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Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalysts: Principles, mechanisms and 1 photocatalytic applications 2 3 Tamer M. Khedr<sup>1,2,\*</sup>, Kunlei Wang<sup>1,3</sup>, Damian Kowalski<sup>4</sup>, Said M. El-Sheikh<sup>2</sup>, Hany M. Abdeldayem<sup>5</sup>, Bunsho Ohtani<sup>1</sup> and Ewa Kowalska<sup>1,\*</sup> 4 5 6 <sup>1</sup>Institute for Catalysis, Hokkaido University, N21, W10, Sapporo 001-0021, Japan 7 <sup>2</sup>Nanomaterials and Nanotechnology Department, Advanced Materials Institute, Central Metallurgical Research and Development Institute (CMRDI) P.O. Box: 87 Helwan, Cairo 11421, 8 9 Egypt 10 <sup>3</sup>Northwest Research Institute, Co. Ltd. of C.R.E.C., 730000, Lanzhou, P.R. China <sup>4</sup>Faculty of Chemistry and Biological and Chemical Research Centre, University of Warsaw, Zwirki 11 12 i Wigury 101, 02-089 Warsaw, Poland 13 <sup>5</sup>Chemistry Department, Faculty of Science, Ain Shams University, 11566 Abassia, Cairo, Egypt 14 \* Correspondence: tamerkhedr56@gmail.com (T.M.K.) and kowalska@cat.hokudai.ac.jp (E.K.) 15 16 17 **Abstract:** The development of novel photocatalysts for efficient utilization of solar energy 18 is highly essential for the most critical humanitarian challenges, i.e., energy and water crises as well as environmental pollution. Bismuth tungstate (Bi<sub>2</sub>WO<sub>6</sub>), an outstanding Aurivillius 19 phase perovskite, has attracted intensive attention as a visible-light-responsive photocatalyst 20 21 because of its non-toxicity, low cost, and outstanding physicochemical characteristics, i.e., nonlinear dielectric susceptibility, ferroelectric piezoelectricity, pyroelectricity, catalytic 22 behavior, modifiable morphology, strong oxidation power, and good photochemical 23 stability. However, the photocatalytic activity of bare  $Bi_2WO_6$  is restricted because of the 24 inherent drawbacks such as poor light-harvesting efficiency, weak reduction potential, 25 relatively low specific surface area, the fast recombination rate of photoinduced charge 26 carriers, and thus poor quantum yields of photocatalytic reactions. Moreover, the 27 impossibility of simultaneous strong redox ability (demanding wide bandgap) and high 28 29 light-harvesting efficiency (requiring narrow bandgap) is considered as a big challenge for the practical application of Bi<sub>2</sub>WO<sub>6</sub>. Undeniably, the construction of Z-scheme 30 photocatalytic systems is recommended strategy to overcome the above-mentioned 31

32 disadvantages because of the efficient spatial separation of photogenerated charge carriers and the boosting the redox performance. This review summarizes the principles and recent 33 developments on Z-scheme photocatalytic systems with special emphasis on the Bi<sub>2</sub>WO<sub>6</sub>-34 based photocatalysts, including the types, photocatalytic mechanisms and practical 35 applications. Moreover, major differences between type-II heterojunction and Z-scheme 36 photocatalyst have also been discussed. Additionally, the significant role of unique 37 38 structures (e.g., core-shell and 2D/2D) for the improvement of photocatalytic activity of Zscheme photocatalyst has been presented. Indeed, Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalysts 39 40 have exhibited superior photocatalytic activity for various applications. For example, they show high photocatalytic activity towards water/wastewater treatment (removal of organic 41 and inorganic pollutants, as well as microorganisms), air purification (decomposition of 42 volatile organic compounds and inorganic matters), "green" energy conversion (e.g., 43 44 generation of H<sub>2</sub> and CH<sub>4</sub> fuels under solar irradiation), and organic synthesis. It is thought that this remarkable activity of Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalysts might be attributed to 45 46 the efficient solar light harvesting, separation and further transfer of charge carriers and 47 strong redox ability. To the best of our knowledge, the present paper is the first attempt to summarize the Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalytic reactions, providing important insights 48 and up-to-date information for the scientific community to fully explore the potential of Bi<sub>2</sub>WO<sub>6</sub>-49 50 based photocatalysts for renewable environmental remediation, energy conversion, and chemical 51 synthesis.

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53 Keywords: Bi<sub>2</sub>WO<sub>6</sub>; Z-scheme photocatalyst; environmental purification, energy
 54 conversion; organic synthesis; green energy; solar photocatalysis, coupled semiconductors

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## 57 **1. Introduction**

Fast-growing environmental pollution and the global energy crisis are probably the most 58 serious threats to human life and property. Therefore, various methods of environmental 59 purification and energy generation have been intensively examined in recent times. For 60 example, advanced oxidation processes (AOPs), also known as advanced oxidation 61 technologies (AOTs), have been proposed as efficient methods, especially for environmental 62 63 treatment, since reactive oxygen species (ROS, mainly hydroxyl radicals), formed in-situ, are highly active against organic and inorganic compounds, as well as microorganisms. 64 Usually, ROS, e.g., hydroxyl radical ('OH), superoxide radical ('O2), sulfate radical 65  $(SO_4^{-})$ , ferrate radical (FeO<sub>4</sub><sup>2-</sup>) and ozone (O<sub>3</sub>), are generated using primary oxidants, e.g., 66 hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), persulfate, ferrate, permanganate, and oxygen or via physical 67 68 methods. Unfortunately, majority of these methods, e.g., supercritical water oxidation (SCWO), wet air oxidation (WAO) and UV-based irradiation systems (UV/O<sub>3</sub> and 69  $UV/H_2O_2$ ), require high energy demands, continuous feeding of reagents (e.g.,  $H_2O_2$ ,  $O_3$ ) 70 71 and high investment and operating costs. It has been thought that heterogeneous photocatalysis is probably the best AOP method, as solar energy might be used for 72 photocatalyst excitation even for wide-bandgap semiconductors (e.g., TiO<sub>2</sub> and ZnO). 73 Accordingly, various reports have been published on the synthesis, property-governed 74 75 activities, mechanism clarifications, activity improvements and possible applications of 76 heterogeneous photocatalysis, especially for TiO<sub>2</sub> photocatalysts (the most broadly 77 investigated photocatalyst) [1-4]. However, titania must be excited with UV irradiation, and thus only slight part of solar radiation (ca. 2%) might be efficiently used for photocatalytic 78 79 reactions. Therefore, various methods of wide-bandgap semiconductors' modification have 80 been proposed, including doping, surface modification and formation of composite photocatalysts. It is thought that the last-listed above, i.e., composite photocatalysts, might 81

be the most attractive since both inhibition of charge carriers' recombination and appearance
of vis activity might be achieved by proper selection of constituent parts. It should be
pointed out that not only the modification of well-known materials, but also preparation of
novel photocatalysts with vis response has been intensively studied, e.g., graphitic carbon
nitride (g-C<sub>3</sub>N<sub>4</sub>), WO<sub>3</sub>, CuO, Cu<sub>2</sub>O, SrTiO<sub>3</sub>, BiVO<sub>4</sub>, Ta<sub>3</sub>N<sub>5</sub>, TaON, CaTaO<sub>2</sub>N, SrTaO<sub>2</sub>N,
BaTaO<sub>2</sub>N (e.g., [5-9]).

88 First, the basic mechanism of heterogeneous photocatalysis should be presented. Three sequential and integral steps are as follows: 1) photocatalyst (semiconductor) 89 90 activation under irradiation with energy equal or larger than its energy gap (Eg), resulting in charge carriers' generation, i.e., electrons in the conduction band (CB) and holes in the 91 92 valence band (VB), 2) charge carriers' transfer, i.e., migration to the photocatalyst surface or 93 recombination (bulk/surface), 3) surface redox reactions (Fig. S1). It should be pointed out 94 that the photocatalysis thermodynamics, which is governed by the band levels (CB and VB) and redox potentials of adsorbed reactants (acceptor and donor), and the photocatalysis 95 96 kinetics (i.e., charge generation, migration and consumption), are highly critical for the photocatalytic efficiency [10,11]. Thermodynamically, the redox potential of oxidant 97 (acceptor) should be below (more positive than) the conduction band minimum (CBM) of 98 the semiconductor to proceed the reduction reactions, whereas the redox potential of the 99 100 reductant (donor) should be more negative than the valence band maximum (VBM) to attain 101 the oxidation reactions. Therefore, for superior photocatalytic activity, the semiconductor needs to have strong redox ability, which demands a wide bandgap (from the 102 thermodynamic viewpoint) and wide light-harvesting range, which requires a narrow 103 104 bandgap (from the kinetic viewpoint). Obviously, the balance between thermodynamics and kinetics to attain high-performance photocatalysts is impossible to be achieved in traditional 105 106 (i.e., single component) photocatalyst. Therefore, the multi-component (heterojunctions)

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photocatalysts, composed of two or more semiconductors with different oxidation and redox potentials, have been proposed for efficient light harvesting and high quantum efficiencies.

Three types of heterojunction between two semiconductors could be distinguished, 109 i.e., i) type-I ("straddling gap", Fig. 1a, e.g., α-Fe<sub>2</sub>O<sub>3</sub>/TiO<sub>2</sub>), ii) type-II ("staggered gap", Fig. 110 1b, e.g., Cu<sub>2</sub>O/TiO<sub>2</sub>), and iii) type-III ("broken gap", Fig. 1c). Obviously, the type-II 111 heterojunction is the most recommended for photocatalytic applications, because of the 112 113 spatial separation of charge carriers (an opposite direction of electron and hole transfer), resulting in inhibition of charge carriers' recombination, and thus an increase in the 114 115 photocatalytic efficiency. Typically, the type-II heterojunction photocatalysts contain two semiconductors (SCI and SCII) with staggered band structure configuration, as shown in 116 Fig. 1b. Under irradiation (with an energy equal or larger than their band gaps), both SCI 117 118 and SCII are simultaneously excited, and thus formed charge carriers (electrons in CB and holes in VB) might migrate between semiconductors, i.e., electrons from SCI (of more 119 negative CB) to SCII (of less negative CB), and holes from SCII (of more positive VB) to 120 SCI (of less positive VB), resulting in accumulation of electrons and holes in SCII and SCI, 121 respectively. Although, the separation of charge carriers in type-II heterojunction results in 122 inhibition of charge carriers' recombination, the redox ability (the thermodynamic 123 viewpoint), is lower than that in singlet-component photocatalysts since the holes and 124 125 electrons are accumulated at the lower and higher potential of VB and CB (i.e., less positive 126 and less negative), respectively. Additionally, the electrostatic repulsions between symmetrically charged components restrict the charge carriers' migration between two 127 semiconductors (the kinetic viewpoint). 128



**Fig. 1.** The schematic drawings for semiconductor-based heterojunctions: type I (**a**), type II (**b**) and type III (**c**). Ox: oxidant, Red: reductant.

