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Fabrication of Visible Light Active Mn-Doped Bi2WO6-GO/MoS2 Heterostructure for Enhanced Photocatalytic Degradation of Methylene Blue

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Research Article

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1 Fabrication of visible light active Mn-doped Bi₂WO₆-GO/MoS₂ heterostructure for

2 enhanced photocatalytic degradation of methylene blue

3

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8 Abstract

9 The increase in environmental pollution has led to an increased investigation in the development of 10 novel ternary photocatalytic systems for remediation. These Photocatalytic systems exhibit superior photocatalytic action for the removal of pollutants because of their visible light active bandgaps. A 11 12 highly effective visible light active ternary heterojunction was fabricated using a hydrothermal method assisted by ultrasonication. Herein, we report the Insitu hydrothermal synthesis of Mn-doped Bi₂WO₆-13 GO/ MoS₂ photocatalyst, efficiently exhibiting greater photocatalytic activity for the wastewater 14 15 treatment under solar light. The binary metal sulphide (MoS_2) used as a co-catalyst, acted as electron 16 collector and Graphene oxide (GO) as a support material for interfacial electron transfer to and from 17 bismuth tungstate and MoS₂. The as-prepared samples were characterized using SEM-EDX, FT-IR, 18 XRD, XPS, and UV–Vis techniques. The bandgap of the novel photocatalyst was found in the visible 19 region (2.2 eV) which helped in suppressing photoinduced electron-hole pairs recombination. The 20 ternary Mn-BMG showed 99% methylene blue removal after 60 minutes of sunlight irradiation at the optimum conditions of pH 8, catalyst dose 50 mg/100ml and initial MB concentration of 10ppm under 21 22 sunlight irradiation. The Mn-BMG has proved to be an effective sunlight active photocatalyst that can 23 be reused without substantial loss in photocatalytic efficiency.

24

25 Keywords: Heterogenous Photocatalysis; Wastewater treatment; Semiconductor metal oxides;
26 nanocomposites; doping; graphene Oxide.

28 **1. Introduction:**

29 Water pollution and energy crisis has become a global threat due to rapid industrialization. Various 30 industries like textile, pharmaceutical, paper, cosmetics and food industry utilize organic dyes for refining the quality of their manufactured products. These industries frequently produce an extensive 31 32 load of untreated dye wastewater, making dyes a major component of water pollution (Aziz et al., 2020). 33 Methylene Blue (MB) dye is an immensely used coloring agent for coloring silk, cotton and wood 34 products. Owing to the complex aromatic structure, the dye methylene blue is resistant to oxidation and 35 degradation by conventionally available methods. Exposure and utilization of water having traces of 36 this dye may cause serious health hazards in humans and animals including nausea, difficult breathing, 37 cyanosis, jaundice, and irritation on the skin and in the eyes. Hence, an appropriate method for the 38 removal of such poisonous dyes is crucial in present times of water shortage (Nawaz et al., 2020). 39 Advanced oxidation processes have received considerable importance because of their role in 40 wastewater treatment and water disinfection. Among Advanced oxidation processes, photocatalysis is 41 considered a green technology because of the utilization of solar energy at ambient conditions (Mushtaq 42 et al., 2020). Heterogeneous Photocatalysis is now a promising strategy to address the wastewater crisis 43 owing to the generation of light-induced electrons and holes which can easily scavenge H₂O molecules 44 O₂ and produce reactive oxygen species. Hence, Using solar energy one can convert recalcitrant organic 45 pollutants into harmless compounds (Mudhoo et al., 2020). Ternary metal tungstates have been used for water purification owing to their narrow bandgaps, low-cost applicability, crystallinity and effective 46 47 utilization of sunlight because of their narrowband gaps (Singh et al., 2020). These ternary compounds 48 suffer from the limitation of high charge carrier's recombination. The photocatalytic efficiency of the 49 ternary metal oxides irradiation can be enhanced by coupling with other semiconductor materials having 50 a narrow bandgap and support materials or via doping with transition or plasmonic metals under sunlight 51 (Saher et al., 2021). Since the formation of heterojunctions via coupling and doping both can be 52 effective for suppression of $e^{-/h^{+}}$ recombination by providing new pathways to charge carriers (Mafa 53 et al., 2019). Bismuth tungsten oxide (Bi₂WO₆), is an important member of ternary Aurivillius metal 54 oxides having perovskite layered crystal structure. The sandwich structure consists of alternating layers of $(Bi_2O_2)^{2+}$ and octahedral WO₄⁻². This interlayer spacing offers a large surface area and adequate active 55 sites for an efficient photocatalytic process and also prevents electrons and holes from recombining. 56 57 Bi₂WO₆(BWO) has a narrow bandgap of (2.80 eV) which makes it efficient as a visible light active photocatalyst as compared to binary compounds (Wang et al., 2017). Although BWO exhibits excellent 58 59 photodegradation efficiency towards organic pollutants, its activity is still hindered by some factors 60 including narrow photo-response limiting maximum utilization of solar light and high recombination 61 rate of charge carriers. Among various approaches being proposed in previous researches, the construction of ternary novel heterojunctions via coupling and doping has widely been used to enhance 62 63 the charge separation efficiency (Li et al., 2018). Hence, the key step for the formation of novel ternary 64 heterojunctions is the selection of a suitable material for coupling with BWO ensuring maximum visible 65 light utilization. Among all methods to increase the visible light response of ternary compounds, doping 66 is a relatively effective method. These ternary hybrid photocatalytic systems can separate electrons and 67 holes more efficiently than binary nanocomposites. Doping alters the bandgap and atomic structure of 68 host material by the introduction of foreign ions. The introduction of additional energy levels helps in 69 trapping and separating electrons for a longer period of time hence preventing recombination. Transition 70 metal doping includes use of transition metals like Cobalt, niobium, manganese, zinc, tungsten, iron 71 and molybdenum for modifying the d-orbital configuration, the fermi level and the bandgap (Huang et al., 2016). The transition metals have partially filled d orbitals which helps in the formation of new 72 73 energy bands below the conduction band of host material. These newly formed energy bands are 74 responsible for redshift in the bandgap energy enabling photocatalysts to act efficiently in the visible 75 range of light spectra (Ahmad, 2019).

