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Grain boundary engineering in electrospun ZnO nanostructures as promising photocatalysts
Grain boundary engineering in electrospun ZnO nanostructures as promising photocatalysts†

Anitha Senthamizhan,*a Brabu Balusamy,*a Zeynep Aytacab and Tamer Uyar*ab

Electrospun ZnO nanofibers (ZNF) have received increased attention as photocatalysts owing to their potential for incredible performance. However, uncertainty still exists in determining the correlation between grain boundaries (GBs) and photocatalytic activity. Therefore, effective thought has been put into engineering the GBs to convert ZNF into a promising photocatalyst. Herein, the obtained electrospun ZnO structures are composed of nanograins, which are connected to each other in an ordered manner. In-depth studies have revealed that the growth of nanograins severely altered the morphology of ZNF and GB areas at higher annealing temperatures ranging from 500 °C to 1000 °C. Based on the morphological features and their structural evolution, the obtained structures are named as ZnO nanofibers-1 (ZNF-1, 500 °C), ZnO hollow tubes (ZHT, 600 °C), ZnO nanofibers-2 (ZNF-2, 700 °C), ZnO bamboo structured fibers (ZBF, 800 °C), ZnO segmented fibers (ZSF, 900 °C) and ZnO nanoparticles (ZNP, 1000 °C). A strong correlation between the inherent emission features of ZNF and their peak positions have been detected with the GB. The comparative degradation efficiency of methylene blue (MB) has been studied and the results showed that the ZNF-1 with highly stacked nanograins containing rich grain boundaries demonstrated ~6 times higher efficiency than other structures. In addition, it has been shown to have a strong effect towards the degradation of Rhodamine B (Rh B) and 4-nitro-phenol (4-NP). A critical parameter for improving the photocatalytic activity is found to be the GB mediated defects, which are proposed to be oxygen/zinc vacancies at nanograin fusion interfaces, while supposedly maintaining its fibrous structure, wherein no relationship has been drawn implying the direct domination of morphology, surface area and defect.

Introduction

Incredible regard has been drawn towards electrospun zinc oxide nanofibers (ZNF) owing to their functional diversity, which has been proven to be an efficient photocatalyst for the effective degradation of organic pollutants in water.1–5 This has been attributed not only to their low cost, but also it is listed as a generally regarded as a safe (GRAS) material for human beings by the Food and Drug Administration (21 CFR 182.8991). Existing reports have already shown that ZNF are polycrystalline, composed of numerous single crystalline grains.6,7 For example, Ren et al. prepared large-pore mesoporous ZnO nanofibers using a foaming-assisted electrospinning approach and demonstrated their enhanced photocatalytic performance compared to conventional solid nanofibers.4 Sufficient evidence makes it notable here that the grain size properties and their consequent orientation are proven to have a strong impact on the functional properties of nanofibers.8,9 In the past few years, intensive research to evaluate the growth of grains and their functionalities has been observed. Surface controlled reactions are expected to play a vital role in sensors and catalyst applications that largely accept electrospun nanofibers.10–15 Important factors, including grain boundaries (GB) and their domain characteristics, have still been unable to comprehend their value and importance in photocatalytic activity (PCA).16,17 Therefore, understanding the GB mediated ZNF characteristics will pave us a path to attain a higher performance of PCA.

It is important to recall at this point that the ZNF are generally obtained by calcination of the composite fibers in which the temperature has a strong impact on determining the structure, which is impelling to comprehend the growth processes of the nanograins to attain a perfect stand on the functional performance.18 In depth reports have been provided on the growth of nanograins at higher calcination
temperatures and longer calcination times. In another study, Liu et al. showed that the electrospun La$_2$Mo$_2$O$_9$ nanowires with a bamboo-wire microstructure exhibited giant ionic conductivity of 0.20 Scm$^{-1}$. A recent study was concluded on the amount of grain boundaries in $\alpha$-Fe$_2$O$_3$ structures prepared via electrospinning for gas sensing properties of ethanol. Investigation by Katoch et al. determined the crystallinity effect of ZNF treated at different temperatures and suggested the concurrent optimization of size and crystallinity of grains as vital parameters to maximize the sensing abilities of oxide nanofibers.