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Based on these aspects, a dilemma on the efficient heterojunctions has been raised, i.e., how to prepare efficient and stable heterojunction photocatalysts with efficient separation of charge carriers and strong redox ability. Therefore, it has been proposed that the new system must be designed to overcome these limitations. Accordingly, the nature has inspired scientists as the photosynthesis occurring in the green plants might be adapted for
the photocatalytic reactions by the construction of the artificial photosynthesis systems,
named as Z-scheme heterojunctions [5,12-14]. The Z-scheme mechanism occurs naturally in
green plants through photosynthesis process ("natural Z-scheme photocatalysis"), in which
the charges migration route shows a two-step photoexcitation, which looks like the letter
"Z" turned 90° counterclockwise ("the zigzag"), as shortly presented in SI (Fig. S2).

The artificial Z-scheme photocatalytic system (ZSPS) is generally composed of 143 144 two semiconductors (SCI & SCII) and charge mediator, classified into three groups, as follows. (1) Liquid-phase Z-scheme/traditional Z-scheme (1<sup>st</sup>-generation Z-scheme, Fig. 2a), 145 which contains an aqueous redox-pair mediator, i.e., shuttle redox mediator, including 146  $\text{Fe}^{3+/2+}$ ,  $\text{IO}_3^{-}/\text{I}^{-}$  and  $[\text{Fe}(\text{CN})_6]^{3-/4-}$ . (2) All-solid-state Z-scheme (2<sup>nd</sup>-generation Z-scheme, 147 Fig. 2b), which consists of a solid mediator, such as metals (Ag, Pt, Au), reduced graphene 148 oxide (RGO) and carbon nanotubes (CNTs). (3) Direct Z-scheme (3<sup>rd</sup> generation Z-scheme, 149 Fig. 2c), which does not contain any charge mediator. Upon irradiation, electron/hole pairs 150 are generated in the CB/VB of both semiconductors. Then, the photogenerated low-energy 151 152 electrons (at the CB of SCII) recombine with low-energy holes (at the VB of SCI) directly (direct Z-scheme) or with the help of redox-mediator, whereas the high-energy electrons 153 (strong reducer) and holes (strong oxidizer) in the CB of SCI and the VB of SCII, 154 155 respectively, react with adsorbed compounds via respective redox reactions (Table S1). As mentioned above, the Z-scheme photocatalysts outperform the type-II heterojunction due to 156 strong redox ability, since the photogenerated electrons and holes are more negative and 157 158 more positive, respectively, and they are spatially separated.

159 It should be pointed out that Z-scheme photocatalytic systems might be constructed 160 from semiconductors with visible-light (vis) absorption (narrower bandgap than that in 161 titania), and thus efficient light harvesting at broad solar radiation range could be achieved.

Accordingly, visible-light-responsive (VLR) bismuth-based photocatalysts, such as Bi<sub>2</sub>O<sub>3</sub>, 162 BiOX (X = Br, I), BiVO<sub>4</sub>, Bi<sub>2</sub>MoO<sub>6</sub>, Bi<sub>2</sub>WO<sub>6</sub>, have attracted great attention because of their 163 significant photocatalytic activity for environmental decontamination as well as oxygen 164 evolution during water splitting [15,16]. For example, bismuth tungstate (Bi<sub>2</sub>WO<sub>6</sub>, n-type 165 semiconductor) has recently been considered as a promising material, especially for 166 photocatalytic oxidation reactions, because of its excellent intrinsic physicochemical 167 168 characteristics, such as optical/electronic properties, including VLR (bandgap of ca. 2.8 eV), nonlinear dielectric susceptibility, ferroelectric piezoelectricity, pyroelectricity, strong 169 170 oxidation potential, catalytic behavior, photostability, low cost, and non-toxicity [17-19]. However, the practical application of Bi<sub>2</sub>WO<sub>6</sub> (BWO) is limited by the rapid electron-hole 171 recombination (low efficiency of charge carriers' separation) and low photo-harvesting 172 173 ability. As a class of highlighted strategies, the construction of BWO-based Z-scheme photocatalytic systems with an effective spatial separation of charge carriers as well as 174 strong redox properties and high light harvesting efficiency, has attracted substantial 175 176 research interest for efficient solar-energy conversion (i.e., as broad "green" applications).

Accordingly, this review presents the development of BWO-based photocatalysts, including their types, preparation, characterization, mechanism clarifications and various applications, including environmental purification, solar-fuel generation and synthesis of organic compounds.

Although, it is of importance to follow the historical development of Z-scheme photocatalytic systems (from the first-generation to the third-generation ones) to understand the principles, mechanisms, nano-architecture design, advantageous and shortcomings of various materials, due to journal policy ("to keep the review manuscripts concise and readable") this part has been placed in SI (including Fig. S3-S7).



Fig. 2. The schematic drawings showing artificial Z-scheme photocatalytic systems: liquid-phase Z scheme (a), all-solid-state Z-scheme (b) and direct Z-scheme (c). A: acceptor, D: donor, Ox: oxidant,
 Red: reductant.

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### **192 2. Bi**<sub>2</sub>**WO**<sub>6</sub>**-based Z-scheme photocatalysts**

Bismuth tungstate (Bi<sub>2</sub>WO<sub>6</sub>; BWO) has been considered as a novel and promising VLR photocatalyst for photooxidation of organic pollutants because of its unique structure (a perovskite-type layered), superb physicochemical characteristics, chemical stability, nontoxicity, low cost, strong oxidation power, and vis harvesting ability (band gap of ca. 2.8 eV) [17-19]. BWO can be found in nature as mineral russellite, named in honor of Sir Arthur Edward Ian Montagu Russell (1878-1964), British mineral collector (arthurite mineral was also named in his and Arthur W. G. Kingsbury honor). Russellite is an orthorhombic – pyramidal mineral of various tints (brown-orange, greenish, greenish yellow and light yellow) and ferroelectric properties [20]. Russellite is the simplest member of the Aurivillius phase compounds, i.e., a form of perovskite with the general formula of  $Me_2O_2(Me'_{n-1}R_nO_{3n+1})$ , where Me is a large 12 co-ordinate cation (here Bi), and R is a small 6 co-ordinate cation (here W). The structure of russellite is built up of  $Bi_2O_2^{2+}$  slabs between  $WO_4^{2-}$  layers.

206 The Aurivillius phases are the compounds having (Bi<sub>2</sub>O<sub>2</sub>)(A<sub>n-1</sub>B<sub>n</sub>O<sub>3n+1</sub>) structure, composed of alternating layers of  $(Bi_2O_2)^{2+}$  and perovskite-like blocs. One of this type of 207 208 structures having A site absent in the oxide with a general formula of (Bi<sub>2</sub>O<sub>2</sub>)(BO<sub>4</sub>) is bismuth tungstate Bi<sub>2</sub>WO<sub>6</sub>. For bismuth tungstate, the WO<sub>6</sub> units construct two-dimensional 209 network of corner-sharing octahedra, which is one WO<sub>6</sub> layer thick. At ambient temperature 210 211 the Bi<sub>2</sub>WO<sub>6</sub> crystallizes in orthorhombic crystal system with 29 P2<sub>1</sub>ab symmetry [21,22]. The transition from the space group 29 P2<sub>1</sub>ab to higher symmetry 41 B2cb has been suggested to 212 occur at approximately 660 °C [23], whereas the earlier studies reported high symmetry at 213 lower temperatures [24]. This transition corresponds to the loss of one octahedral tilt mode in 214 the pseudo-perovskite layer [23,25]. The additional phase transition occurs at 960 °C, which 215 corresponds to ferroelectric Curie point. The phase structure formed at T > 960 °C is under 216 debate; according to Watanabe the structure might be similar to BiLaWO<sub>6</sub>, in which the 217 Bi<sub>2</sub>O<sub>2</sub> layers essentially remain unchanged, whereas pseudo-perovskite blocks, constructed at 218 219 the corner-sharing WO<sub>6</sub> octahedra, are reconstructed to WO<sub>4</sub> tetrahedra [26]. In contrast to this observation, McDowell et al. have found that high temperature phase is composed of 220 alternating Bi<sub>2</sub>O<sub>2</sub> layers and perovskite-like layers reconstructed from corner-sharing into 221 edge-sharing WO<sub>6</sub> octahedra [23,27]. The medium temperature phase of Bi<sub>2</sub>WO<sub>6</sub> having 41 222

B2cb symmetry is shown in Fig. 3. The unit cell has lattice parameters of a = 5.5340 Å, b = 5.4998 Å, c = 16.5507 Å and cell volume of 503.7381 Å<sup>3</sup>.





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**Fig. 3.** Structure of  $Bi_2WO_6$  displaying the  $WO_4^{2-}$  and  $Bi_2O_2^{2+}$  layers.

BWO has already been proposed for various applications, including: i) piezoelectric devices as a lead-free material, because of high ferroelectric Curie point (Tc  $\approx$  960 °C) and high electromechanical coupling coefficient, ii) an electrode material for solid oxide fuel cells due to its high oxide ion conductivity, iii) medical applications (tumor radiosensitization, and photothermal/photodynamic therapy of cancer), iv) catalyst, e.g., oxidation of methane to methanol, and v) photocatalyst, as discussed further in this paper [28-30].

For example, Lai et al. prepared nanospherical Bi<sub>2</sub>WO<sub>6</sub> by solvothermal method for 234 photocatalytic degradation of Erichrome Black T dye under solar radiation [30]. However, 235 the rapid recombination of the photogenerated charge carriers, low reduction ability and 236 237 limited light-harvesting efficiency have resulted in low interest for the practical applications of a one-component BWO for photocatalysis [17-19]. Accordingly, increasing efforts have 238 been made to overcome these shortcomings, e.g., by construction of BWO-based Z-scheme 239 240 photocatalysts, including mediated-Z-scheme (all-solid-state Z-scheme) using different solid mediators, such as Ag [31-34], Au [35,36], Pt [37,38], Bi [39-42], Cu [43], Zn [44], oxygen 241

vacancies (OV) [45], polypyrrole (PPy) [46], RGO [47-51], CNT [52-54], and Mxenes
(Ti<sub>3</sub>C<sub>2</sub>) [55], and mediator-free Z-scheme (direct Z-scheme) [56–68] (Fig. 2a).