Molybdenum disulfide (MoS₂) is a typical layered two-dimensional layered binary compound particularly used as a hybrid co-catalyst material. A large number of nanocomposites coupled with MoS₂ have been reported which showed enhanced activity because of the existence of S-atoms on exposed ends of MoS₂ (Lv et al., 2017). But, the overall photocatalytic reaction activity of MoS₂ alone is reduced due to poor electrical conductivity. Besides the effectiveness of Bi₂WO₆ as a visible light active catalyst and its coupling with MoS₂, a suitable mediator is required for transferring electrons efficiently.

83 The presence of Graphene oxide provides enhanced surface area because of the two-dimensional 84 assembly of carbon atoms which are all Sp^2 hybridized and covalently linked. The electrically 85 conductive large surface area of graphene oxide helps in effective charge migration and acts as a mediator for shuttling electrons between available active sites of photocatalyst and co-catalysts, 86 87 subsequently suppressing the electrons and holes from reuniting, accelerating the overall catalytic 88 process (Tabasum et al., 2020). Also, Graphene oxide adsorbs organic pollutants well on the surface of photocatalysts besides enhancing their photo-response. Hence, the graphene supported ternary 89 90 heterojunctions greatly improve the overall activity of ternary nanocomposites under visible light 91 irradiation.

92 Herein, we report the synthesis of Mn-doped Bi₂WO₆-supported over Graphene oxide and coupled with 93 binary metal sulphide MoS_2 by adjusting the weight percentage of each component in the ternary 94 composite. A simple in situ hydrothermal synthesis process was employed for the fabrication of novel 95 Mn-BMG and was employed for the cationic dye Methylene blue (MB) degradation under sunlight. 96 Meanwhile, the characterization studies of the prepared catalysts were done by techniques like FTIR, 97 SEM-EDX, XPS, XRD and UV-Vis spectroscopy. The studies showed that Graphene oxide supported 98 Mn-Bi₂WO₆ coupled with MoS₂ provided enhanced surface area and an easy pathway for the transport 99 and separation of photogenerated charge carriers effectively increasing the visible light response of

100 photocatalyst.

101

102 2. Experimental and synthesis

103 2.1. Materials and Reagents:

Sodium tungstate dihydrate (Na₂WO₄.2H₂O), Sodium molybdate dihydrate (Na₂MoO₄·2H₂O), Bismuth
nitrate pentahydrate Bi (NO₃)₃.5H₂O, Sodium Nitrate (NaNO₃), Potassium permanganate (KMNO₄),
Ethylene glycol (analytical grade), Sulphuric acid (H₂SO₄) and Hydrogen peroxide were obtained from
(Sigma Aldrich), Manganese chloride MnCl₂.4H₂O were obtained from UNI-Chem reagents. Graphite
Powder and thioacetamide C₂H₅NS were obtained from Sharlau. Dye methylene blue was purchased
from Fischer scientific company. All chemicals used were of analytical grade and used without any
further purification. Distilled water was used for all the reaction solutions throughout the study.

111

112 **2.2.** Synthesis of Graphene Oxide:

Graphene oxide (GO) was synthesized by a modified hummers method. as previously reported in our
work (Tabasum et al., 2019). The detailed information is present in the supplementary material.

115

116 **2.3.** Synthesis of Mn-BWO-GO/MoS₂ (Mn-BMG)

Initially, Bi (NO₃)_{3.5}H₂O (5mmol) was dissolved in 30ml Ethylene glycol (Solution A) and 117 Na₂WO₄.2H₂O (2.5mmol) was dissolved in 30ml distilled water termed Solution B. The prepared 118 119 solution B was added dropwise in solution A under vigorous stirring. Mn doping (10 mol%) was done 120 by adding precursor $MnCl_{2.4}H_{2}O(0.0178gm)$ directly in the above solution and the whole mixture was 121 magnetically stirred for two hours and poured in a Teflon lined stainless steel autoclave for hydrothermal treatment for 24 hours at 180°C temperature. A white powder was obtained after 122 123 centrifugation and washing thrice with distilled water and ethanol was oven-dried at 60°C. Next, an in-124 situ hydrothermal method was performed in which 2.5mmol of Sodium molybdate and 5mmol of thioacetamide was dissolved in 30ml distilled water to form MoS₂(Solution C). The whole mixture was 125 stirred magnetically for two hours. The above prepared Mn-BWO in 50wt% was dissolved with 126 magnetic stirring in 20 ml distilled water and added dropwise to solution C. Meanwhile, Graphene oxide 127 128 20wt% (0.2gm), prepared above was ultrasonically dispersed in 20ml distilled water for about 30 minutes and added dropwise to Solution C. The whole mixture was vigorously stirred for another three 129 130 hours until a homogenous solution was obtained. The whole mixture was again given hydrothermal treatment at 180°C for 24 hours by pouring the reaction mixture in a 250ml capacity autoclave. The 131 prepared black powder was centrifuged, washed thrice using distilled water and ethanol and dried in an 132 133 electric oven at 60°C for 12 hours. The ternary BWO-GO/MoS₂ (BMG) was prepared by a similar method except for the addition of MnCl₂.4H₂O. The binary BW-GO (BG) was prepared by a method 134 135 reported earlier by Nguyen and his fellows (Nguyen et al., 2018).