Summarizing all the abovementioned factors, electrospun based nanostructures prepared at different temperatures displayed improved performances in various applications with respect to their morphology, size, crystallinity and other parameters. This implies that the structure type is crucial dependent upon the application and not a definite structure that improves the overall performance. Among the most significant factors affecting the photocatalytic performance, morphology and surface area are considered to be the primary factors. However, factors affecting the stacking behaviours of adjacent nanograins and GB have been left unnoticed. In most of the cases, the existence of defects at the GBs and their correlation is not well known. For defects, such as interstitials and vacancies, the GBs act as efficient sinks. Therefore, the necessity to study the interplay between defects and GBs in-depth, and further how these correlate with the functionality of the system. This plays a crucial role in determining the controllable parameters of luminescent features and further enhancing the degradation of dye pollutant in water.

The main goal of this study is to highlight the temperature induced structural evolution of ZNF and how the growth of grains and GBs affect their performance towards photo-degradation of dyes. From these results, one might be able to clearly perceive and develop ZNF with controlled properties through GB engineering. An important consideration to be made here is the non-correlation effects from surface area and defect related emissions on the photocatalytic performance. In contrary, efforts have been made to apprehend the effect of GBs, their arrangement and their correlation in photoluminescence properties. Due to the grain surfaces being defect rich, we strongly believe that engineering the grain surface provides a scope to improve PCA performance. The present research dictates the analysis of the structural evolution of ZnO nanofibers under temperatures ranging from 500 $^\circ$C to 1000 $^\circ$C, prepared by an efficient and cost effective method called, electrospinning. Apart from the attained morphologies of electrospun ZnO, the luminescence of ZnO is also severely affected by GBs. Extensive studies have been carried out to understand the effect of GB on the degradation efficiency of dye pollutants, in addition to its features, including morphology, high surface area and defects. Surprisingly, superior performance is shown by firmly stacked nanofibers for the efficient degradation of methylene blue (MB) compared to nanofibers and hollow tubes.

### Experimental section

#### Materials
Polyvinyl alcohol (PVA, Mw: 125 000, Scientific Polymer), zinc acetate dihydrate (Sigma-Aldrich) and methylene blue (MB, Sigma-Aldrich), Rhodamine B (Rh B, Sigma-Aldrich), and 4-nitro-phenol (4-NP, Alfa Aesar) were procured.

#### Electrospinning

The electrospinning solution was prepared by dissolving polyvinyl alcohol (PVA, 7.5% w/v) in water at 80 $^\circ$C followed by addition of zinc acetate dihydrate (4%, w/v). The resulting solution was stirred for 2 hours to obtain a homogeneous solution. Furthermore, the solution was loaded in a 3 mL plastic syringe with needle diameter of 0.8 mm and placed horizontally on a syringe pump (KD Scientific, KDS101). The flow rate of the polymer solution was controlled by the syringe pump and fixed at 1 mL h$^{-1}$. The grounded metal collector was covered with aluminium foil and placed at a distance of 17 cm from the needle tip. The electric field (15.5 kV) was applied from a high voltage power supply (Spellman, SL series, USA). Electrospinning procedure was carried out at 22 $^\circ$C and 19% relative humidity in a Plexiglas box. The obtained zinc acetate/PVA composite nanofibers were further calcined at 500 $^\circ$C, 600 $^\circ$C, 700 $^\circ$C, 800 $^\circ$C, 900 $^\circ$C and 1000 $^\circ$C separately for 3 h in air.