For instance, novel Z-scheme In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> core-shell photocatalyst was controllably 244 fabricated for the improvement of photocatalytic degradation of tetracycline hydrochloride 245 (TC) antibiotic under vis irradiation (500-W Xe lamp with a UV cut-off filter,  $\lambda > 420$  nm) 246 [69]. First, the pristine In<sub>2</sub>S<sub>3</sub> micro/nanospheres were synthesized by hydrothermal method, 247 248 followed by self-etching. Then, the binary In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> photocatalyst was prepared via template/hydrothermal method with different compositions (In<sub>2</sub>S<sub>3</sub>: 0-40 wt%). The 249 photocatalytic activity of these photocatalysts (In<sub>2</sub>S<sub>3</sub>, Bi<sub>2</sub>WO<sub>6</sub>, and In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> composites: 250 20%In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub>, 30%In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> and 40%In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub>) are shown in Fig. 4a. 251 Although the photocatalytic activity of bare samples  $(In_2S_3 \text{ and } Bi_2WO_6)$  is not high (only 252 40.0% and 45.0% degradation rate, respectively), probably due to the rapid recombination of 253 photoinduced charge carriers, the 30%In<sub>2</sub>S<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> composite shows 2.4- and 2.1-fold 254 higher activity, respectively, and high stability (Fig. 4b). It has been proposed that the activity 255 improvement has been caused by the core-shell morphology, polycrystalline structure, 256 257 enhanced light absorption efficiency, effective charge carriers' transfer and separation through Z-scheme mechanism. The interface in core-shell structure might ensure an 258 interfacial transfer of charges, i.e., from the VB of In<sub>2</sub>S<sub>3</sub> to the CB of Bi<sub>2</sub>WO<sub>6</sub>, generating the 259 built-in electric field, and thus facilitating the migration of charge carriers in accordance with 260 Z-scheme mechanism, as shown in Fig. 4c. Moreover, the core-shell structure (Bi<sub>2</sub>WO<sub>6</sub> shells 261 on the  $In_2S_3$  core) could provide more reactive sites, hence boosting the photocatalytic 262 performance. 263



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Fig. 4. (a) Photocatalytic activity (tetracycline hydrochloride antibiotic degradation during 2-h vis irradiation) data of  $Bi_2WO_6$ ,  $In_2S_3$ , and  $In_2S_3/Bi_2WO_6$  composites prepared for different contents of  $In_2S_3$  (20-40 wt.%), (b) stability data for fresh and recycled sample (30 wt% $In_2S_3/Bi_2WO_6$ ), and (c) schematic drawing showing Z-scheme mechanism for  $In_2S_3/Bi_2WO_6$  composite under vis irradiation. Reproduced from Ref. [69], copyright 2022, with permission from Elsevier.

# 302 2.1. Methods used for Z-scheme mechanism elucidation of Bi<sub>2</sub>WO<sub>6</sub>-based photocatalysts

Various methods have been proposed for the clarification of the mechanism (mainly charge carriers' transfer) of Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalysts, including experimental methods, such as, reactive species trapping (using different scavengers), 5,5-dimethylpyrroline N-oxide (DMPO)-ESR analysis, terepthalic acid-photoluminescence (TA-PL), photocatalytic reduction tests, in-situ X-ray photoelectron spectroscopy (XPS), and band 308 structure characterizations as well as corresponding theoretical analysis (Mott-Schottky 309 analysis and density functional theory (DFT) calculations), as shortly presented in the 310 following subsections. It should be pointed out that using only one method might not give 311 accurate elucidating for the charge carriers' transfer mechanism in the Z-scheme 312 photocatalytic system. Therefore, the combination of different approaches is required to 313 prove the construction of Z-scheme.

# 314 2.1.1. Experimental methods

It is well known that photocatalytic oxidation reactions (oxidative decomposition of organic 315 compounds and microorganisms, and O<sub>2</sub> generation) on the surface of photocatalyst by 316 photogenerated charge carriers ( $e^{-}$  and  $h^{+}$ ) can only proceed when there is sufficient potential 317 to produce reactive oxygen species (ROS), including 'OH (standard redox potential of 318 'OH/H<sub>2</sub>O of 2.4 V vs. NHE) and 'O<sub>2</sub><sup>-</sup>(standard redox potential of O<sub>2</sub>/'O<sub>2</sub><sup>-</sup> of -0.33 V vs. 319 NHE) radicals [70,71]. Therefore, the mechanism of the Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme 320 photocatalytic systems could be elucidated by detecting these ROS generated during 321 photocatalytic reactions. Commonly, ROS could be estimated by following experiments: (i) 322 trapping tests, in which different scavengers are used to capture  $h^+$ , 'OH and 'O<sub>2</sub><sup>-</sup>, (ii) electron 323 spin-resonance (ESR) analysis (5,5-dimethyl-pyrroline N-oxide, DMPO, is used to probe 'OH 324 and 'O<sub>2</sub><sup>-</sup>, producing DMPO-'OH and DMPO-'O<sub>2</sub><sup>-</sup>, respectively), and (iii) photoluminescence 325 (PL) experiments in the presence of nonflourescent terepthalic acid (TA), which reacts with 326 'OH, resulting in formation of highly fluorescent 2-hydroxyterephathalic acid (HTA). 327

For instance, the photocatalytic degradation of antibiotic tetracycline over the binary Bi<sub>2</sub>WO<sub>6</sub>/CuBi<sub>2</sub>O<sub>4</sub> nanocomposite was examined in the absence and the presence of various scavengers, such as disodium ethylenediamine tetraacetic acid (EDTA), tert-butanol (t-BuOH), and 1,4-benzoquinone (BQ) to capture h<sup>+</sup>, 'OH and 'O<sub>2</sub><sup>-</sup>, respectively, and also N<sub>2</sub> purging was performed to confirm the importance of O<sub>2</sub> during reaction, and obtained data

are shown in Fig. 5a [72]. Obviously, the photocatalytic activity was significantly decreased 333 from 86% to 57%, 56% and 54% after addition of BQ and EDTA, and by N<sub>2</sub> purging, 334 respectively. However, there was only slight change after t-BuOH addition, indicating that 335  $O_2^-$ , h<sup>+</sup> and O<sub>2</sub> were crucial for tetracycline degradation, whereas OH radicals had a minor 336 role. These results were further investigated by ESR analysis, as shown in Fig. 5 (b and c). It 337 was found that pristine  $Bi_2WO_6$  was inactive for production of 'O<sub>2</sub><sup>-</sup> (Fig. 5b), whereas 338 339 significant 'OH signals were observed for Bi<sub>2</sub>WO<sub>6</sub>/CuBi<sub>2</sub>O<sub>4</sub> composite (Fig. 5c). These results have indicated that the photogenerated electrons and holes in the Bi<sub>2</sub>WO<sub>6</sub>/CuBi<sub>2</sub>O<sub>4</sub> 340 341 composite photocatalyst are able to form  $O_2^-$  and OH, whereas the potential of photogenerated electrons in  $Bi_2WO_6$  is not sufficient to produce ' $O_2$ '. It is not surprising after 342 consideration that CB and VB levels of Bi<sub>2</sub>WO<sub>6</sub> and CuBi<sub>2</sub>O<sub>4</sub> are 0.33 V and -0.42 V, and 343 3.11 V and 1.37 V (vs. NHE), respectively. Therefore, only photogenerated electrons in 344 CuBi<sub>2</sub>O<sub>4</sub> could reduce O<sub>2</sub> to produce  $O_2^-$  (E (O<sub>2</sub>/ $O_2^-$ ) = -0.33 V vs. NHE), and similarly only 345 photogenerated holes in  $Bi_2WO_6$  could oxidize  $H_2O$  to generate 'OH (E ( $H_2O$ /'OH) = 2.4 V 346 vs. NHE), as shown in Fig. 5d. These results demonstrate that the electrons could not migrate 347 from CB of CuBi<sub>2</sub>O<sub>4</sub> to CB of Bi<sub>2</sub>WO<sub>6</sub> via type-II heterojunction, but rather the Z-scheme 348 mechanism should be considered (Fig. 5d). 349

The 'OH radicals were also detected by PL of HTA ( $\lambda_{exc.}$  = 315 nm) for C-350 dots@TiO<sub>2</sub>/Bi<sub>2</sub>WO<sub>6</sub> composite photocatalyst, as shown in Fig. 6a [73]. It was found that PL 351 352 intensity of HTA increased with an increase in the irradiation time, indicating that the photocatalyst possessed adequate oxidation potential to generate 'OH radicals (conversion of 353 TA into HTA). Considering that the CB and VB potentials of Bi<sub>2</sub>WO<sub>6</sub> are 0.595 V and 3.2 V, 354 respectively, and -0.21 V and 2.8 V are respective values for TiO<sub>2</sub>, it has been suggested that 355 only photogenerated holes at the VB of Bi<sub>2</sub>WO<sub>6</sub> could produce 'OH, whereas O<sub>2</sub> could be 356 only reduced into  $O_2^-$  on the CB of TiO<sub>2</sub> (Fig. 6b), and thus Z-scheme mechanism was also 357

suggested in this system. Although Z-scheme mechanism is highly recommended for high activity, and obtained data could indirectly confirm it, it should be remembered that  $TiO_2$  is known to be effective to form hydroxyl radicals (slight differences in VB bottom have been reported). Moreover, the generation of hydroxyl radical by both redox pathways should be also considered, i.e., via photogenerated holes (as discussed here), but also due to further reaction of  $O_2^-$  radicals (formed from adsorbed oxygen and photogenerated electrons) with water (via H<sub>2</sub>O<sub>2</sub>).



**Fig. 5.** (a) The photodegradation activity of tetracycline (15 mg L<sup>-1</sup>) over  $Bi_2WO_6/CuBi_2O_4$ photocatalyst in the presence of different scavengers (BQ, EDTA and t-BuOH) or N<sub>2</sub> purging, (b-c) DMPO spin-trapping ESR spectra of: (b)  $Bi_2WO_6$  in methanol dispersion for DMPO-'O<sub>2</sub><sup>-</sup> and (c)  $Bi_2WO_6/CuBi_2O_4$  in aqueous dispersion for DMPO-'OH, and (d) the schematic drawing showing the Z-scheme mechanism of tetracycline degradation on  $Bi_2WO_6/CuBi_2O_4$ . Reproduced from Ref. [72], copyright 2019, with permission from Elsevier.

