, an absorption band edge at around 380 nm in porous $TiO₂$ beads sample may be associated with the charge transfer properties⁴⁸ and the anatase phase of the $TiO₂$.⁴⁹ Furthermore, the bandgaps assumed by an indirect transition between valence and conduction bands of PAM-TiO₂ NC and porous TiO₂ bead were estimated to be 3.44 and 3.10 eV, respectively (Fig. S- $1B[†]$), using reflectance (%) data (Fig. S-1A \dagger).

The chemistry of PAM, PAM-TiO₂ NC and porous NC TiO₂ macrobeads was verified by the FTIR (see Fig. 2(B)). In transmission spectra of PAM beads sample, the signals at 3342 cm^{-1} and 3192 cm^{-1} can be ascribed to the asymmetric and symmetric NH₂ stretching, respectively, while the appearance of the peaks at 2944 cm^{-1} and 1451 cm^{-1} denotes the stretching and bending vibrations of $CH₂$ group, respectively.⁵⁰ An intense peak at 1649 cm⁻¹ may be attributed to the C=O stretching, whereas the shoulder at 1606 cm^{-1} and a weak signal at 1525 cm^{-1} can be assigned to N-H plane bending.^{51,52} The C-N stretching vibration at 1311 cm^{-1} may be consigned to the amine group. In the case of PAM-TiO₂ NC beads sample, the suppression of PAM peaks verifies its loading with $TiO₂$. Moreover, the inaccessibility of all PAM signals in calcined porous NS TiO₂ beads sample may not just justify its removal, but a highly intense signal at 494 cm^{-1} is considered as the signature of the characteristic Ti-O-Ti bonds.^{53,54} Paper

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The PXRD patterns were recorded to determine the structural properties of PAM, PAM-TiO₂ NC and porous TiO₂ beads. The flat lines with small humps were appeared in PAM and PAM-TiO₂ NC beads due to the absence and presence of amorphous $TiO₂$ respectively. However, the amorphous nature of PAM-TiO₂ NC beads was changed by removing PAM and converting the remaining amorphous $TiO₂$ into orderly crystalline phase at once through calcination. The well-resolved peaks originating from (101), (004), (200), (105), (211), (204), (116), (215) and (220) planes correspond to the anatase phase (JCPDS: $00-002-0406$) in calcined porous NS TiO₂ macrobeads sample (see Fig. 3(A)).

The thermal behaviour of PAM, PAM-TiO₂ NC and porous $TiO₂$ beads was analysed by the TGA (see Fig. 3(B)). A step-wise weight losses of about 12 and 8% at around 25-200 $^{\circ}$ C in PAM and PAM-TiO₂ NC beads samples, respectively, display the evaporation of the physisorbed solvents, accompanied by subsequent weight losses of about 85% at \sim 565 °C and 99% at \sim 650 °C due to the removal of chemisorbed water and organics in PAM and PAM-TiO₂ NC beads samples, respectively.⁴⁸

Fig. 3 (A) PXRD patterns, and (B) TGA curves of (a) PAM, (b) PAM-TiO₂ NC and (c) porous NS TiO₂ macrobeads.

However, no significant weight loss up to \sim 990 °C in porous TiO2 beads sample advocates its purity and thermal stability. Additionally, these results also provide an estimate of the loading of $TiO₂$ into the PAM matrix.

BET surface areas of PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads were mainly concomitant with their pore size distributions. A lower surface area (4.14 $\mathrm{m^2\,g^{-1}}$) of PAM beads was associated with vast majority of macropores ranging from \sim 5 to 295 µm [Fig. S-2(A)–(C) and S-3A†]. While the pores of PAM beads were blocked and reduced in sizes from 295 to 90 μ m [Fig. S-2(D)–(F) and S-3B†] by its maximum loading with TiO₂, which in turn led to the reduction in surface area (1.60 m²) $\rm g^{-1})$ of the resultant PAM-TiO $_2$ NC beads. However, the removal of PAM after calcination produced hierarchically porous NS TiO₂ macrobeads [Fig. 4(A)–(F)] having large number of relatively smaller macropores of around 1 to 16 μ m sizes [Fig. 4(A)– (D) and S-3C†]. Subsequently, a 41.16 m^2 g⁻¹ surface area of porous NS $TiO₂$ beads was comparatively higher than those of the PAM and PAM-TiO₂ NC beads, which may be attributed to the highly crystalline (anatase) nature and nanoscale features of TiO₂ building units/blocks (90.16 \pm 24.84 nm) [Fig. 4(E) and (F) and S-3D†]. **EXAMENCE ARTICLE ARTICLE ARTICLE CONSULTS ARTICLE CONSULTS ARTICLE IS CONSULTS ARTICLE CONSULTS ARTICLE CONSULTS ARTICLE CONSULTS ARTICLE CONSULTS ARTICLE CONSULTS ARE CONSULTS ARE CONSULTS ARE CONSULTS ARE CONSULTS ARE**