#### Characterization

The morphological and elemental composition analysis were performed using field emission scanning electron microscopy (FESEM, Quanta 200 FEG) equipped with an energy-dispersive X-ray spectrometer (EDS). The crystal structures of the annealed ZnO samples were measured by a PANalytical X'Pert multipurpose X-ray diffractometer with Cu Ka radiation at 2$\theta$ angle ranging from 20° to 65°. X-ray photoelectron spectroscopy (XPS, Thermo K-alpha-monochromated) was employed to analyze the surface chemical composition. Photoluminescence (PL) measurements were performed by using a time-resolved fluorescence spectrophotometer (FL-1057 TCSPC). The absorbance spectra of organic pollutants were obtained using a UV-vis Spectrophotometer (Varian Cary 100). The Brunauer, Emmett and Teller (BET) surface area analyzer (Micromeritics TriStar 3000) was used to measure the surface area of ZNF-1, ZNF-2 and ZNP. The data collected at 77 K were in the range of 0.1–0.9 relative pressure.

#### Photocatalytic performance

The catalytic performance of electrospun ZnO structures was evaluated by measuring the degradation of methylene blue (MB) dye under UV irradiation (Ultra-Vitalux Ultraviolet high-pressure lamp (300 W, Osram, sunlight simulation)) at room temperature. Typically, 1 mg mL$^{-1}$ of electrospun ZnO structures was placed into quartz cuvettes containing MB dye solution (5 ppm) separately, and then placed in the dark for 30 minutes to establish absorption/desorption equilibrium of
dye on the sample surface prior to irradiation. The samples were placed at a working distance of 15 cm from the lamp. A series of samples and controls (without catalyst) were simultaneously irradiated. The degradation of MB dye was monitored by measuring their absorbance as a function of irradiation time at predetermined time intervals using a UV-vis spectrophotometer. The reusability of the sample was performed during five consecutive cycles. The degradation efficiency of MB dye was calculated by \( \frac{(C_0 - C)}{C_0} \times 100 \), where \( C \) and \( C_0 \) indicates the absorption peak intensities before and after UV irradiation. The same procedure has been followed for Rh B (5 ppm) and 4-NP (5 ppm).

**Results and discussion**

Fig. 1 represents the schematic of the procedure for preparing various electrospun ZnO structures through a combination of electrospinning and heat treatment approaches. The SEM image of as-electrospun PVA/zinc acetate composite nanofibers clearly show the bead free and smooth fibrous nature with an average fiber diameter of 355 ± 85 nm as illustrated in Fig. S1.† Furthermore, the as-prepared composite fibers are subjected to thermal treatment (calcination) for obtaining ZNF from the total degradation of the polymer matrix. The high priority has been given to the effect of temperature, heating rate, time and environment during this period, in attaining the optimal structure and desired properties of ZnO. Our present study investigates the structural variation of electrospun ZnO nanostructures that are exposed to calcination at different temperatures ranging from 500 °C to 1000 °C for a period of 3 h. Clear indications can be sought out from the representative SEM images exhibiting the entire feature and its associated morphological variations, as illustrated in Fig. 2. Post process of calcinations, the fibrous surface is shown to turn rough, as compared to as-electrospun PVA/zinc acetate composite nanofibers. In short, it is

![Fig. 1 Schematic of the procedure for preparing various electrospun ZnO structures through the combination of electrospinning and heat treatment approaches. The obtained structures are named as ZnO nanofiber-1 (ZNF-1), ZnO hollow tube (ZHT), ZnO nanofiber-2 (ZNF-2), ZnO bamboo structured fiber (ZBF), ZnO segmented fiber (ZSF) and ZnO nanoparticles (ZNP).](image-url)
apparent that the composition of the nanofibers is mainly made up of ZnO nanograins, which further deters the fibrous morphology on increasing the temperature.