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Fig. 6. (a) PL spectra of the C-dots@TiO<sub>2</sub>/Bi<sub>2</sub>WO<sub>6</sub> photocatalyst under different duration of solar
 radiation in a basic solution of terephthalic acid, and (b) the proposed mechanism of levofloxacin
 degradation on the direct C-dots@TiO<sub>2</sub>/Bi<sub>2</sub>WO<sub>6</sub>Z-scheme photocatalyst [73].

In contrast to photocatalytic oxidation reactions, photocatalytic reduction reactions 378 (e.g., reduction of organic and inorganic compounds, H<sub>2</sub> generation, CO<sub>2</sub> reduction and 379 organic synthesis) involve the reactions on the surface of the photocatalysts by the 380 photogenerated electrons with a suitable potential for specific reaction, and thus 381 photogenerated holes are usually scavenged during these experiments, e.g., by alcohol 382 addition. Accordingly, the Z-scheme formation could be confirmed by detecting the 383 accumulation of the photogenerated electrons on a specific semiconductor of the Z-scheme 384 photocatalytic system during photoreduction. For example, it was suggested that during 385 overall water splitting over the binary Bi<sub>2</sub>WO<sub>6</sub>/Cu<sub>3</sub>P composite under solar light irradiation 386 only Cu<sub>3</sub>P could cause water reduction, considering the CB and VB levels of Bi<sub>2</sub>WO<sub>6</sub> (0.22 V 387 and 3.02 V, respectively) and Cu<sub>3</sub>P (-0.64 V and 1.02 V, respectively) [74]. These results 388 have suggested that in the case of Bi<sub>2</sub>WO<sub>6</sub>/Cu<sub>3</sub>P photocatalyst, the heterojunction (type-II) 389 (Fig. 7a) could only result in water oxidation, whereas the overall water splitting might be 390 only achieved via Z-scheme mechanism (Fig. 7b). 391





Fig. 7. The schematic drawings of charge carriers' transfer for Bi<sub>2</sub>WO<sub>6</sub>/Cu<sub>3</sub>P photocatalyst in the case
of: (a) type-II heterojunction and (b) Z-scheme photocatalyst. Reproduced from Ref. [74], copyright
2018, with permission from RSC.

397 X-ray photoelectron spectroscopy (XPS) analysis is usually used to investigate the surface properties of materials, such as chemical composition and oxidation states of 398 elements. Recently, XPS analysis has also been applied to confirm the formation of Z-scheme 399 photocatalytic systemes via evaluation of the shift in binding energies, i.e., positive/negative 400 through losing/receiving electrons, respectively, as well as the changes in the oxidation states 401 of elements after irradiation. Accordingly, after the formation of Z-scheme photocatalytic 402 system between two semiconductors with different electronic properties, the electrons are 403 transfered from the semiconductor of higher Fermi level to that with lower one till two Fermi 404 405 levels are aligned to the same level, resulting in the change of binding energy of the specific elements. Furthermore, when the Z-scheme photocatalyst is photoexcited, photogenerated 406 charge carriers' migration takes place at the interface between two semiconductors, which 407 408 might also result in the change of oxidation state of elements (in the case of incomplete charge carriers' consumption). Therefore, XPS could be also used to investigate the pathway 409 of charge transfer in the Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalytic systems. For example, Yuan 410 et al. have confirmed the formation of Bi<sub>2</sub>WO<sub>6</sub>/CuBi<sub>2</sub>O<sub>4</sub> Z-scheme by the shift towards the 411 higher binding energy for Cu2p (Fig. 8a), Bi4f (Fig. 8b) and O1s (Fig. 8c) in comparison with 412 the pristine CuBi<sub>2</sub>O<sub>4</sub> photocatalyst [72]. It was found that the binding energy of Cu2p in the 413 pure CuBi<sub>2</sub>O<sub>4</sub> could be assigned to Cu<sup>2+</sup>, whereas, that in the Bi<sub>2</sub>WO<sub>6</sub>/CuBi<sub>2</sub>O<sub>4</sub> composite 414

415 could be attributed to  $Cu^+$ . It has been proposed that the electrons are transferred from 416  $CuBi_2O_4$  to  $Bi_2WO_6$  after heterojunction formation for creating an electric field between 417  $CuBi_2O_4$  to  $Bi_2WO_6$ . During XPS analysis, both semiconductors are photoexcited, and hence 418 the photogenerated electrons might migrate from  $Bi_2WO_6$  to  $CuBi_2O_4$  under the effect of 419 internal electric field. Accordingly, the formation of  $Bi_2WO_6/CuBi_2O_4$  Z-scheme has been 420 confirmed by XPS analysis and trapping tests, as discussed above (Fig. 5d).



421

- 422 **Fig. 8.** XPS spectra of binary  $Bi_2WO_6/CuBi_2O_4$  (BWO-CBO-10) and bare  $CuBi_2O_4$  (CBO)
- photocatalysts: Cu 2p (a), Bi 4f (b), and O 1s (c). Reproduced from Ref. [72], copyright 2019, with
  permission from Elsevier.

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#### 428 2.1.2. Band structure characterization and corresponding theoretical analysis

It should be pointed out that the Fermi level ( $E_f$ ) positions of the heterojunction components have a significant effect on the bending mode of energy band and charge transfer direction between these components. The flat-band position (close to that of Fermi level equilibrium) could be obtained from Mott-Schottky plots using Eq. (1) [75]:

433  
434 
$$\frac{1}{C^2} = \frac{2}{A^2 e \varepsilon_0 N_A} \left( E - V_{fb} - \frac{k_B T}{e} \right)$$
 Eq. (1)

435 where, C (interfacial capacitance), e (electronic charge),  $N_A$  (carrier concentration),  $\epsilon$ 436 (dielectric constant of the semiconductor),  $\epsilon_0$  (permittivity of free space), A (electrode 437 surface), E (applied potential),  $V_{fb}$  (falt-band potential),  $k_B$  (Boltzmann constant), and T 438 (absolute temperature).

The V<sub>fb</sub> could be estimated from the potential (E) intercept by plotting  $1/C^2 vs$  E. The 439 positive/negative slope of the Mott-Schottky curve of semiconductors indicates n/p-type 440 character. The V<sub>fb</sub> is closer to CB for n-type semiconductors and to VB for p-type ones. 441 Typically, in the heterojunction system, the electrons could migrate from the semiconductor 442 of higher E<sub>f</sub> to that of lower E<sub>f</sub>, resulting in upward and downward band bending (at the side 443 of high E<sub>f</sub> and low E<sub>f</sub>, respectively), forming a new equilibrated Fermi level, causing a shift 444 in the flat-band position. This phenomenon controls the direction of the transfer of the photo-445 446 induced charge carriers (it could be confirmed by XPS analysis). Therefore, the Mott-Schottky measurements might help to prove the Z-scheme formation. For example, the Mott-447 Schottky analysis has indicated that both Bi<sub>2</sub>WO<sub>6</sub> and g-C<sub>3</sub>N<sub>4</sub> are n-type semiconductors (a 448 positive slope in the Mott-Schottky curve, Fig. 9a-b), and the energy bands are shifted 449 downward and upward, respectively (Fig. 9c), due to the transfer of electrons from  $g-C_3N_4$ 450  $(V_{fb} = -1.03 \text{ eV})$  to  $Bi_2WO_6$   $(V_{fb} = 0.36 \text{ eV})$  [76]. Therefore, under irradiation the 451 452 photogenerated electrons should migrate from the Bi<sub>2</sub>WO<sub>6</sub> to g-C<sub>3</sub>N<sub>4</sub>, suggesting the Zscheme mechanism, as shown in Fig. 9d. 453

It is worth to mention that the loading of metal and metal oxide could also be useful to confirm the charge carriers' transfer mechanism in the Z-scheme photocatalytic systems since it is known that metals and metal oxides are selectively deposited on the electron-rich and electron-deficient sites/components (easily confirmed by the microscopic investigations [77]). Unfortunately, there are no reports on selective deposition of metal/metal oxides on Bi<sub>2</sub>WO<sub>6</sub>-based Z-scheme photocatalytic systems.



460 **Fig. 9. (a-b)** Mott-Schottky plots of  $g-C_3N_4$  (a) and  $Bi_2WO_6$  (b), (c) the diagram of the band bending 461 formed in the  $Bi_2WO_6/g-C_3N_4$  hetrojunction, and (d) the schematic drawing for charge carriers' 462 transfer in  $Bi_2WO_6/g-C_3N_4$  Z-scheme. Reproduced from Ref. [76], copyright 2018, with permission 463 from Elsevier.

464

465 Moreover, the theoretical simulations are significantly useful to estimate the 466 mechanism of charge carriers' transfer in heterojunction systems. Usually, the first-principles simulation based on DFT calculation could be performed to evaluate the photogeneration and
transfer of the charge carriers in a specific photocatalytic system by calculating the effective
mass of charge carriers (m<sup>\*</sup>), as shown in Eq. (2) [77,78]:

470

$$m^* = \frac{h}{\frac{d^2 E_k}{dk^2}}$$
Eq. (2)

472

473 where,  $m^*$  (effective mass of  $e^-$  or  $h^+$ ),  $\hbar$  (reduced Plank constant), k (wave vector), and  $E_k$ 474 (energy corresponding to the wave vector).

It is worth noting that the charge carriers' migration (mobility) might be faster with 475 decreasing the effective mass. For example, the theoretical simulation was used to confirm 476 the formation of Bi<sub>2</sub>WO<sub>6</sub>/SnS Z-scheme photocatalytic system [78]. Firstly, the electronic-477 band structure of Bi<sub>2</sub>WO<sub>6</sub> and SnS were estimated by using first-principles simulation based 478 on DFT calculation, as shown in Figs. 10a-b. Thereafter, the effective masses of electrons and 479 480 holes for Bi<sub>2</sub>WO<sub>6</sub> and SnS were calculated. It has been found that the mobility of electrons (0.192) in Bi<sub>2</sub>WO<sub>6</sub> is much higher than that in SnS (0.010), and hence the electrons would 481 transfer from Bi<sub>2</sub>WO<sub>6</sub> to SnS, resulting in Z-scheme mechanism, as shown in Fig. 10c. 482



**Fig. 10. (a-b)** Band structure of SnS (**a**) and Bi<sub>2</sub>WO<sub>6</sub> (**b**), and (**c**) the diagram of charge carriers' transfer in the Bi<sub>2</sub>WO<sub>6</sub>/SnS Z-scheme photocatalyst. Reproduced from Ref. [78], copyright 2019, with permission from Elsevier.