136 **2.4.** Characterization and equipment.

The microstructure, morphology and elemental analysis of as prepared ternary Mn-BMG and BMG 137 were characterized using a Scanning electron microscope equipped with EDX (SEM-EDX; FEI NOVA 138 450 NANOSEM). The crystalline properties of all powder samples were measured by XRD (Bruker D8 139 140 Advanced equipment operated) over a range of 2θ values from 5-85 degrees. The presence of several functional groups on the surface of prepared photocatalysts was studied by Fourier transform infrared 141 142 spectroscopy (FTIR, Thermo Nicolet). The XPS analysis was done to study the elemental states and the 143 surface chemical compositions investigated (Escalab 250 XPS system, Thermo Fisher Scientific UK). 144 The bandgap analysis of the ternary doped and undoped samples was done using UV-VIS 145 spectrophotometer (Cecil CE 7200) in the 200-800 nm range by dispersing catalysts in water under 146 ultrasonication.

147 **2.5.** Evaluation of photocatalytic degradation

148 Various experiments on the photocatalytic degradation of cationic dye methylene blue (MB) were 149 carried out under sunlight in batch mode using ternary Mn-BMG, BMG and binary BG nanocomposites 150 for one hour. The nanocomposites in different concentrations were initially screened in Ultraviolet and 151 Visible irradiation to check the feasibility of the MB degradation. The effect of several parameters 152 including pH (2–9), catalyst dosage (10–100mg/100ml), initial dye concentration (2–20 ppm) and 153 irradiation time (10-120 min) on MB photodegradation was studied. For comparison, a single 154 experiment was performed under UV irradiation for each catalyst at their optimum conditions. The control experiments of Dye/Sunlight, Dye/UV, Dye/UV light/catalyst were also run to examine the dye 155 156 removal under these conditions also. For the photocatalytic degradation of MB under sunlight, the 157 desired quantities of catalysts were dispersed via ultrasonication in 100 mL of 10 ppm dye solution. Before exposing the solutions to sunlight, the solutions were stirred mechanically in dark to obtain 158 159 adsorption-desorption equilibrium for nearly thirty minutes. Before exposure to any light source, the adsorption capacity of samples was examined. After checking the adsorption capacity of photocatalysts, 160 the dye solutions were kept under sunlight irradiation for about 60 min at 180 rpm in electrical orbital 161 shakers. After each trial, the nanocomposites were separated from the solution by centrifugation of 5ml 162 dye solution and the concentration of the remaining dye was determined by taking absorbance of clear 163 solution using a UV–VIS spectrophotometer at 665nm. A blank beaker of dye solution was also kept in 164 165 dark and sunlight to investigate dye degradation in the presence and absence of light. The % degradation 166 was calculated using the following formula

167

% Degradation =
$$1 - \frac{A}{Aq} \times 100$$
 Eq. (1)

Here, Ao is the initial absorbance and A is absorbance after treatment under solar light. The pH was
maintained by using different molar concentration solutions of HCl and NaOH using a pH meter (Ohaus
ST3100, USA). The sunlight intensity was measured by a solar power meter (SM206), and the

- brightness was measured by a light meter (HS1010A). The reusability of catalysts was investigated by
 5 times recycling the catalysts and reusing at optimum conditions under sunlight each time with freshly
- 173 prepared 10 ppm MB solution.
- 174

175 **3. Characterization**

176 **3.1. FTIR Analysis:**

FTIR analysis was employed to elucidate the existence of several functional groups and chemical bond 177 formation in the doped and undoped ternary BMG samples as shown in figure 3.1. Transmittance peaks 178 were observed around 645, 948 1024, 1069, 1380, 1450 and 1620 cm^{-1} for the ternary nanocomposites. 179 The peaks at around 1024 cm^{-1} and 1069 cm^{-1} have appeared because of O=S=O asymmetric stretching 180 vibrations and S-O-S symmetric vibrations, respectively. Additionally, the presence of a peak at 1185 181 cm^{-1} shows asymmetric vibrations of Mo-O bonds confirming the synthesis of MoS₂ as described in 182 previous studies (Zolgharnein et al., 2018). An elongated broad band from 645 cm⁻¹ and 845 cm⁻¹ is 183 accredited to Bi–O and W–O stretching vibrations. A peak at 715.46 cm⁻¹ explains the antisymmetric 184 bridging mode which is associated with tungstate chains. An intense peak present at 1384 cm⁻¹ explains 185 the bending vibrations of W–O–W. These characteristic bands of BW were observed for all samples. 186 (Khojeh et al., 2017). The absorption band appearing at 1450 cm¹, 1557.9 cm⁻¹, and 1630.75 cm⁻¹ 187 indicates the formation of COO⁻, C=C and C=O stretching vibrations indicating successful fabrication 188 of GO and incorporation of GO into BW (Hu et al., 2019). The skeletal vibrations of graphene sheets 189 appearing at 1630.75 cm⁻¹ indicate carbonyl stretching that the oxygen-comprising groups in GO are 190 191 decomposed upon exposure to a hydrothermal environment. But no other obvious absorption peak 192 related to GO could be observed in the composites due to the low content of GO (Liu et al., 2017). These vibrations appearing at 1630cm⁻¹ may also indicate adsorbed water molecules and unoxidized 193 graphitic domains (Hou et al., 2020) A very small peak is visible at 947 cm⁻¹ in Mn-doped ternary 194 composite as compared to undoped ternary sample, which is ascribed to Mn-O vibrations as confirmed 195 196 from previous studies (Ahmad et al., 2019). Besides, the peak intensity is reduced in the doped ternary 197 composite, which supports the successful insertion of dopant ions into the host material.