At 500 °C, the nanofiber (diameter of 180 ± 50 nm) reveals the packing of dense small grains of size 35 ± 10 nm, as shown in Fig. 2a-a1. It is found that the grains are bound to each other throughout the fiber and exhibit solid morphology. Fig. 2b-b1 demonstrates the sample obtained at 600 °C, which is found to be cylindrically connected and displaying a hollow tube morphology. Fig. 3 evidently proves the existence of the solid and hollow nature of electrospun ZnO nanofibers calcined at 500 °C and 600 °C, respectively. The marked portion in the images clearly illustrates the solid and hollow nature of the obtained electrospun ZnO nanofibers. The cross

![Fig. 2 Morphology evaluation of electrospun ZnO nanostructures at various calcination temperatures. (a-f and a1-f1) representative SEM images of electrospun ZnO nanostructures. (a and a1) 500 °C, the attained morphology is denoted as ZnO nanofiber-1 [ZNF-1]; (b and b1) 600 °C, ZnO hollow tube [ZHT]; (c and c1) 700 °C, ZnO nanofiber-2 [ZNF-2]; (d and d1) 800 °C, ZnO bamboo structured fiber [ZBF]; (e and e1) 900 °C, ZnO segmented fiber [ZSF]; (f and f1) 1000 °C, ZnO nanoparticles [ZNP] and their corresponding EDX mapping [Zn (a2-f2) and O (a3-f3)] and, representative TEM images (a4-f4) and HR-TEM images (a5-f5).](image)

![Fig. 3 Comparative SEM images of (a) ZNF-1 and (b) ZHT. The marked portion in the images clearly illustrates the solid and hollow nature of obtained electrospun ZnO nanofibers.](image)
sectional view represents that the nanograins are tightly packed in ZNF-1 along the fibers, whereas ZHT possesses distinctly packed nanograins, which creates a hollow tube morphology as observed.

A detailed analysis on the image of the single hollow tube implies an increase in the grain size (60 ± 10 nm), with well-defined gaps between the nanograins and as a result the diameter is increased to 200 ± 50 nm. In addition, an increase in the temperature to 700 °C causes shrinkage in the fiber diameter, measuring 130 ± 30 nm, as illustrated in Fig. 2c-e1. Moreover, the composition of the fibers comprising larger nanograins (120 ± 40 nm) is due to significant growth and coalescence of the grains. A worthy mention has to be made here that the annealing temperature has a severe effect on the distribution of grains in a consistent manner. Evidence proves that the fibrous morphology of the grown nanograins are bounded very tightly to each other and make up a solid fiber appearance.

Upon calcination at 800 °C, the fibers exist as a bamboo-like network structure by arranging themselves as single chain nanograins of size 180 ± 80 nm, as depicted in Fig. 2d-d1. It is found that the grains are well grown at this temperature and a clear change in the morphology is observed. Severe breakage of the fibers is noticed, with every short segment composed of tightly packed particles upon calcination at 900 °C, each with a grain size of 300 ± 110 nm, as illustrated in Fig. 2e-e1. The agglomeration of ZnO nanograins with complete destruction of the fibrous structures were noted following a thermal treatment at 1000 °C as clearly demonstrated in Fig. 2f-f1. Based on the morphological features and their structural evolution upon calcination (500 °C to 1000 °C), the obtained structures are named as ZnO nanofibers-1 (ZNF-1, 500 °C), ZnO hollow tubes (ZHT, 600 °C), ZnO nanofibers-2 (ZNF-2, 700 °C), ZnO bamboo structured fibers (ZBF, 800 °C), ZnO segmented fibers (ZSF, 900 °C) and ZnO nanoparticles (ZNP, 1000 °C). The measured mean grain diameters are plotted against the annealing temperature shown in Fig. S2.t