Summarizing, it should be pointed out that despite various methods have been proposed for mechanism confirmation; many of them do not give the direct proof for Zscheme migration of charge carriers. For example, the shift in XPS suggests only the strong interaction between components under dark conditions, which does not unequivocally confirm the direction of charge carriers` migration under irradiation. Therefore, it is proposed that in-situ methods under irradiation would be the most recommended, such as EPR studies.

494 Similarly, the comparison of properties of photocatalysts before and after irradiation might provide some insights on the mechanism of charge carriers' migration, e.g., the change 495 496 of oxidation states of elements (by XPS as shown for other materials [79]). However, it should be pointed out that such changes should not be observed in perfectly stable materials, 497 i.e., without photo-corrosion. The microscopic investigations for the selective deposition of 498 499 metals and/or metal oxides on the surface of photocatalyst components with reductive and oxidative properties, respectively, might be also helpful, but it should be remembered that the 500 presence of such deposits does not unequivocally prove their selective deposition during 501 irradiation since the post-deposition migration towards the most stable component 502 (adsorption-governed) must also be considered, as recently reported for faceted titania [80]. 503

504 Scavenger tests and photoluminescence experiments might provide some insights on 505 the possible direction of charge carriers, but all materials/components must be carefully 506 analyzed (with reference experiments; not only for the most active sample), e.g., for ternary 507 A/B/C photocatalyst, the activity for A, B, C, A/B, A/C, B/C and A/B/C should be compared.

It should be pointed out that though dyes are commonly used as testing molecules because of their convenient and cheap analysis (UV/vis), the sensitization of semiconductor by them should be included in the overall mechanism, which might be quite challenging without action spectrum analysis [81]. Additionally, special care should be taken for vis inactive components, e.g.,  $TiO_2$  and ZnO, since their vis response would be mainly caused by

dye-sensitization mechanism (self-doping could also give vis activity), and thus other 513 compounds (colorless) should be selected for activity testing instead of dyes [82]. Moreover, 514 for reliable comparison of different materials, the same conditions of light absorption should 515 be provided, i.e., at the maximum of photoabsorption (usually at 1-2 g/L of photocatalyst), as 516 pointed by H. Kisch [83]. It is thought, that the following experiments might be useful to 517 prove the Z-scheme mechanism: (i) action spectrum analysis, (ii) irradiation by two sources 518 519 with different energy (to activate one or more components), (iii) light-intensity dependence, (iv) selective redox reactions (e.g., hydrogen generation on the photocatalyst containing of 520 521 only one component able to reduce proton), and (v) in-situ characterization of photocatalyst properties under irradiation, e.g., EPR. 522

Accordingly, it might be concluded that still a lot of work must be done to prove the mechanism of Z-scheme charge carriers` migration for BWO-based materials using novel and powerful characterization tools. Furthermore, the advanced theoretical calculations and modeling are greatly needed for deeper understanding of the mechanism of charge carriers' transfer kinetics in the Z-scheme systems.

## 528 2.2. Applications of Bi<sub>2</sub>WO<sub>6</sub>-based photocatalysts

Based on the above-mentioned properties of BWO-based photocatalysts, including an efficient spatial separation of photogenerated  $e^{-}/h^{+}$  pairs and good redox properties, the efficient solar-energy conversion for wide range applications has already been reported, including environmental purification, solar fuel generation, and photocatalytic organic synthesis, as briefly presented in the following sections.

# 534 2.2.1. Environmental purification

535 With the rapid industrial development and population growth, numerous harmful organic 536 compounds (HOCs) enter the environment every day. These HOCs have negative impacts on humans, animals, plants, and even whole ecosystem. Therefore, efficient, cheap and green methods of HOCs removal must be applied worldwide. Although, physical methods, e.g., adsorption, filtration, coagulation, separation, flotation and air stripping, are quite efficient, the pollutants are not decomposed, but only moved to another form/state (e.g., water/wastewater/air  $\rightarrow$  solid waste; water/wastewater  $\rightarrow$  gas phase). Therefore, other methods based on decomposition of pollutants (preferably complete degradation – mineralization) are highly recommended, such as AOPs/AOTs (as discussed in section 1).

Accordingly, BWO-based photocatalysts have shown to be a perspective material for 544 545 the decomposition of organic pollutants in water/wastewater, including dyes (methylene blue, methyl orange, methyl green, rhodamine B, reactive blue 19, basic blue 41, acid fuchsin, 546 malachite green, auramine-O, crystal violet and procion Red MX-5B), phenolic compounds 547 (phenol, bisphenol, 4-nitrophenol, 2,4-dichlorophenol, 2,4,6-trichlorophenol 548 and pentachlorophenol), pharmaceuticals, personal care products and endocrine disruptors 549 (tetracycline, oxytetracycline, metronidazole, salicylic acid, levofloxacin, norfloxacin, and 550 ciprofloxacin, cephalexin, enrofloxacin, fluoroquinolones, glyphosate, chlorpyrifos, 2-551 mercaptobenzothiazole, 17  $\beta$ -estradiol) [31-34,36-42,44,46,47,49,50-55,57,59,60,63-552 69,72,73,84-104] (see Table 1). 553

555 **Table 1.** Examples of  $Bi_2WO_6$ -based Z-scheme photocatalysts for degradation of 556 environmental pollutants

Photocatalytic system (PS)		Electron	Light	Application	Activity	Ref.
PSI	PSII	mediator	source			
(reduction)	(oxidation)					
g-C <sub>3</sub> N <sub>4</sub>	Bi <sub>2</sub> WO <sub>6</sub>	Ag	500 W-Xe	rhodamine B	$\eta = 92\%;$	[32]
			lamp (full-	degradation	K = 0.013	
			spectrum)		min <sup>-1</sup>	
					(150 min)	
$g-C_3N_4$	Bi <sub>2</sub> WO <sub>6</sub>	Au	300 W-Xe	rhodamine B	$\eta = 88.7\%;$	[36]
			lamp (UV-	degradation	$K = 0.07 \text{ min}^{-1}$	
			cut-off		1	
			filter, $\lambda >$		(30 min)	

			400 nm)			
g-C <sub>3</sub> N <sub>4</sub>	Bi <sub>2</sub> WO <sub>6</sub>	Pt	500 W-Xe	rhodamine B	$\eta = 100\%;$	[38]
-			lamp (UV-	degradation	K = 0.067	
			cut-off	-	min <sup>-1</sup>	
			filter, $\lambda >$		(70 min)	
			420 nm)			
BiOBr	Bi <sub>2</sub> WO <sub>6</sub>	Bi	350 W-Xe	rhodamine B	$\eta = 98.02\%;$	[41]
			lamp (UV-	degradation	K = 0.046	
			cut-off		min <sup>-1</sup>	
			filter, $\lambda >$		(60 min)	
			420 nm)			
g-C <sub>3</sub> N <sub>4</sub>	Bi <sub>2</sub> WO <sub>6</sub>	Zn	300-W Xe	bisphenol A	$\eta = 93\%;$	[44]
			lamp (UV-	degradation	K = 0.021	
			cut-off		min <sup>-1</sup>	
			filter, $\lambda >$		(120 min)	
			400 nm)			
$g-C_3N_4$	Bi <sub>2</sub> WO <sub>6</sub>	PPy	500-W Xe	rhodamine B	$\eta = 98\%;$	[46]
			lamp (UV-	degradation	$K = 0.04 \text{ min}^{-1}$	
			cut-off		1	
			filter, $\lambda >$		(100 min)	
			420 nm)			
AgBr	Bi <sub>2</sub> WO <sub>6</sub>	RGO	350 W-Xe	tetracycline	$\eta = 84\%;$	[50]
			lamp (UV-	degradation	K = 0.052	
			cut-off		min <sup>-1</sup>	
			filter, $\lambda >$		(60 min)	
			420 nm)			
TiO <sub>2</sub>	$Bi_2WO_6$	CNT	350 W-Xe	cephalexin	$\eta=89.7\%$	[54]
			lamp (full-	degradation	(100 min)	
			spectrum)			
$g-C_3N_4$	$Bi_2WO_6$	Mxene	300-W Xe	ciprofloxacin	$\eta = 87.4\%;$	[55]
		$(Ti_3C_2)$	lamp (full-	degradation	K = 0.058	
			spectrum)		min <sup>-1</sup>	
					(70 min)	
$g-C_3N_4$	$Bi_2WO_6$	-	300 W-Xe	17β-estradiol	$\eta = 100\%$ (50)	[63]
			lamp (UV-	degradation	min)	
			cut-off			
			filter, $\lambda >$			
			420 nm)			
ZnCdS	$Bi_2WO_6$	-	1K W Xe	malachite green	$\eta = 94\%;$	[86]
			lamp (full-	degradation	K = 0.0534	
			spectrum)		min <sup>-1</sup>	
-					(50 min)	
$Bi_2WO_6$	$P25-TiO_2$	-	300 W-Xe	fluoroquinolones	$\eta = 80\%$	[90]
			lamp (UV-	degradation	(90 min)	
			cut-off			
			filter, $\lambda >$			
			420 nm)			