Fig. 1. FTIR spectra of (a) BMG and (b) Mn-BMG ternary nanocomposites

200 **3.2 XRD analysis:**

The phase purity and crystalline structures of all the ternary and binary samples were studied by XRD 201 characterization in 2θ range of 5-85° as shown in Figure 3.2. The influence of manganese doping and 202 GO/MoS₂ coupling on the crystalline structure of bismuth tungstate were also affirmed through XRD. 203 All the strong diffraction peaks of Bi_2WO_6 are readily indexed at 20 values 28.02°, 32.8°, 47.12° and 204 205 55.7°, well assigned to (131), (200), (260) and (331) crystal planes indicating the successful formation of Bi₂WO₆ in orthorhombic phase. The sharp diffraction peaks of bismuth tungstate in all samples 206 including Mn-BMG, BMG, and BG are the same showing formation of high purity and crystalline 207 208 single-phase orthorhombic bismuth tungstate, showing that the addition of MoS_2 And GO has not 209 affected the orthorhombic structure of bismuth tungstate. The peaks at 2 θ values of 39.2° and 52.2° 210 corresponding to (100) and (110) planes, indicating the formation of amorphous molybdenum 211 disulphide. No other visible peaks of MoS₂ are observed suggesting low crystallinity and trace loading of MoS_2 on the GO support. The reported peak of Graphene oxide at 10 disappears and appears at 23.5° 212 indexed (002), exhibiting that the Graphene oxide is a highly oxidized product. No extra visible peaks 213 214 of graphene oxide could be detected in all the XRD patterns owing to the less content and stacking of GO in the ternary nanocomposite, (Zhou et al., 2019). The fewer XRD patterns of GO in ternary hybrids 215 216 shows stacking of GO and its function as a substrate for nucleation and growth of MoS₂ upon exposure to hydrothermal treatment (Li et al., 2015) The absence of any impurity peak in the Mn-BMG sample 217 infers the effective substitution of divalent cations (Mn^{2+}) in the bismuth tungstate matrix. The peaks in 218 the XRD pattern of Mn-doped samples are lower in intensity as compared to the undoped ternary 219

220 composite sample. Additionally, a peak shift to higher diffraction angles position has also been observed in the Mn-doped sample as compared to the undoped sample suggesting that the Mn^{2+} ions successfully 221 doped the host ternary composite. This peak shift suggests changes in interstitial Bi₂WO₆ unit cell 222 volume as a result of doping. The ionic radius of Bi⁺² ions is larger than that of Mn²⁺ions, hence Mn²⁺ 223 ions can enter Bi₂WO₆ host lattice and substitute Bi⁺² ions, causing the XRD pattern to shift to higher 224 225 diffraction angles (inset of fig.3.2.) Similar outcomes of Mn doping in host lattice were also observed 226 by Reddy et al (Reddy et al., 2019). The average particle size was calculated using Debye-Scherrer 227 formulae equation 3:

$$D = 0.9\lambda/\beta cos\theta \qquad \text{Eq.....(2)}$$

D is the particle size, λ is the wavelength of CuK α (i.e., 0.154 nm), β explains the intensity at full-width half maximum of diffraction line which is expressed in radians and θ refers to Bragg's angle. The average particle size of undoped and doped particles was found to be 24.2nm and 20.1nm respectively. The smaller crystallite size of Mn-doped nanocomposite can be referred to as the better growth of nanoparticles onto support media but having restricted dimensions which may distort host lattice due to insertion of Mn⁺² ions decreasing nucleation and grain growth (Chanu et al., 2019).



235 236

Fig.2 X-ray diffraction patterns of (a) BG (b) BMG and (c) Mn-BMG nanocomposites

237 **3.3 SEM analysis:**

238 The morphology and microstructure of as-synthesized pristine and doped samples were investigated by 239 Scanning electron microscope (SEM) at various resolutions. SEM images determined the morphological changes after doping in the surface of the ternary nanocomposite. After combining Mn-240 BWO with MoS₂ and GO the ternary hybrid showed a hierarchical structure showing the assembly of 241 plenty of bismuth tungstate nanorods interlaced and flower-like nanostructured MoS₂ distributed over 242 stacked graphene sheets fig 3(c). The MoS₂ exhibited nanoflowers-like morphology as in fig 3.3 (a) and 243 are clearly shown growing on the edges alongside stacking of graphene oxide ternary heterostructure 244 245 of SEM images fig (3.3b). The undoped ternary composite exhibited severe agglomeration due to the stacking of graphene oxide. The exterior coarse texture of BW nanoplates is because of coverage and 246 stacking of GO sheets However, this agglomeration notably weekend after the introduction of Mn in 247 the nanocomposite. This suggests that Mn doping effectively suppressed the severe aggregation of 248 particles (Dou et al., 2020). The Mn-doped BWO-graphene shows uniformed surface nanorods and less 249 250 agglomeration. The ternary doped nanocomposite with all the components present, well separated with 251 minimized aggregation, uniform structure and Mn particles uniformly attached to BW rods fig 3.3 (c 252 &d). Hence, the incorporation of Mn in the host lattice influenced the morphology of the particles.