Further elemental mapping has been performed to evaluate the presence of zinc (Zn) and oxygen (O). The observed results clearly show the homogeneous distribution of both Zn and O throughout structures, as shown in Fig. 2a2–f2 (Zn) and a3–f3 (O). The observed structural variation were further investigated using TEM, which is presented in Fig. 2a2–f4, highlighting the coincidence of change in morphology of the fibers with SEM results. The HR-TEM Fig. 2a2–f5 represents the lattice spacing of ZnO nanograins and the corresponding results suggest that they are polycrystalline. The lattice spacing of two consecutive planes are measured to be 0.26 nm, 0.58 nm, 0.52 nm, 0.24 nm, 0.28 nm and 0.58 nm for ZNF-1, ZHT, ZNF-2, ZBF, ZSF and ZNP, respectively (Fig. 2a2–f5). The BET specific surface area of ZNF-1, ZNF-2 and ZNP were measured to be 6.44 m² g⁻¹, 5.07 m² g⁻¹ and 1.61 m² g⁻¹, respectively. The results have shown that the surface area of electrospun ZnO structures are decreased upon increasing the calcination temperatures. The higher surface area observed in ZNF-1 compared to ZNF-2 might be due to the smaller particle size and voids amongst the nanograins. A clear indication is shown by the HR-TEM image of ZNF-1 about the existence of well grown nanograins with an intense population that are found to be robustly overlapped along the fibers, as shown in Fig. 4a. Contrary to this, the size of the ZHT grains that are found to be smaller in size, less populated and incoherently arranged. Importantly, it is found that the grains exhibit strong crystallinity, as they most likely exist singularly, with considerable distance between each other. Moreover, it is noticed that there is no definite GB due to their arrangement characteristics. Therefore, this is found to have an impact on the grain connectivity of ZNF-1 from the other two structures, as they have a more compact structure. ZBF grains are pragmatically found in random shapes and sizes, tightly bonded to each other without secondary particle arrangement. The appropriate formation of GB in ZNF-2 is clearly shown in Fig. 4b. These obvious variations in the GBs infer the difference in the luminance characteristics as well.

XRD analysis gives us inputs on the formation of ZnO and their crystalline nature. As shown in Fig. 5, the peaks in the diffraction patterns are indexed to (100), (002), (101), (102), (110) and (103) planes of ZnO highlighting their Wurtzite hexagonal crystal structure (JCPDS No. 36-1451). Conspicuously, no evident peaks of polyvinyl alcohol (PVA) or any other impurities have been observed emphasizing the complete removal of the polymer and the simultaneous decomposition and formation of zinc acetate into pure ZnO.36 On increasing the temperature, we noticed a higher intensity and sharpness in the diffraction peaks suggesting an increase in the grain size and crystal quality improvement.

Detailed information on surface composition and the oxidation states of electrospun ZnO nanostructures calcined at different temperatures have been obtained from XPS analysis. As illustrated in the Fig. 6, the high resolution scan of O 1s profile can be composed of multiple peaks and fitted as three peaks with the Gaussian function, denoting the existence of three different O species named lattice oxygen (O_l), oxygen vacancies (O_v) and chemisorbed oxygen species (O ads). The peak at 530.1 ± 0.2 eV in all samples is attributed to O^{2−} ions in a Wurtzite structure composed of a hexagonal Zn^{2+} ion.
array, which are surrounded by zinc atoms with the full supplement of nearest-neighbour O$_{2}^{−}$ ions. They are generally known as lattice oxygen (OL) and their percentage represents the relative amount of the O at the ideal lattice positions. The centred peak located at 531 ± 0.1 eV is attributed to the O$_{2}^{−}$ ions in the oxygen deficient regions within the matrix of ZnO. The third peak around 532.1 ± 0.2 eV is typically ascribed to OH$^{−}$ groups, chemisorbed oxygen or dissociated oxygen on the surface of the ZnO.37,38 The observed variations in the intensity of the peaks might be connected to the concentration of oxygen vacancies. The calculated relative percentage of OL, O$_{Ch}$ and OV for all samples are noted in their corresponding Figures. The highest intensity of OL as compared with O$_{Ch}$ and OV indicates the strong Zn–O bonding in the prepared samples. The gradual increases in OV is noted for all samples except ZBF, which might be due to the reduced amount of GB. It is realized that the peak O$_{Ch}$ that is related to the chemisorbed oxygen, cannot be removed even at higher temperatures. The estimated relative percentage of oxygen vacancies (OV) following the trend as ZNP > ZSF > ZNF-2 > ZHT > ZNF-1 > ZBF. The intese characteristic peaks observed at 1021.5 ± 0.3 eV and 1044.6 ± 0.2 eV in XPS spectra of Zn 2p are assigned to Zn 2p$_{3/2}$ and Zn 2p$_{1/2}$, respectively (Fig. S3†). The spin–orbit splitting of $\sim$23 eV between two core level components of Zn 2p$_{3/2}$ and Zn 2p$_{1/2}$ are in good agreement with the reported literature values.39 The GBs are considered to be an obvious sink for defects and facilitates more chemisorption for oxygen-related species, as discussed in the previous section. This is the reason why we have observed a higher percentage of O$_{Ch}$ in ZNF-2 as compared with ZNF-1 and ZHT. This would be more advantageous in enhancing dye degradation processes because the presence of hydroxyl groups can effectively facilitate the trapping of photo-induced electrons and holes.37 If the performances are strongly correlated with oxygen vacancies, then there would be a higher degradation rate in defect rich samples. We will observe in future sections, whether these results follow the same trend for enhancing the PCA or not.