For example, Li et al. reported the photodegradation of rhodamine B (RhB) under vis 557 irradiation on the direct Z-scheme photocatalyst Bi<sub>2</sub>Fe<sub>4</sub>O<sub>9</sub>/Bi<sub>2</sub>WO<sub>6</sub> (BFWO, synthesized with 558 different contents of Bi<sub>2</sub>Fe<sub>4</sub>O<sub>9</sub> (BFO)) [58]. In this work, 100 mL of aqueous suspension of 559 photocatalyst (30 mg) and RhB (10 mg L<sup>-1</sup>) was stirred for 30 min in the dark 560 (adsorption/desorption equilibrium), and then irradiated using 300-W Xe lamp and UV cut-561 off filter ( $\lambda > 420$  nm) as a vis irradiation source. It was found that the best composite 562 photocatalyst (BFWO-7) decomposed RhB completely (100%, 0.0380 min<sup>-1</sup>) leading to its 563 mineralization (75.9% total carbon (TOC) removal), which was much higher than that by 564 pristine Bi<sub>2</sub>Fe<sub>4</sub>O<sub>9</sub> (15.2%, 0.0015 min<sup>-1</sup>, TOC=8.6%) and Bi<sub>2</sub>WO<sub>6</sub> (62.5%, 0.0122 min<sup>-1</sup>, 565 TOC=36.9%), respectively, as shown in Fig. 11a. Additionally, the BFWO-7 photocatalyst 566 showed much higher activity (ca. one order in magnitude) than other one-component 567 photocatalysts, such as TiO<sub>2</sub>, CdS, ZnO, and NiO (Fig. 11b), and good photostability during 5 568 cycles (Fig. 11c). Additionally, the effect of coexisting ions (i.e., Na<sup>+</sup>, Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, and 569  $CO_3^{2-}$ ) was also investigated (Fig. 11d). For this purpose, the model inorganic salts (NaCl, 570 NaNO<sub>3</sub>, Na<sub>2</sub>SO<sub>4</sub>, and Na<sub>2</sub>CO<sub>3</sub>) at concentration of 0.1 mol L<sup>-1</sup> were used. It was found that 571 NaCl and NaNO<sub>3</sub> displayed a negligible effect on RhB degradation, probably due to their 572 neutrality. However, a decrease in RhB degradation efficiency might arise from the 573 competition between sodium cation and dye molecule for the limited reactive sites on the 574 photocatalyst surface. The lower degradation efficiency in the presence of NaCl than NaNO<sub>3</sub> 575 576 could be caused by the ability of generation of hydroxyl radical ('OH) by the latter, and thus increasing the degradation of RhB. On the other hand, Na<sub>2</sub>SO<sub>4</sub> and especially Na<sub>2</sub>CO<sub>3</sub> 577 showed negative effects, resulting from an increase in the pH value of the reaction 578 579 environment. The significant inhibition of RhB degradation by Na<sub>2</sub>CO<sub>3</sub> is caused by generation of  $CO_3^{2-}$  and  $HCO_3^{-}$  (the product of Na<sub>2</sub>CO<sub>3</sub> hydrolysis) acting as hole 580 scavengers. 581





**Fig. 11. (a)** Photocatalytic activity of bare  $Bi_2WO_6$  (BWO) and  $Bi_2Fe_4O_9$  (BFO), and binary Bi\_2Fe\_4O\_9/Bi\_2WO\_6 composites (BFWO) prepared with different BFO/BWO mass ratios, **(b)** comparative photocatalytic activity of BFWO-7 and conventional photocatalysts (TiO<sub>2</sub>, ZnO, CdS and NiO), **(c)** reusability experiments of BFWO-7 photocatalyst for RhB (20 mg L<sup>-1</sup>) degradation under vis irradiation, and **(d)** effect of coexisting ions on RhB degradation activity on BFWO-7 under vis irradiation. Reproduced from Ref. [58], copyright 2018, with permission from ACS.

In another study, the all-solid-state Z-scheme g-C<sub>3</sub>N<sub>4</sub>/RGO/Bi<sub>2</sub>WO<sub>6</sub> photocatalyst 590 (prepared with different contents of g-C<sub>3</sub>N<sub>4</sub>) showed to be highly efficient for 591 photodegradation of 2,4,6-tricholorophenol (TCP) under vis irradiation [47]. In this report, 592 250 mL of aqueous suspension of photocatalyst (0.25 g) and TCP (20 mg L<sup>-1</sup>) was stirred for 593 594 30 min in the dark, and then irradiated with 500-W Xe lamp and cut-off filter (420 nm) as a vis irradiation source. As shown in Fig. 12a, the composite photocatalyst (100% g-595 C<sub>3</sub>N<sub>4</sub>/RGO/Bi<sub>2</sub>WO<sub>6</sub>) exhibited much higher photocatalytic activity (98%) than its 596 components (62% by binary g-C<sub>3</sub>N<sub>4</sub>/Bi<sub>2</sub>WO<sub>6</sub> composite, and 58% and 52% for single 597

photocatalysts, i.e.,  $g-C_3N_4$  and  $Bi_2WO_6$ , respectively) after 120-min vis irradiation. Moreover, high stability during five photodegradation cycles was also proven, as shown in Fig. 12b.



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**Fig. 12.** Photocatalytic activity of  $Bi_2WO_6$ ,  $g-C_3N_4$ , binary  $g-C_3N_4/Bi_2WO_6$  composite, and ternary  $g-C_3N_4/RGO/Bi_2WO_6$  composite, prepared with different contents of  $g-C_3N_4$  for TCP degradation (**a**) and photocatalytic degradation cycles of TCP over  $g-C_3N_4/RGO/Bi_2WO_6$  photocatalyst (**b**) under vis irradiation for 120 min. Reproduced from Ref. [47], copyright 2016, with permission from Elsevier.

Interestingly, two-dimensional (2D) Bi<sub>2</sub>WO<sub>6</sub> exhibits higher photocatalytic activity 607 608 than other morphologies, probably due to larger specific surface area and more photocatalytic active sites, greatly reducing the distance of charge transfer and facilitating the carrier 609 migration from the interior to the photocatalyst surface. However, the photocatalytic activity 610 611 of pure 2D Bi<sub>2</sub>WO<sub>6</sub> is low (as clarified in Introduction), and thus 2D/2D materials have attracted much attention. Accordingly, Jiang et al. fabricated 2D/2D SnNb<sub>2</sub>O<sub>6</sub>/Bi<sub>2</sub>WO<sub>6</sub> direct 612 Z-scheme photocatalysts (with different contents of SnNb<sub>2</sub>O<sub>6</sub>) for photodegradation of 613 antibiotic norfloxacin under vis irradiation [105]. In this study, the aqueous suspension of 614 photocatalyst (50 mg) and norfloxacin (10 mg L<sup>-1</sup>) was stirred for 30 min in the dark, and 615

then irradiated using 300-W Xe lamp and UV cut-off filter ( $\lambda > 420$  nm). The binary SnNb<sub>2</sub>O<sub>6</sub>/Bi<sub>2</sub>WO<sub>6</sub> composite (3% SnNb<sub>2</sub>O<sub>6</sub>/Bi<sub>2</sub>WO<sub>6</sub>) exhibited the best activity, which was 9.3 and 2.3 times higher than its components (SnNb<sub>2</sub>O<sub>6</sub> and Bi<sub>2</sub>WO<sub>6</sub>, respectively), as shown in Fig. 13a-b. Moreover, the high stability during four subsequent experiments was also proven (Fig. 13c).



Fig. 13. (a-b) Photocatalytic degradation of norfloxacin over Bi<sub>2</sub>WO<sub>6</sub>, SnNb<sub>2</sub>O<sub>6</sub>, and
SnNb<sub>2</sub>O<sub>6</sub>/Bi<sub>2</sub>WO<sub>6</sub> composites (synthesized with different contents of SnNb<sub>2</sub>O<sub>6</sub>), under vis irradiation
for 60 min (a) with the corresponding pseudo-first-order reaction kinetics (b), and (c) the
photodegradation cycles of norfloxacin over 3% SnNb<sub>2</sub>O<sub>6</sub>/Bi<sub>2</sub>WO<sub>6</sub> Z-scheme. Reproduced from Ref.
[105], copyright 2019, with permission from Elsevier.

Recently, great attention has been paid to the fabrication of metal-organic frameworks (MOFs) as a class of crystalline micro/mesoporous hybrid materials, composed of metal ions or metal clusters interconnected by organic linkers, because of their numerous structures and functions, which results in a new type of photoactive materials for diverse photocatalytic applications. More recently, nickel-based MOFs (Ni-MOFs), particularly 2D Ni-MOFs, have attracted intensive research interest because of their non-toxicity, low cost, different and 643 tailored topological structures, and high stability [65]. Nevertheless, the bare Ni-MOFs showed low photocatalytic activity, because of the rapid recombination of photogenerated 644 charges. The construction of Z-scheme from Ni-MOFs and other semiconductors of suitable 645 energy bandgaps might be an effective strategy to solve this issue. Accordingly, Cheng et al. 646 synthesized the direct Z-scheme of Bi2WO6/Ni-MOF sheets (NiBWO, with different 647 compositions) for photodegradation of methylene blue (MB) dye under vis irradiation [65]. In 648 this work, the aqueous suspension of photocatalyst (0.5 g) and MB (20 mg  $L^{-1}$ ) was 649 magnetically stirred for 30 min in the dark, and then irradiated with 300-W Xe lamp and UV 650 cut-off filter ( $\lambda > 420$  nm). It has been shown that the composite (NiBWO) could get 24- and 651 6.4-times higher activity than its components (Ni-MOF and BWO, respectively; Fig. 14a-b), 652 reaching 91.1% removal of total organic carbon (TOC) during 30 h of irradiation (Fig. 14c). 653 654

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**Fig. 14. (a-b)** Photocatalytic degradation of MB over Bi<sub>2</sub>WO<sub>6</sub>, Ni-MOF, and NiBWO composites (synthesized with different contents of Ni-MOF: 10, 20, 30 wt.%), under vis irradiation for 60 min (a) with the corresponding pseudo-first-order reaction kinetics (b), and (c) the TOC removal within MB degradation over NiBWO (20wt% of Ni-MOF) under vis irradiation [65]. Copyright 2022, with permission from Elsevier.

Summarizing, it is obvious that BWO-based Z-scheme photocatalysts have exhibited 664 remarkable photocatalytic activity towards photodegradation of organic pollutants (dyes, 665 phenolic compounds, pharmaceuticals and pesticide). However, the physicochemical 666 characteristics of BWO-based Z-scheme photocatalyst should be further investigated. 667 Additionally, the photocatalytic degradation mechanism of organic compounds and the 668 combination of outstanding activity with photostability and reusability need further 669 670 clarifications (only limited number of reports dealing with these aspects). Moreover, the design and operative optimization of BWO-based photocatalysis processes should be fully 671

672 investigated, considering the effluent properties to enable the use of BWO-based Z-scheme673 photocatalytic systems at large scale (industrial scale) with high performance and low cost.