Fig.3 SEM images of (a) MoS₂ nanoflowers (b) BMG (c &d) Mn-BMG

256 **3.4 EDX analysis:**

EDX elemental mapping confirmed the chemical configuration and distribution of Bi, W, C, S, N, O according to their nominal weight percentages as shown in fig 3.4. The undoped ternary BWO-GO/MS did not show the presence of any impurity atom whereas the EDX pattern of doped ternary hybrid shows Mn atom in I.68 Wt% which is also confirmed by the XPS spectra, suggesting successful incorporation of Mn in ternary nanocomposite according to their nominal chemical stoichiometry amounts. It also confirms the presence of all other elements including Bi, W, O, C, Mo, S and Na. Little changes in the wt% of all elements is observed after the hydrothermal processing.



- 265
- 266

Fig.4 EDX spectra of (a)BMG (b) Mn- BMG ternary nanocomposites

3.5 XPS analysis:

The composition and chemical states of all the elements present and the interactions among all the three 268 269 components in the ternary nanocomposite is analyzed by XPS spectra. The elemental composition of 270 BMG and Mn-BMG was investigated through XPS survey scan which revealed the peaks of Bi (165.06eV), W(37.81eV), O1s (534.82eV), S (163 eV), C (284.17eV) and Mn as shown in fig (3.5a) 271 while the atomic percentage of detected elements were shown in fig (3.5b). The Bi 4f peak appearing 272 at a binding energy of 165.067eV ascribed to Bi 4f₇ and Bi 4f₅ Bi⁺³ ions in Bi₂WO₆ nanoparticles. The 273 O1s peak in the ternary component at a binding energy of 536.0eV is assigned to the metal-oxygen 274 275 bonds of Bi₂WO₆ and also to the adsorbed water molecules on the catalyst's surface. 276 The C1s peak at 283.74eV is possibly credited to the adventitious residual carbon as also confirmed

from previous studies (Yang et al., 2015). A peak earing at a binding energy of 1011eV is assigned to

278 Mn^{2+} , respectively, indicating the presence of Mn^{+2} . Characteristic peaks appearing at 648.04eV and



684.27eV are ascribed to Mn2p_{3/2}, Mn2p_{1/2}, Mn2s respectively. Thus XPS data confirms the presence 279 280 of Mn ions in doped ternary nanocomposite (Dou et al., 2020). The XPS of Wf₄ can be detected at 37.81eV suggesting W $4f_{5/2}$ of W₆₊. The peak at 163 eV is the S2p peak matched exactly well with the 281 S^{2-} binding energy which is required for MoS2 formation. Also, the peaks at 229.60 confirm the 282 presence of Mo 3d5/2 and Mo $3d_{3/2}$, indicating the dominant presence of Mo⁴⁺ species (Senthil et al., 283 2019). The atomic percentage of S2p, O1s, C1s, W4f, Bi4f, Mo3d, Mn2p3 is 40.4%, 26.9%, 24.2%, 284 3.3%, 2.1%, 1.6%, 1.5% respectively in Mn-BMG, while the undoped counterpart comprised of C1s, 285 O1s, S2p, Bi4f, Mo3d, W4f in 37.1%, 28.7%, 27.4%, 4.6%, 2.1%, 0.2% atomic percentages 286 respectively. 287

288

289

Fig.5 XPS survey spectrum of (a) BMG (b) Mn-BMG and (c) atomic percentages of BMG and Mn-BMG

292

293 **3.6 Optical Analysis**

294 The optical response was measured by observing UV-VIS absorption spectra and determine the 295 bandgap energies of pure and Mn-doped ternary nanocomposite in the range of 200-900nm. The 296 absorption edge of the Mn-doped BWO-GO/MoS₂ nanocomposite was found in the visible region using 297 the tauc plot. The Mn-doped BMG showed an even greater range of light absorption to the entire visible 298 spectrum in comparison to the pristine ternary sample. The Mn doping in ternary heterojunctions 299 improved the optical absorption providing a possibility to enhance photo-induced electron-hole pairs generation ultimately enhancing the photodegradation in the visible region. The bandgap energies of 300 301 the catalysts were estimated using the formula given in equation 4.

$$(\alpha h v)^2 = B (h v - Eg) \qquad \qquad \text{Eq } \dots (3)$$

Where Eg =hv when $\alpha hv = 0$ and the $(\alpha hv)^2$ is plotted against energy in eV. The absorption energy taken at hv value where it then extrapolates to α =0 as shown in Fig. 5. The estimated band gap energies were found to be 2.4 eV and 2.2 eV for BMG and Mn-BMG. Here, α , B and v are described as the absorption coefficient, proportionality constant and the light frequency. The optical studies show that all the photocatalysts show strong absorption in the visible light region.

309



310

Fig.6 Bandgap energies estimation of (a) BG and (b) Mn-BMG (c) BMG by Tauc plot method with
inset showing the UV-VIS absorption spectra of the BMG and Mn-BMG

313

4. Effect of Operational parameters:

315 **4.1 Effect of pH:**

In the photodegradation process, the key factor that directly affects the photoefficiency of the 316 317 photocatalytic system is the pH of the aqueous solution. The pH governs the surface charge 318 characteristics of the photocatalysts according to their point of zero charges (pH_{PZC}) and also the adsorption capacity of the organic molecules on the surface of the catalyst (Ahmadi et al., 2020). The 319 320 effect of pH on the photodegradation of MB was examined by changing pH values from 2 to 9 keeping 321 all other factors constant (catalyst dose=50 mg/100mL for doped and undoped ternary and 40mg/100ml 322 for binary BG, Initial dye concentration: 10ppm, and irradiation time 60 min). The pH of dye solutions was maintained using 0.1 M and 1mM solutions of sodium hydroxide hydrochloric acid. Methylene 323