Fig. 7 showcases the photoluminescence spectra of ZnO nanostructures at different temperatures. Deconvolution of the spectra shows a strong influence of photoluminescence for ZnO by their grain arrangements and the presence of GBs. The presence of defects has confirmed the broad range of defect emissions. Evidence has already proved the correlation of defects and their position in ZnO, including oxygen vacancies (V$_{O}$), zinc vacancies (V$_{Zn}$), zinc interstitials (I$_{Zn}$) to the preparation methods, annealing temperature, and other conditions.38–40 However, information on the exact defect centres and the electron transition pathways are still unknown. The dominant peak in the UV range called near band edge emission (NBE) is related to the direct recombination of photo generated carriers, denoted as P$_{1}$. Illustration proves the occurrence of a slight shift in the NBE spectra of all other samples to longer wavelength as compared with the ZNF-1 due to the increase in the grain size. Apart from this, an obvious change in the peak intensity can be noticed among the spectra. Numerous studies have illustrated that the controlled density of the defects and their position is important for efficient photocatalysis.37,38,41–45

From the conducted experiment, we have clearly observed that Zn and V$_{O}$ showcase dominant defects in the prepared electrospun ZnO nanofibers, apparently visible at the GB regions. On performing a comparative analysis with ZNF-1, it is observed that the intensity of defect level emissions (DLE) gradually decreases emphasizing the well separated and crystallized nature of the formed grains, thereby significantly reducing their defect related emission in ZHT. In addition, it is also noticed that the intensity of DLE in ZBF is significantly higher than that of either ZNF-1 or ZNF-2. Thus, this analysis implies that with an increase in the density of the GB, the defect concentration increases. Theory also suggests that for the stability of defects at the GB regions.46 Therefore, the successful combination of both structural changes and
luminescent properties leads to developing interesting facts on the GB engineering in electrospun ZnO nanostructures. The violet (P2) and blue (P3) emissions are attributed to the transition from IZn (IZn $\rightarrow$ VB) and extended IZn states (ex-IZn $\rightarrow$ VZn) to the valance band, respectively.  

Another dominant peak existing in all the samples is related to the green emission comprising two components (V$^*$O $\rightarrow$ VB [P4], and CB $\rightarrow$ VO$^{++}$ [P5]). All these cases show competency in the peak intensity of V$^*$O $\rightarrow$ VB with ex-IZn $\rightarrow$ VZn. However, the peak intensity of CB $\rightarrow$ VO$^{++}$ is found to be lower than V$^*$O $\rightarrow$ VB in all cases. Apparently, on increasing the temperature, the intensity of violet and blue emissions are increased, whereas in the case of ZNP, the emission from ex-IZn $\rightarrow$ VZn is found to be superior than the emission from V$^*$O $\rightarrow$ VB. Importantly, the peak positioned at $\sim$465 nm corresponds to the blue emission, which is significantly red-shifted to 480 nm on increasing the temperature. Clear evidence is provided from the morphology evaluation of ZnO nanostructures that the alteration of GBs leads to the intensity variation of peak and their position. ZNF-1 and ZHT shows superior UV emission, denoting good optical quality. Observations prove the reduction of defects in well-ordered grains and their separated nature. Additionally, both XRD and HR-TEM results show that high temperature annealing enhances the crystallinity of ZnO. However, in contrary to this, the observed PL spectra does not confirm the reduction of defects with high temperature annealing. Therefore, persistent results imply that higher temperature annealing promotes grain growth and GB related emissions occur with enhanced intensity thereby highlighting the importance of GB on the luminescent characteristics of ZnO.