Degradation of MB

Prior to illumination, all MB solutions each containing PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads were first kept in the dark to rule out the effect of adsorption wherein a saturated

Fig. 4 SEM images (cross-sectional view) of porous NS TiO₂ macrobead. (A–D) Hierarchically organized macroporous structure of TiO₂ bead. (E and F) Aggregation of $TiO₂$ NBUs.

adsorption was achieved after 1 h in almost all samples. It is pertinent to mention that the PAM beads showed an extremely low dark adsorption of 0.37% [see Fig. 5(A) and (B)] that is obvious due to their no net surface charge and thus can just be associated with the entrapment of MB molecules into its pores. On the contrary, the relatively higher dark adsorption of MB by both PAM-TiO₂ NC (0.98%) and porous NS TiO₂ macrobeads (6.13%) [see Fig. 5(A) and (B)] may be best explained by the electrostatic interaction between the positively charged MB and the negatively charged $TiO₂$ surface,⁵⁵ as verified by the zeta potential value $(-30 \pm 5 \text{ mV})$ at pH 10.

Upon illumination, MB in PAM samples was reduced up to 0.86 and 0.98% under simulated solar and UV light conditions, respectively [see Fig. 5(A) and (B)]. This slight destruction of MB molecules under illumination may not be logically linked to the photo-inactive PAM, but is most likely expected from its direct photolysis by both simulated solar and UV lights.⁵⁶ The simulated sunlight-driven degradation of MB reached to 16.20 and 32.88% by PAM-TiO₂ NC and porous NS TiO₂ macrobeads, respectively. However, PAM-TiO₂ NC and porous NS TiO₂ macrobeads were found much efficient to degrade MB up to 29.08 and 86.87% under UV light conditions, respectively [see Fig. 5(A) and (B)]. The improved catalytic activities of PAM-TiO₂ NC and porous NS $TiO₂$ macrobeads samples under light conditions may be ascribed to the prevalence of the hydroxyl radicals which are considered to be the major species for the $TiO₂$ catalysed photodegradation of organics.⁵⁷

Furthermore, the addition of H_2O_2 significantly enhanced the degradation of MB by all of PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads under dark and light conditions. In the presence of H₂O₂, the dark adsorption capacities of PAM, PAM- $TiO₂$ NC and porous NS TiO₂ macrobeads were increased up to 0.74, 8.71 and 9.82%, respectively [see Fig. $5(C)$ and (D)], which can most probably be assumed by the bleaching or oxidative adsorption process.⁵⁸ The oxidant also raised the photocatalytic performance of PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads to remove MB up to 3.19, 22.45 and 42.09% under

Fig. 5 Photodegradation efficiencies of (a) PAM, (b) PAM-TiO₂ NC, and (c) porous NS TiO₂ macrobeads, (A and B) without H_2O_2 , and (C and D) with H_2O_2 .

simulated sunlight, respectively, and 3.44, 34.44 and 98.53% under UV light, respectively [see Fig. $5(C)$ and (D)]. The remarkably higher photocatalytic performance particularly by porous NS TiO₂ macrobeads with H_2O_2 under light conditions can be attributed to the effective promotion of \cdot OH generation.

The photocatalytic performance of PAM, PAM-TiO₂ NC and porous NS $TiO₂$ macrobeads was also evaluated in terms of a pseudo-first-order reaction rate constant (k) as the basic kinetics parameter (see Fig. 6). After subtracting MB removal in the dark, the reaction rate constant values of PAM, PAM-TiO₂ NC and porous $TiO₂$ beads came out to be 0.00016, 0.00248 and 0.00667 min⁻¹ under simulated sunlight, respectively, and 0.00018, 0.000624 and 0.03015 min^{-1} under UV light, respectively [see Fig. 6(A) and (B)]. However, the addition of H_2O_2 significantly increased the reaction rate constant values of PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads up to 0.00055, 0.00482 and 0.00882 min^{-1} under simulated sunlight, respectively, and 0.00060, 0.00820 and 0.05041 min^{-1} under UV light, respectively [see Fig. 6(C) and (D)].