324 blue is a cationic dye hence it is positively charged upon dissociation in water. The pH_{PZC} value for Mn-325 BMG was determined at pH 6.2. At pH values lower than the pH_{PZC} , the surface of the catalyst is 326 positively charged and the positively charged MB dye molecules are electrostatically repulsed. In the acidic medium, the active sites on the catalyst's surface are weak to produce hydroxl radicals, 327 consequently decreasing the dye degradation (Nie et al., 2014). At pH values above 6.2, the catalyst's 328 surface is negatively charged and the cationic MB dye molecules were electrostatically attracted 329 towards the catalyst surface, enhancing the adsorptive property of dye molecules. The active sites on 330 331 the surface of the catalyst showed increased hydroxyl radicals production resulting in the increased MB degradation in the basic medium. Hence, the ternary Mn-BMG showed 99% photocatalytic degradation 332 in an alkaline medium at optimum pH of 8 under visible light irradiation of 60 minutes as shown in fig. 333 7(a) The undoped ternary BMG at pH 8 and binary BG at pH 7 showed 95% and 92% degradation 334 335 respectively in the presence of sunlight for about the same time duration.

4.2 Effect of catalyst dose:

The amount of catalyst used is another important parameter for the evaluation of photocatalytic 337 338 performance and economic cost. To understand the relation among catalyst loading and photodegradation efficiency, degradation of 10ppm MB dye solution was investigated in the range of 339 340 10-80mg/100mL, keeping all others factors constant (pH=8) for Mn-doped and undoped ternary,7 for 341 binary BG, catalyst dose= optimum of each catalyst, and irradiation time 60 min. Photocatalyst dose 342 ranging from 10mg-100mg/L were used to study the influence on degradation. The photodegradation efficiency of Mn-BMG enhanced from 69% to 99% as the catalyst dose was increased from 10mg to 343 344 50mg. The increase in catalyst dose enhances the number of active sites on the catalyst's surface, 345 consequently enhancing the production of hydroxyl and superoxide radicals. The undoped and doped ternary composites showed enhanced photodegradation at an optimum catalyst dose of 50mg/ml and 346 347 binary BW-GO at 40mg/100ml as shown in graph b of fig 7(b). An increase in catalyst dose above 50 348 mg showed a slight increase up to 97% and started decreasing when it went beyond 100mg. This 349 reduction in degradation is owing to the interception of light preventing light photons to reach the 350 surface of adsorbed contaminant because of agglomeration of catalysts at their higher doses and 351 subsequent unavailability of active sites (Hu et al., 2019). Hence, catalyst dose decides the limits of a 352 catalyst for particular organic pollutants in wastewater above which the rate of degradation eventually decreases. Graphene oxide-supported composites prevent agglomeration and show effective 353 354 degradation even at higher concentrations (Tabasum et al., 2021).

4.3 Effect of initial dye concentration:

356 The dye degradation is highly dependent upon the concentration of dye adsorbed on the catalyst surface.

- 357 The effect of initial dye concentrations ranging from 2-20ppm was studied for the photodegradation of
- 358 MB by Mn-BMG and all other nanocomposites keeping all other factors constant at their optimal values.
- 359 At initial low MB load, maximum degradationn was attained which is linked to the availability of

360 maximum active sites and also due to the reaction between dye molecules and hydroxyl radicals formed 361 on the catalyst's surface. As the dye concentration is increased, active sites on the catalyst surface become saturated as more dye molecules are adsorbed on the surface. The number of photons reaching 362 363 the surface of the catalyst becomes fewer causing less generation of hydroxyl radicals for effective 364 degradation of MB molecules (Nawaz et al., 2020). The performance of all photocatalysts is shown in graph c of fig 4.1. All catalysts showed maximum degradation at a methylene blue concentration of 365 10ppm. The ternary Mn-BMG nanocomposite showed above 90% degradation even at higher MB 366 367 concentrations. This is because the ternary heterojunction has enhanced surface area and can adsorb enhanced concentration of MB molecules. The vacant active sites available after the degradation of 368 369 adsorbed dye molecules on the surface of the photocatalyst are rapidly occupied by other dye molecules near the catalyst surface (Mushtag et al., 2020). This results in enhanced MB degradation even at higher 370 371 dye loads by the novel Mn-BWO-GO/MoS₂ as compared to other photocatalysts



The maximum energy that is required for the initiation of photocatalytic reaction comes directly from the light energy hence, the conditions of light illumination have a great impact on the overall performance of photocatalysts. Fig shows the decomposition efficiency of MB over Mn-BMG, BMG and BG under sunlight irradiation. Under the optimum conditions for all catalysts, The dye solutions were exposed to light irradiation for various for about an hour. The UV-VIS absorption spectra of treated MB solutions after irradiation under sunlight were recorded in the 400-800nm range each after

388 ten minutes. The spectral changes in the treated dye solution were observed by plotting absorbance data 389 as a function of time. The trends show an increase in the degradation as time increases as shown in fig S1(a, b, c for BMG, Mn-BMG, and BG) of supplementary information. The dye color starting 390 disappearing right after 30 minutes of irradiation and complete degradation was observed around 60 391 392 minutes of irradiation using novel Mn-BMG whereas, the decrease in the intensity was gradual with 393 time using ternary BMG and binary BG. The rapid decrease in intensity upon light irradiation is 394 signifying that chromophore groups responsible for imparting colour are breaking down slowly. 395 Further, the kinetic models are applied to this data.