To date, electrospun nanofiber based catalytic systems display enhanced photocatalytic performance owing to the existence of a larger surface area and other dominant features.  

Fig. 6 XPS spectra of O 1s in electrospun derived ZnO structures annealed at various temperatures over a range from 500 °C to 1000 °C. The observed high resolution scan of O 1s spectra is composed of multiple peaks and fitted as three peaks denoting the existence of three different O species named lattice oxygen (O$_L$), oxygen vacancies (O$_V$) and chemisorbed oxygen species (O$_{ch}$). The details on the position and relative percentage of peaks are given in corresponding spectrum.
process on the photocatalytic performance rather than solid fibers. However, we would like to stress again that the core theme of our study is not in line with effects of defect related emission on photocatalytic performance, but it is elucidate the effect of GBs. The observance of the growth of nanograins results in an obvious change in the surface of the fiber, thereby influencing the GB area. It has to be noted here that the surface morphology of the nanofibers plays a crucial role in highlighting surface related attributes as an excellent feature. To summarize, GBs play an important role in determining specific physical and chemical properties of nanofibers in addition to higher surface area and aspect ratios.

Controlled morphological features facilitated by prepared ZnO nanostructures expect to showcase an enhanced performance. Therefore, this is accomplished by the photodegradation of a cationic textile dye, MB performed under UV irradiation. The diverse feasible mechanism, including the production of reactive oxygen species involved in the enhancement of photocatalytic activities, have been already well studied and reported. In the outset, the MB degradation experiment was conducted in the presence of ZNF-1 under dark conditions and the respective absorbance spectra (Fig. S4†) clearly demonstrated that there is no significant effect on the degradation even up to 90 min. This confirms that the dye was not adsorbed onto the surface of the catalyst and it could reach the adsorption equilibrium in 30 min. Photodegradation efficiencies of electrospun ZnO nanostructure samples with different morphologies as a function of irradiation time are presented in Fig. 8a. The recorded absorption spectra of MB as a function of irradiation time during the degradation processes are presented in Fig. S5†. The self-degradation efficiency of the control dye solution without any catalyst was also presented for comparison and was found to be ~16% under irradiation for 90 min. A worthy note here is the correlation of distinct particle sizes and morphology of the calcined electrospun ZnO nanostructure samples. Therefore, observations prove that the MB dye (~5 ppm) has undergone complete degradation for the ZNF-2. In contrast, such photodegradation efficiency has not been observed in the other structures. Analysis proves that calcined samples at 500 °C (ZNF-1), 600 °C (ZHT), 700 °C (ZNF-2), 800 °C (ZBF), 900 °C (ZSF) and 1000 °C (ZNP) exhibited about 85%, 76%, 98%, 74%, 59% and 49% degradation, respectively, as can be observed from Fig. 8b.

The photo-degradation of MB follows the pseudo-first-order rate law (Fig. 8c) and the rate expression is given by the equation \( \ln(C_0/C) = kt \) where \( k \) is the degradation rate constant (min\(^{-1}\)), and \( C_0 \) and \( C \) represents the initial and final concentration of MB at a given time period. The observed values of the rate constants are 0.0199, 0.0169, 0.0419, 0.0154, 0.0087 and 0.0062 min\(^{-1}\) for ZNF-1, ZHT, ZNF-2, ZBF, ZSF and ZNP, respectively. As observed in Fig. 8d, the ZNF-2 shows a much higher rate constant which is about 6.7 times higher than the ZNP, which exhibits the lowest rate constant. This implies the fact that the ZNF-2 displays higher degradation efficiency when compared to other ZnO nanostructures. Furthermore, the photocatalytic degradation efficiency of ZNF-2 for Rh B and 4-NP were studied and results are shown in Fig. S6 and S7.† The decolourisation of pollutants was evaluated through monitoring the absorption spectra, as shown Fig. S6a and S7a.† It is noted that the intensity of the peaks gradually decrease upon increasing the irradiation time. The corresponding digital images of pollutants taken at definite time interval are shown in Fig. S6b and S7b.† The degradation efficiency of ZNF-2 for Rh B and 4-Nitrophenol were calculated to be ~91% in 3 h and ~92% in 4 h, respectively. The above results clearly show that ZNF-2 has excellent photocatalytic activities for a wide range of pollutants.