By comparison, the overall performance of porous $NS TiO₂$ macrobeads to remove 86.87 and 98.53% of MB with k values of 0.03015 and 0.05041 min^{-1} under UV light in the absence and presence of H_2O_2 , respectively, is even superior to many recently reported state-of-the-art materials (see Table S-1†). Additionally, the dark adsorption subtracted MB degradation efficiencies of porous NS TiO₂ macrobeads up to 60.89 and 64.24 mg g^{-1} under the UV light condition in the absence and presence of H_2O_2 , respectively (see Fig. S-4†), are also higher than the one and only reported material (see Table S-1†) ²⁸ because of their comparatively greater affinity for water and MB, higher probability to directly contact MB, hierarchical porosity, higher surface area, narrower bandgap (see Fig. S-1†), and orderly crystalline anatase phase (known to be more photoactive). All these factors jointly facilitate an effective production of hydroxyl radicals to elevate the removal of MB.⁵⁹ Owing to their pure $TiO₂$ -based chemical nature, these nanomaterials are more beneficial than

Fig. 6 Plots of $ln(C_0/C)$ versus illumination time representing the fitting results to deduce the pseudo-first-order reaction rate constants of (a) PAM, (b) PAM-TiO₂ NC and (c) porous NS TiO₂ macrobeads, (A and B) without H_2O_2 , and (C and D) with H_2O_2 .

the other previously reported $TiO₂$ -based composites while preventing the loss of accessible active sites, unwanted catalysts release, and so forth. Furthermore, the macrosize of these nanostructured $TiO₂$ macrobeads offers easier handling and subsequent separation with no or low energy than the conventional $TiO₂$ nanoparticles and microstructures that makes the remediation process more cost-effective and practically useful.

Disinfection of E. coli and S. aureus

Bactericidal properties of PAM, PAM-TiO₂ NC and porous NS $TiO₂$ macrobeads were evaluated in the dark and under UV illumination. The PAM beads showed no activity against both E . coli and S. aureus in the dark as well as under UV-light irradiation [Fig. 7(B) and (C) and S-5†]. The 0.2 mg mL⁻¹ of PAM-TiO₂ NC and porous $TiO₂$ beads killed 10 and 13% S. aureus in the dark, respectively (Fig. S-5B†), while 35 and 66% S. aureus after exposure to UV light, respectively (Fig. S-5A†). However, the same dosage of PAM-TiO₂ NC and porous NS TiO₂ macrobeads showed 12 and 15% efficiency to kill E. coli in the dark, respectively (Fig. S-5D†), and 46 and 72% under UV light, respectively (Fig. S -5C \dagger). After increasing doses up to 0.8 mg $\mathrm{mL}^{-1},$ PAM-TiO $_2$ NC and porous NS TiO $_2$ macrobeads caused 77 and 100% reduction, respectively, in CFUs of S. aureus under UV light after 60 min [Fig. 7(A) and (B)]. Whereas the 0.8 mg mL^{-1} of PAM-TiO₂ NC beads caused death of 80% E. coli after 60 min and the bactericidal efficiency of porous NS $TiO₂$ macrobeads reached to 100% even after 50 min of exposure to UV light [Fig. 7(A) and (C)]. Paper

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In general, PAM, PAM-TiO₂ NC and porous NS TiO₂ macrobeads were found to be inactive, fairly-active and potentiallyactive against both bacteria under all the tested conditions. The higher bactericidal efficiency of porous NS $TiO₂$

Fig. 7 (A) Digital photographs of bacterial colonies in the Petri dishes. Time-wise killing of (B) S. aureus and (C) E. coli by (b) PAM, (c) 0.2, (e) 0.4 and (g) 0.8 mg mL⁻¹ of PAM-TiO₂ NC, and (d) 0.2, (f) 0.4, and (h) 0.8 mg mL $^{-1}$ of porous NS TiO₂ macrobeads and their comparison with (a) the control containing only bacteria.