396

397 4.5 Kinetics of Photodegradation reaction.

Two kinetic models first-order and second-order were studied for the photodegradation of dye
methylene blue by the doped and undoped ternary nanocomposites. The expression for each model is
expressed in equations 3-4

401

403

402	First-order	kinetics:

 $ln\frac{co}{ct} = k_1 t \qquad \qquad \text{Eq.....} \tag{3}$

404 405

406 Second-order kinetics:

- 407 $\frac{1}{ct} \frac{1}{co} = k_2 t$ Eq..... (4) 408
- 409

410 Fig. 8(b), shows a linear relationship between ln(CO/C) and reaction time, where Co and C are the initial concentrations, indicating that the photodegradation of methylene blue by doped and undoped ternary 411 nanocomposites follows first-order kinetics. The plot of Co/C versus time represents a straight line 412 413 where the slope of which upon linear regression equals the apparent first-order rate constant k. The higher rate constants k of Mn-BMG for the first-order reaction than undoped ternary indicates that Mn-414 doped BWO-GO is more effective under sunlight than BWO-GO/MoS₂. The values of R² for the first-415 order reaction for doped and undoped ternary composites are 0.9967 and 0.993 respectively, suggesting 416 417 that the ternary hybrids effectively follow the first order reaction. The rate constant k of the doped component is more than that of undoped ternary revealing that doping has caused an acceleration in the 418 419 degradation rate under visible light A. The higher rate constant of doped BMG than other photocatalysts 420 explains that the Mn-BMG shows more photoefficiency than others. Comparison of correlation 421 coefficients for a first order and second order are shown in table 1.



Fig.8 MB % removal as a function of time (a) photocatalytic degradation of MB in the presence of
 sunlight, catalysts under optimum conditions (a) First order kinetic model fitting

426

423

427	Table.1 Correlation coefficient	s (R ²) an	d kinetic parameters	s for MB	(Co=10ppm)	degradation
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	Experi	mental conditio	ns	First-order kinetics		Second-order kinetics	
Photocatalysts	pН	Photocatalyst	Initial dye	$k1(\min^1)$	R^2	K2	R^2
		dose(mg/L)	concentration(ppm)			$(L\mu mol^1 min^1)$	
Mn-BMG	8	50mg	10ppm	0.0866	0.9967	0.43215	0.729
BMG	8	50mg	10ppm	0.02899	0.993	0.14207	0.8616
BW	7	40mg	10ppm	0.0255	0.987	0.0886	0.8487

428

429 **4.6** Type of the Photocatalyst and the role of dopant species:

430 The type of photocatalyst used is the most imperative parameter for greater photocatalytic degradation. 431 The bandgap, crystallinity, porosity, surface area and nature of dopant all determine the effectiveness of catalyst in the visible light region. The ternary nanocomposites as compared to binary composites 432 show increased degradation due to the synchronized activation of the charge transfer mechanism 433 434 between the various semiconductor materials. The transfer of charges between the conduction bands of 435 two metal oxides leads and a support material leads to enhanced charge separation and less agglomeration by increasing the specific surface area ultimately offering more active sites. The ternary 436 437 undoped BMG showed 94% degradation in visible light as compared to the binary counterpart. 438 The addition of dopant species in the semiconductor ternary system further improved the photocatalytic 439 activity which is mainly ascribed to alteration in the bandgap energy, following the transfer of energy.

- 440 The addition of dopant creates additional energy levels and leading to better separation and transfer of
- 441 photogenerated charge carriers to the surface of the catalyst. This separation mainly depends upon two
- 442 factors including large surface area and the light absorbed by the catalyst (Aydoghmish et al. 2019).
- 443 Therefore, the doped ternary Mn-BMG showed strong visible light region absorption and greater
- degradation for methylene blue up to 99.7% in one hour.

445 **4.7 Degradation under UV irradiation**

A sole experiment was done in UV radiations considering the optimized conditions specific for all the photocatalyst. Methylene blue was degraded almost completely in UV irradiation only after exposure of half an hour by Mn-BMG. Undoped BMG and binary BG also performed well under ultraviolet light and their ternary heterojunction has shown more than 90% degradation under UV light. Similar results of greater MB degradation by ternary nanocomposite were shown earlier in UV light degradation (Aydoghmish et al., 2019).

452

453 5. Radical Scavenging and proposed mechanism

- 454 The determination of active species involved in the photocatalytic process is significant for understanding the degradation mechanism of organic pollutants. In the degradation of dyes, several 455 456 active radical species such as hydroxyl radicals (OH \cdot), holes, (h⁺), electron (e⁻) superoxide anion radical $(\cdot O^{-2})$ play a vital role (Rajendran et al., 2018). Hence, to investigate the key active species 457 involved during the photocatalytic process, a radical scavenging experiment was performed. 5mM of 458 459 EDTA (ethylene-diamine-tetra-acetate), K₂Cr₂O₇ (potassium dichromate), DMSO (dimethyl sulfoxide) 460 were used to scavenge holes, electrons and hydroxyl radicals. The experiment was performed under 461 sunlight. From fig.10(b) it is clear that EDTA is the main radical scavenger in the degradation process of MB by MnBW-GO/MoS₂, as the addition of EDTA decreased the degradation values from 99 to 462 463 48.83. The addition of DMSO decreased the degradation from 99 to 55% indicating that hydroxyl 464 radicals also have a contributing role in addition to holes. Very little change was observed by the 465 addition of K₂Cr₂O₇ indicating that electrons do not play a major role in MB degradation by Mn-BWO-466 GO/MoS_{2.}
- The proposed mechanism of MB degradation by Mn-BWO-GO/MoS₂ suggests that holes and hydroxyl 467 radicals are the main active species. Holes are directly responsible for degradation and producing more 468 hydroxyl radicals. Upon sunlight irradiation, both Bi_2WO_6 and MoS_2 are excited and produce electrons 469 470 and holes. The photoexcited electrons get transferred from the conduction band of MoS₂ having a 471 negative potential towards the conduction band of Bi_2WO_6 having a positive potential. While the 472 photoinduced holes accumulate above the Valence band of MoS₂. The reactive holes accumulated on the surface of MoS₂ directly oxidize MB dye molecules adsorbed on the catalyst surface. Graphene 473 oxide has a large surface area and electrical conductivity acts as a mediator and transfers electrons to 474 and from MoS₂ and Bi₂WO₆. Mn doping introduces additional energy levels in BWO, hence photo-475 476 induced electrons and holes take longer to recombine. Eventually, the photo-response of the ternary