Another kind of pollutants, which has been treated by BWO-based photocatalyst, is 674 the group of heavy metals. Although BWO-based Z-scheme photocatalysts have been 675 suggested to be a future perspective for the removal of heavy metals, to date, only Cr removal 676 has been reported [99,106-110]. For instance, Z-scheme Co<sub>3</sub>O<sub>2</sub>/Ag/Bi<sub>2</sub>WO<sub>6</sub> photocatalyst 677 678 was used for photocatalytic reduction of Cr(VI) in an aqueous phase under vis irradiation (300-W xenon lamp with UV cut-off filter,  $\lambda > 420$  nm) [108]. Accordingly, it was found that 679 680 the ternary photocatalyst (Co<sub>3</sub>O<sub>2</sub>/Ag/Bi<sub>2</sub>WO<sub>6</sub>) exhibited the best photocatalytic activity (58.0% (Fig. 15a), 0.0153 min<sup>-1</sup> (Fig. 15b)), which was 1.3 and 2.1 times higher than those of 681 the binary photocatalysts (Ag/Bi<sub>2</sub>WO<sub>6</sub> (43.4%, 0.0102 min<sup>-1</sup>) and (Co<sub>3</sub>O<sub>2</sub>/Bi<sub>2</sub>WO<sub>6</sub> (27.5%, 682  $0.00598 \text{ min}^{-1}$ ), respectively, and 3.4 times higher than that by the single photocatalyst 683  $(Bi_2WO_6 (16.9\%, 0.00328 \text{ min}^{-1}), \text{ during 60 min-vis irradiation, as shown in Fig. 15.$ 684



**Fig. 15.** Photocatalytic activity (**a**) and reaction rate constants (**b**) of  $Bi_2WO_6$ ,  $Co_3O_2/Bi_2WO_6$ , Ag/Bi<sub>2</sub>WO<sub>6</sub>, and  $Co_3O_2/Ag/Bi_2WO_6$  photocatalysts for removal of  $Cr^{6+}$  under vis irradiation. Reproduced from Ref. [108], copyright 2019, with permission from Elsevier.

689 BWO-based photocatalysts have also been proposed for removal of microorganisms 690 from water/wastewater. However, only two studies by Li et al. [78], and Yang et al. [88] have 691 been reported up to date [67]. For example, Li et al. reported photo-inactivation of *E. Coli K*- 692 12 on direct Z-scheme SnS/Bi<sub>2</sub>WO<sub>6</sub> (SB) photocatalysts (with different contents of SnS) using 300-W halogen tungsten lamp (with UV cut-off filter,  $\lambda > 410$  nm) [78]. It was found 693 that there was no noticeable change in the cell survival ratio in the dark condition, 694 demonstrating that "dark" cytotoxicity of the photocatalyst was negligible, whereas under 695 irradiation, bacteria inactivation was significantly enhanced, especially in the presence of SB 696 photocatalyst, as shown in Fig. 16a (Fig. 16b presents the photographs of cultured E. coli 697 698 colonies collected at different time during photocatalytic disinfection process.). Although  $SnS/Bi_2WO_6$  (0.50 SB) photocatalyst has exhibited the highest photocatalytic activity 699 700 (survival ratio = 32%) as compared to its components (survival ratios on SnS and Bi<sub>2</sub>WO<sub>6</sub> equal to 81% 50%), the inactivation rate is quite low, i.e., much lower than that by modified 701 702 titania photocatalysts (in logarithmic scale [111-113]), and hence further improvements in 703 antibacterial performance (and against other microorganisms) are necessary.



Fig. 16. Photocatalytic disinfection effect of SnS, Bi<sub>2</sub>WO<sub>6</sub>, and SnS/Bi<sub>2</sub>WO<sub>6</sub> (0.05 SB) photocatalysts
 under vis irradiation (a) with photos of *E. coli* colonies, cultured by the samples collected at different
 time during photocatalytic disinfection process (b). Reproduced from Ref. [78], copyright 2019, with
 permission from Elsevier.

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710 Another application of BWO-based photocatalysts for environmental purification has been focused on air treatment [114-119]. For example, Hu et al. prepared Z-scheme 2D/2D 711 black phosphorus/monolayer Bi2WO6 nanosheets (with different contents of black 712 phosphorus (BP), coded as x% BP/MBWO, where x was the weight ratios of BP to MBWO) 713 for photocatalytic removal of NO under vis irradiation (300-W Xe lamp and UV cut-off 714 filter) [114]. It was found that Z-scheme BP/Bi<sub>2</sub>WO<sub>6</sub> (12% BP/MBWO) photocatalyst 715 716 possessed the best photocatalytic removal efficiency (61%), which was about 2.3-times higher than that of pristine  $Bi_2WO_6$  (Fig. 17a), and good photostability even after six recycles 717 718 (Fig. 17b).





Fig. 17. Photocatalytic activity of monolayer Bi<sub>2</sub>WO<sub>6</sub> (MBWO) and BP/MBWO (with different BP contents, 3–15%) for photo-removal of NO under vis irradiation (a); and cyclic photocatalytic NO degradation reactions on 12% BP/MBWO (b). Reproduced from Ref. [114], copyright 2019, with permission from Wiley.

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Zhang et al. reported the UV photodegradation of gaseous benzene (as one of VOCs)
over direct Z-scheme Bi<sub>2</sub>WO<sub>6</sub>/palygorskite photocatalyst (prepared with different loadings of
palygorskite (Pa)) [115]. It was found that the Bi<sub>2</sub>WO<sub>6</sub>/Pa (8:2) composite exhibited the

highest photocatalytic activity of 62.7%, which was much better than that by its components,
i.e., 46.0% by Bi<sub>2</sub>WO<sub>6</sub> and 35.8% by Pa.

In summary, though BWO-based photocatalysts have exhibited superior performance in comparison to their components towards photocatalytic removal of indoor air pollutants, this activity is still low, and hence more improvements are necessary for boosting the photocatalytic activity under vis irradiation. Moreover, there are only few reports dealing with photocatalytic removal of indoor air contaminants over BWO-based photocatalysts, and thus more study is highly needed.

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# 737 2.2.2. Solar fuel generation

Among different Z-scheme photocatalytic systems, BWO-based photocatalysts have also 738 739 been successfully used for H<sub>2</sub> generation [32,48,49,55,59,60,62,74,76,114,120,121]. For example, Qiang et al. reported the photocatalytic H<sub>2</sub>-evolution over I-doped Z-scheme 740 Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> photocatalyst (prepared with different contents of iodine) under vis 741 742 irradiation (300-W Xe lamp, UV cut-off filter,  $\lambda > 420$  nm) [62]. In this report, photocatalyst (25 mg) was dispersed into 50 mL aqueous suspension of methanol (10 vol%), and then 743 stirred for 30 min in the dark. Before irradiation, the suspension was bubbled with argon to 744 remove oxygen (an electron scavenger). It was found that the I-doped Z-scheme 745 Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> photocatalyst (S5 sample) showed high stability (Fig. 18c) and best 746  $(186.22 \mu mol g^{-1}h^{-1})$  when compared with non-doped 747 photocatalytic activity  $Bi_2O_2CO_3/Bi_2WO_6$  composite (S10 sample, 40.67 µmol g<sup>-1</sup>h<sup>-1</sup>), pristine  $Bi_2O_2CO_3$  (S9 748 sample, 8.33  $\mu$ mol g<sup>-1</sup>h<sup>-1</sup>) and bare Bi<sub>2</sub>WO<sub>6</sub> (inactive S1), as shown in Fig. 18 a-b. Moreover, 749 750 the addition of platinum (3 wt%) increased the photocatalytic activity of S5 sample by 3.6 times (664.5  $\mu$ mol g<sup>-1</sup>h<sup>-1</sup>), reaching AQE of 14.9% (Fig. 18 a-b). 751



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Fig. 18. (a-b) Photocatalytic activities of: Bi<sub>2</sub>WO<sub>6</sub> (S1), and I-doped Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub>
photocatalysts, prepared with different contents of iodide (S2–S8) (a) and non-doped
Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> (S10), I-doped Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> (S5), and bare Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub> (S9) (b), and (c)
photocatalytic activity of I-doped Bi<sub>2</sub>O<sub>2</sub>CO<sub>3</sub>/Bi<sub>2</sub>WO<sub>6</sub> (S5) during 5-cycles. Reproduced from Ref.
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Another example of direct Z-scheme photocatalyst was presented by Rauf et al. for overall water splitting into H<sub>2</sub> and O<sub>2</sub> under AM 1.5 G simulated solar light (100 mW cm<sup>-2</sup>) on Bi<sub>2</sub>WO<sub>6</sub>/Cu<sub>3</sub>P (prepared with different contents of Bi<sub>2</sub>WO<sub>6</sub>) photocatalyst [74]. The photocatalyst (100 mg) was dispersed into 80 mL of buffer aqueous suspension (0.5 M Na<sub>2</sub>HPO<sub>4</sub>/NaH<sub>2</sub>PO<sub>4</sub>) and then bubbled with N<sub>2</sub>. It was obvious that Bi<sub>2</sub>WO<sub>6</sub> (30%)/Cu<sub>3</sub>P composite exhibited the highest photocatalytic activity (H<sub>2</sub> = 9.3 µmol g<sup>-1</sup>, O<sub>2</sub> = 4.6 µmol g<sup>-1</sup>, H<sub>2</sub>/O<sub>2</sub> ~ 2), when compared with bare Cu<sub>3</sub>P (H<sub>2</sub> = 3.5 µmol g<sup>-1</sup>, O<sub>2</sub> = 1.7 µmol g<sup>-1</sup>) and pristine Bi<sub>2</sub>WO<sub>6</sub> (no evolution of H<sub>2</sub> and O<sub>2</sub>), as shown in Fig. 19a. Furthermore, the Zscheme Bi<sub>2</sub>WO<sub>6</sub> (30%)/Cu<sub>3</sub>P photocatalyst exhibited good stability after 3-cycles photoreactions, as shown in Fig. 19b.



<sup>773</sup> 

**Fig. 19. (a)** Time courses of photocatalytic overall water splitting on  $Bi_2WO_6$ ,  $Cu_3P$ , and  $Bi_2WO_6/Cu_3P$  prepared with different  $Bi_2WO_6$  contents (10–40%) under simulated solar light, and (b) reusability experiments for water splitting over  $Bi_2WO_6$  (30%)/ $Cu_3P$  photocatalysts. Reproduced from Ref. [74], copyright 2018, with permission RSC.