Moreover, it has been clearly implied that the photodegradation performance of the samples has been purely based on the GB mediated defect emissions along with their well-maintained morphology. Furthermore, the reusability of the ZNF-2 was evaluated in five consecutive cycles and the results in Fig. S8† shows that there is a slight decrease in the degradation efficiency that could be due to the photo corrosion effect. To attain an in-depth view on the effect of photo corrosion on the catalyst, we have exposed electrospun ZnO nanostructures to UV irradiation for a time period of 24 h by changing the dye solution and studying the changes in surface morphology and their chemical composition. The observed morphological features evidently revealed that the surface of the electrospun ZnO nanostructures is severely damaged, as illustrated in Fig. S9.† Interestingly, the fibrous
structure was not significantly altered and agglomerated (ZNF-1, ZHT, ZNF-2 and ZBF). Conversely, an apparent agglomeration of ZNP has been noticed and exists as a flake-like structure, as clearly shown in Fig. S10.† In addition, high resolution XPS analysis of the samples proved that the drastic shift in O 1s peak to 533.97 ± 1.0 eV. Similarly, the intense characteristic peaks observed at 1021.5 ± 0.3 eV and 1044.6 ± 0.2 eV in Zn 2p were considerably shifted to 1023.3 ± 0.5 eV and 1046.35 ± 0.4 eV, respectively, as illustrated in Fig. S11.† The findings confirmed the photocorrosion of ZnO during the photodegradation process. Further improvements in inhibition of photocorrosion in electrospun ZnO nanostructures present a challenge and much effort is needed to provide a solution to this issue.

The overall analysis of this experiment suggests that the dependence of the enhanced PCA found in ZNF-2 on structural features, as well as defects at the GB region. Moreover, it is imperative to clearly differentiate the properties of both ZNF-1 and ZHT because both of them possess structural features with small grains. Furthermore, the structures of ZBF, ZSF and ZNP represent significantly higher defects than other structures. Thus, samples possessing excellent structural features and defects prove to display improved photocatalysis. Interestingly, it should also be noted herein that ZNF-2 showed superior photo degradation performance compared to than other structures, implying the crucial role of defects in the GB region.66–68 In the current study, we have thus highlighted the creation of defects by engineering the GB in electrospun ZnO structures for higher photocatalytic performance.

Conclusions

We have effectively demonstrated the temperature induced deformation of electrospun ZnO nanostructures from fibers to particles and their comparative degradation efficiency of MB. The grain size and GB mediated emission were concerned as primary factors and evaluated the influence of
photocatalytic efficiency of ZnO nanostructures. It has been shown that the morphology of electrospun ZnO nanofibers are severely altered at higher temperature annealing. Furthermore, it is observed that the density of nanograins are decreased with a simultaneous increase in size. The effect on GB on the emission characteristics of ZnO have been derived and observations have shown that defects are reduced in well-ordered grains and their distinct nature, observed in ZNF-1 and ZHT. Studies from the photocatalytic measurement shows that the ZNF-2 with highly stacked nanograins demonstrates superior performance for the degradation of MB as compared to ZNF-1 and ZHT, which are usually thought to conferring superior performance. Detailed discussion shows that the ZNF-2 with highly stacked nanograins and their boundary areas, in addition to superior features, including morphology, higher surface area and defects. This implies the detailed study of both the GB character distribution and the GB network arrangement for the development of an efficient catalyst.

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Notes and references