- heterojunction is increased towards visible light. A similar mechanism has been proposed by lv et al., 477
- (Lv et al., 2017). Hence, holes are responsible for the degradation of dyes and production of hydroxyl 478 479 radicals whereas, the influence of electrons and superoxide anions is relatively less. Fig 9 shows the 480 degradation mechanism by ternary Mn-BWO-GO/MoS₂.
 - Reduction reaction O₂ ee GO СВ ee e CB MoS₂ Mn



- 481
- 482





484

485 Fig. 10 (a) Study of various radical scavengers (b) reusability of Mn-BMG, BMG, BW nanocomposites

6. Reusability: 486

487 Under the optimized conditions for each photocatalyst, the stability of catalysts was confirmed by using 488 the catalysts repeatedly in five successive trials. For this, the catalysts are separated from the treated 489 MB dye solution by centrifugation then rinsed thrice using distilled water and later dried up at 60°C in an electric oven. The catalysts are then weighed and investigated for their photocatalytic efficiency. All 490 491 the reusability trials are run at the same optimized conditions for each catalyst i.e at pH 8, catalyst dose-50mg/100m, IDC 10ppm for doped undoped ternary nanocomposite, and pH 7, catalyst dose 492 40mg/100ml, IDC 10ppm for binary BW-GO. No substantial loss in photocatalytic activity was 493 observed after five runs with just a 10% decrease in catalytic activity fig.10(b). The results revealed 494 495 that the ternary doped composite exhibits high mechanical stability and reuseability.

496 **Conclusion:**

497 The novel ternary hybrid Mn-BWO-GO/MoS₂ has found valuable applications for Organic pollutants 498 degradation. The doped ternary photocatalyst was synthesized by an insitu hydrothermal method. The 499 ternary hybrid showed excellent degradation for dye Methylene blue under sunlight irradiation. Undoped ternary BW-GO/MoS₂, and binary BW-GO were also prepared to compare the degradation 500 501 efficiencies of doped and undoped ternary hybrids with binary composite. The Mn-BWO-GO/MoS₂ showed 99% MB degradation in an hour. The improved photocatalytic efficiency was credited to the 502 ternary heterojunction formed as a result of BWO, MoS₂, and GO. The ternary heterojunction has the 503 504 characteristics of all three components and the Mn doping further suppresses the electron-holes 505 recombination. The photostability of photocatalysts was evaluated by five consecutive runs with less than 10% reduction in degradation. 506

507 **Declarations:**

508 Ethical Approval and Consent to Participate Not applicable

- 509 *Consent to Publish* All the author agree to pulish this article
- 510 Authors Contributions

511 Noor Tahir: Data curation, Visualization, Investigation, Writing - original draft, Writing -

review & editing. **Muhammad Zahid:** Conceptualization, Project administration, Resources,

513 Supervision, Writing - review & editing. Ijaz Ahmad Bhatti: Conceptualization,

- 514 Methodology, Software, Writing review & editing. Yasir Jamil: Visualization, Investigation,
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- 519 *Competing Interests*
- 520 The authors have no conflicts of interest to declare that are relevant to the content of this article.
- 521 Availability of data and materials

- 522 The datasets used and/or analyzed in this study are available in the manuscript and supplementary
- 523 information additional information can be asked from the corresponding author upon request.

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- 626
- 627



Figure 1

FTIR spectra of (a) BMG and (b) Mn-BMG ternary nanocomposites





X-ray diffraction patterns of (a) BG (b) BMG and (c) Mn-BMG nanocomposites





SEM images of (a) MoS2 nanoflowers (b) BMG (c &d) Mn-BMG



EDX spectra of (a)BMG (b) Mn-BMG ternary nanocomposites

Figure 5

XPS survey spectrum of (a) BMG (b) Mn-BMG and (c) atomic percentages of BMG and Mn-BMG



Bandgap energies estimation of (a) BG and (b) Mn-BMG (c) BMG by Tauc plot method with inset showing the UV-VIS absorption spectra of the BMG and Mn-BMG





Optimization of reaction parameters using Mn-BMG, BMG and BW (a) pH, (b) catalyst dose, (c) Initial dye concentration





MB % removal as a function of time (a) photocatalytic degradation of MB in the presence of sunlight, catalysts under optimum conditions (a) First order kinetic model fitting



Figure 9

Photocatalytic degradation mechanism of Dye Methylene blue by Mn-BWO-GO/MoS2



(a) Study of various radical scavengers (b) reusability of Mn-BMG, BMG, BW nanocomposites

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