Summarizing, it should be pointed out that though the fabrication of Z-scheme has enhanced the photocatalytic activity of  $Bi_2WO_6$  for  $H_2$  evolution and overall water splitting through the improvements of the charge carriers' separation and redox ability, and protecting  $Bi_2WO_6$  against photocorrosion, the amounts of evolved gases are still low. Therefore, more studies are needed for activity improvements, especially in the absence of noble metal cocatalyst to decrease the operation cost.

Among solar-light-activated Z-scheme photocatalysts, BWO-based Z-scheme 785 heterojunctions have also been fabricated for photocatalytic CO<sub>2</sub> reduction [35,43,48,56,122]. 786 For example, Li et al. used direct Z-scheme g-C<sub>3</sub>N<sub>4</sub>/Bi<sub>2</sub>WO<sub>6</sub> heterojunction (prepared 787 with different contents of g-C<sub>3</sub>N<sub>4</sub>) for photoconversion of CO<sub>2</sub> into CO under vis irradiation 788 (300-W xenon lamp, UV cut-off filter,  $\lambda > 420$  nm) [56]. It was found that the g-789 C<sub>3</sub>N<sub>4</sub>/Bi<sub>2</sub>WO<sub>6</sub> (prepared with addition of 0.1 g g-C<sub>3</sub>N<sub>4</sub>) exhibited the best photocatalytic 790 activity with CO generation rate of 5.19  $\mu$ mol g<sup>-1</sup> h<sup>-1</sup>, which was 22 and 6.4 times higher than 791 those of bare g-C<sub>3</sub>N<sub>4</sub> and Bi<sub>2</sub>WO<sub>6</sub>, respectively, as shown in Fig. 20. It should be pointed out, 792 793 that though some BWO-based Z-scheme photocatalysts showed superior photocatalytic activity for photocatalytic conversion of CO<sub>2</sub> than its components, the selectivity, the mass 794 transfer and thermodynamics of photocatalytic CO<sub>2</sub> reduction, as well as the reaction 795 796 mechanism have not been well investigated yet, and thus further studies are needed.



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**Fig. 20.** Photocatalytic CO<sub>2</sub> reduction into CO on  $g-C_3N_4/Bi_2WO_6$  heterojunctions prepared with different  $g-C_3N_4$  contents, and pristine photocatalysts ( $g-C_3N_4$  and  $Bi_2WO_6$ ) under vis irradiation. Reproduced from Ref. [56], copyright 2015, with permission from RSC.

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Among various Z-scheme photocatalytic systems, BWO-based photocatalysts have already been 805 proposed also for photocatalytic organic synthesis. First and only one report on organic synthesis 806 was shown for the photocatalytic selective conversion of 4-nitroaniline into 4-phenylenediamine 807 808 by Yuan et al. over direct Z-scheme Bi<sub>2</sub>WO<sub>6</sub>/TiO<sub>2</sub> (prepared with different contents of BWO) and Au@Bi<sub>2</sub>WO<sub>6</sub>/TiO<sub>2</sub> under UV-vis irradiation (300-W xenon lamp, 320 nm  $< \lambda < 780$  nm) [120]. 809 For photocatalytic reaction, 40 mL aqueous suspension of 4-nitroaniline (10 mg L<sup>-1</sup>) containing 10 810 811 mg catalyst and 40 mg ammonium formate (as a hole scavenger) was stirred in dark for 2 h before illumination. It was found that the binary Bi<sub>2</sub>WO<sub>6</sub>/TiO<sub>2</sub> photocatalyst (prepared with 10 wt% of 812 BWO) showed much higher photocatalytic activity (72% conversion) than its components, i.e., 813 BWO (no activity) and TiO<sub>2</sub> (40% conversion), as shown in Fig. 21a. Additionally, Au (0.1 wt%) 814 loaded on Z-scheme Bi<sub>2</sub>WO<sub>6</sub>/TiO<sub>2</sub> photocatalyst enhanced the activity to reach 100% conversion 815 816 (Fig. 21b). Similar to other photocatalytic reactions, BWO-based photocatalyst exhibited good 817 photostability during 5 reaction cycles, as presented in Fig. 21c. It should be noted that the BWObased Z-scheme photocatalyst showed superior photocatalytic activity when compared to its 818 components towards photoconversion of 4-nitroaniline into 4-phenylenediamine (which is 819 essential industrial intermediate used for manufacturing different chemical compounds). 820 Furthermore, the decoration with metal (Au) was a suitable strategy for boosting the activity of Z-821 scheme photocatalyst, resulting in complete conversion of 4-nitroaniline. However, the 822 improvements for BWO-based Z-scheme photocatalytic system should be performed for vis 823 824 response photocatalytic conversion processes.

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Fig. 21. (a-b) UV/vis photocatalytic conversion of 4-nitroaniline over: (a) bare TiO<sub>2</sub> (TNS) and BWO, and TiO<sub>2</sub>/BWO composites prepared with different contents of BWO (2-50B–T) and (b) TiO<sub>2</sub>/BWO loaded with different contents of Au (10B-T-0.05 to 0.5Au), and (c) recycling experiments over 10B-T-0.1Au. Reproduced from Ref. [120], copyright 2017, with permission from Wiley.

Summarizing, despite the heterogeneous photocatalysis is an effective method (green, 834 sustainable and eco-friendly) for solar driven organic synthesis (through photocatalytic 835 836 transformation), selectivity problems still limit the practical application. Therefore, it is essential to develop the synthetic methods, solvents (green and eco-friendly), as well as to improve the light 837 absorption ability of the photocatalyst for low-cost photo-organic synthesis with high activity, 838 stability and selectivity under vis irradiation. It should be pointed out that the activity of 839 photocatalyst towards organic synthesis could also be improved by increasing the use of 840 photogenerated charge carries and decreasing the use of scavengers by simultaneously selective 841 photoreduction (by photogenerated electrons) and oxidation (by photogenerated holes) in one 842

system. Moreover, it is significant to develop the photoreactor and in-situ detection methods for
the photocatalytic organic synthesis to transform the synthesis from the lab-scale to large-scale
industrial applications.

## 846 **3. Summary and conclusions**

Environmental pollution and energy shortage are considered as the biggest threats to human 847 and every living organism in the world. Solar photocatalysis as green and clean technology is 848 a promising strategy to resolve these issues. The main requirements, which determine the 849 solar conversion efficiency of photocatalyst, are narrow bandgap (to widen the light-850 absorption range) and large bandgap (to achieve strong redox ability). However, these 851 requirements are difficult to occur simultaneously in one-component photocatalyst. 852 Therefore, great efforts have been paid to develop the efficient photocatalyst working under 853 wide range of solar radiation. It should be pointed out that the Z-scheme photocatalyst 854 outperforms type-II heterojunction materials because of strong redox ability since the 855 photogenerated electrons and holes are more negative and more positive, respectively, and 856 857 additionally spatially separated. Accordingly, the construction of Z-scheme photocatalytic systems is regarded as a promising approach to realize solar-light conversion because of their 858 efficient light harvesting ability, effective separation (spatially separated reductive and 859 oxidative active species) and transfer of photogenerated charges, and strong redox ability. 860

Here, the development of Z-scheme photocatalysts from 1<sup>st</sup> to 3<sup>rd</sup> generation has been presented. Accordingly, it is thought that the direct Z-scheme (3<sup>rd</sup> generation) photocatalysts are the most promising since the charge mediators are not required, and thus (i) the building cost of direct Z-scheme can be largely reduced, (ii) the migration of photogenerated electrons is faster, (iii) the backward reactions are prevented, and (iv) light-shielding effect does not occur. Moreover, the band bending, and electrical field formed in the direct Z-scheme system accelerate the transfer of low-energy electrons from CB of semiconductor with low-Fermi level to recombine with holes from VB of semiconductor with high-Fermi level, leaving thephotogenerated high-energy electrons and holes to drive the photocatalytic reactions.

Recently, great attention has been paid to the fabrication of the direct visible-light-870 activated photocatalysts because of their outstanding activity. Among them, BWO is 871 considered as one of highly promising materials because of its non-toxicity, low cost, and 872 outstanding physicochemical characteristics (i.e., nonlinear dielectric susceptibility, 873 874 ferroelectric piezoelectricity, pyroelectricity, catalytic behavior, modifiable morphology, strong oxidation power, and good photochemical stability). However, the photocatalytic 875 876 activity of pristine BWO is low due to its weak reduction potential, poor light-absorption efficiency, low specific surface area, and rapid recombination of photogenerated charge 877 carriers. Moreover, the impossibility of simultaneous strong redox ability (demanding wide 878 879 bandgap) and broad light-harvesting efficiency (requiring narrow bandgap) is a big challenge for the practical application of pure BWO. Therefore, BWO-based Z-scheme photocatalysts 880 have been constructed to overcome these shortcomings. 881

In this review, the recent developments on Z-scheme photocatalytic systems with 882 special emphasis on the Bi<sub>2</sub>WO<sub>6</sub>-based photocatalysts, including the types, photocatalytic 883 mechanisms and practical applications have been presented. It has been proven that BWO-884 based Z-scheme photocatalysts show high photocatalytic activity towards various 885 applications, including environmental purification, solar energy 886 generation, and 887 photocatalytic synthesis of organic compounds. It should be pointed out that the morphology is also crucial, and thus some advanced structures (e.g., 2D/2D and core-shells) could 888 facilitate the transfer and separation efficiency of photoinduced charge carriers, and hence 889 boosting the activity of BWO-based Z-scheme photocatalysts. It should be pointed out that 890 BWO-based Z-scheme photocatalysts have exhibited high vis activity for photocatalytic 891 reduction of inorganic pollutants in water (heavy metals), air purification (greenhouse gases), 892

degradation of organic compounds (in water and air), water splitting, and synthesis of organic 893 compounds. However, the mechanisms of CO<sub>2</sub> conversion and decomposition of organic 894 compounds need further clarification, i.e., Z-scheme mechanism is one of the solutions, but it 895 should be pointed out that oxygen might be reduced in two-electron transfer mode, and thus 896 high conduction-band bottom position is not required (as reported for pristine BWO [123-897 125]). Moreover, the antibacterial activity under vis irradiation is much lower than that by 898 899 modified titania photocatalysts, and thus needs further improvements. To our knowledge, this is the first review on BWO-based Z-scheme photocatalysts to date, and it is thought it should 900 901 be useful for scientific community to fully explore the potential of BWO-based Z-scheme photocatalysts for environmental remediation, energy conversion and organic synthesis. 902

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