macrobeads is attributed to its crystalline nature, greater porosity and higher surface area as compared to $PAM-TiO₂ NC$ beads. Moreover, the activity of porous $TiO₂$ beads to kill E. coli was relatively better than that for S. aureus because the E. coli being Gram-negative bacteria have thin $(\sim 2$ to 3 nm) and delicate peptidoglycan layer situated between the inner (cytoplasmic) and outer cell membranes. Whereas the Gram-positive S. aureus bacteria lack the outer membrane and possess thicker cell wall $(\sim]30 \text{ nm}$ composed of peptidoglycan and lipoteichoic acid multilayers. The denser cell wall renders the killing of S. aureus more difficult.⁶⁰

The killing of bacteria in the darkness can be assumed by their adherence/adsorption onto the surface and entrapment in the interior of the porous PAM-TiO₂ NC and porous NS TiO₂ macrobeads. The direct contact may, most probably, raise an antagonism between the van der Waals forces and electrostatic interaction thereby causing physical stress, deformation and disruption of the cell wall resulting in the membrane permeability ultimately leading to bacterial death. However, disinfection of both bacteria under UV illumination can positively be correlated with the production of highly reactive oxygen species.⁶¹⁻⁶⁴

Proposed mechanism for photocatalysis of dyes/microbes

A thoroughly convincing mechanism for photocatalysis of the organic pollutants and the microorganisms on and/or near the surface of $TiO₂$ is still illusive and needs to be fully explored and elaborated. Nonetheless, many studies demonstrate that the light with energy equal to and/or higher than the bandgap of $TiO₂$ excites electrons from its valence band $(O 2p)$ to the conduction band (Ti 3d). Under UV illumination, the holes (in the valence band of $TiO₂$) react with active hydroxyl ions and/or water molecules to directly produce \cdot OH radicals, while the electrons (in the conduction band of $TiO₂$) reduce dissolved oxygen to \cdot O₂ $^-$ radicals which in turn are converted to \cdot HO₂ radicals. The \cdot HO₂ radicals are further converted to H₂O₂ which finally generate \cdot OH radicals.^{65,66} All of these short-lived and highly reactive chemical intermediates including superoxide $({}^{\cdot}O_2{}^{-})$, hydroperoxyl $({}^{\cdot}HO_2)$, and predominantly the hydroxyl (OH) free radicals contain unpaired electrons which may be

Fig. 8 A proposed mechanism for photodegradation of organic dyes and microbes by porous $NS TiO₂$ macrobead.

able to degrade the organic molecules (MB) into the final minerals and attack the bacterial cell membrane to oxidize its polyunsaturated phospholipids constituents. The proliferation of this peroxidation reaction may not only harm the shape and structure of the cell membrane but also damage the respiratory activity of bacteria and inevitably lead them to their death (see Fig. 8).⁶⁷⁻⁶⁹ The perceived \cdot OH generation by porous TiO₂ sample was further probed by a relatively higher fluorescence intensity shown by $TiO₂$ containing TA suspension as compared to only TA solution (see Fig. S-6†), because of the production of surplus \cdot OH to chemically react with TA to yield more amount of fluorescent hydroxyterephthalic acid (TAOH) which in turn exhibited a stronger fluorescence response.

Conclusions

Herein, the emulsion-templated macroporous polyacrylamide beads were prepared by oil-in-water-in-oil (O/W/O) sedimentation polymerization to be used as the exotemplates. According to the nanocasting technique, PAM beads were soaked in TiO₂ sol NPs, which aggregated to build a compact macrosized network of TiO₂. The process of the calcination not only removed the photocatalytically inactive PAM template but also converted the remaining amorphous $TiO₂$ into its more photoactive anatase phase and hierarchically porous bead architecture with a higher surface area. Consequently, the porous NS anatase $TiO₂$ macrobeads proved their higher potential to degrade the MB and to inactivate E. coli and S. aureus more effectively under UV light irradiation than PAM- $TiO₂$ NC and PAM beads. These porous NS $TiO₂$ macrobeads enjoy the benefits of easier handling and separation for recycling without requiring any external magnetic field and cumbersome separation processes – rendering them to be more efficient, cheaper, and safer for real life photodegradation applications. Furthermore, the current strategy demonstrates a step forward to develop novel porous materials with the improved photocatalytic efficiency. This approach opens up an amenable platform to design a wide range of new visible and NIR-active semiconductors and their noble-metal-based composites for the bandgap engineering to make the photodegradation process more efficient. **BSC Arbaness**

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Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

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