

1 **Visible light responsive heterostructure HTDMA-BiPO₄ modified clays for**
2 **effective diclofenac oxidation: role of interface interactions and basal spacing**

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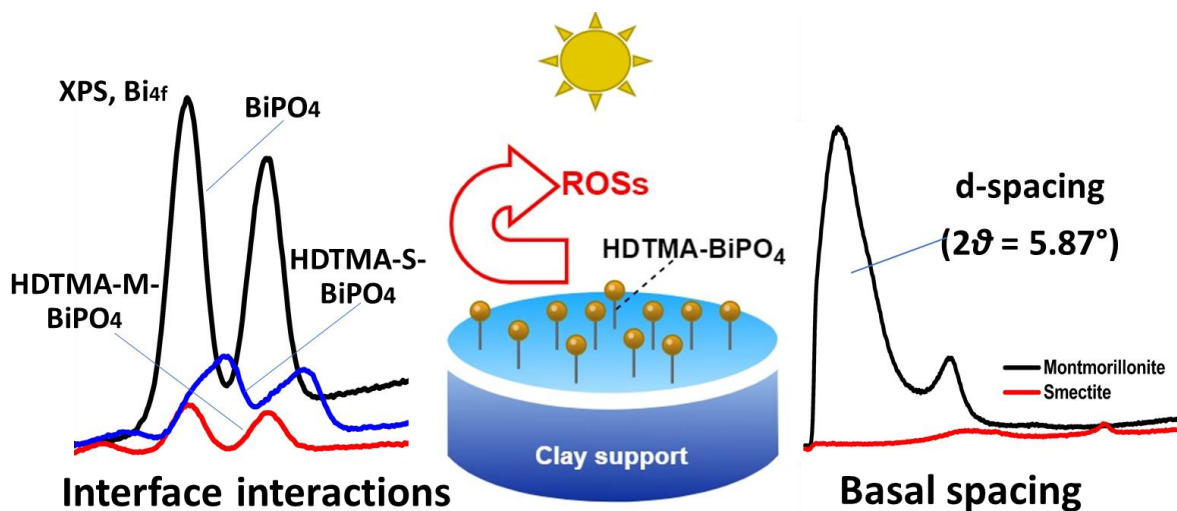
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11 **Graphical abstract**



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21 **Abstract,**

22 In the present study, we investigated comparatively the role of hexadecyltrimethylammonium
23 bromide (HDTMA) in enhancing the adsorption and photocatalytic activity of BiPO₄ coated
24 Montmorillonite (M-BiPO₄), and BiPO₄ coated smectite (S-BiPO₄). Firstly, the direct hybridization
25 of BiPO₄ with clays results in enhanced adsorption and photocatalytic efficiency for the oxidation of
26 diclofenac under solar light compared to bare BiPO₄. Due to the different interlayer spacing of
27 montmorillonite (big) and smectite (small), the hybridization of both BiPO₄ and HDTMA on the
28 surface of clays led to different mechanistic pathways. In terms of montmorillonite, the insertion of
29 BiPO₄ and HDTMA can be realized between the interlayer, while in the case of the smectite, they
30 might be fixed on the external surface. HDTMA reacts better with BiPO₄ set on the external surface
31 to form better heterostructure, as proved by XPS analysis. No peak shifting was observed in Bi_{4f} high
32 resolution in HDTMA-M-BiPO₄, while a strong shift by 1.3 eV along with an obvious change in the
33 shape of peaks was noticed in HDTMA-S-BiPO₄. In addition, the P_{2p} profiles show a positive peak
34 shifting by around 0.7 and 3 eV for HDTMA-M-BiPO₄ and HDTMA-S-BiPO₄, respectively.
35 Regarding the adsorption and photocatalytic tests, at lower concentration (20 ppm), M-BiPO₄ was
36 several times more effective than S-BiPO₄. However, the coating of clays by BiPO₄ and HDTMA
37 showed a different manner, wherein HDTMA-S-BiPO₄ achieved an oxidation rate of around 88%
38 under solar light within 90 min at a concentration of 140 ppm of diclofenac. On the contrary,
39 HDTMA-M-BiPO₄ shows an oxidation rate of only 22 % under the same conditions. It was deduced
40 that the strong surface interactions between HDTMA and BiPO₄ coated on smectite can form a strong
41 interfacial bridge which boosts the visible light response and the separation of photogenerated
42 charges.

43 **Keywords:** Interfacial interaction, HDTMA-BiPO₄, Heterostructures, Solar photocatalysis, Water
44 remediation.

45

46 **Introduction**

47 TiO₂ photocatalysis, since it is discovered in 1972 by Fujishima and Honda [1], has passed by
48 extensive fundamental research and technology pathways towards environmental remediation and
49 energy production [2-6]. The photocatalytic materials fabrication side has taken the most of such
50 scientific and technological research advancements, and many types of photocatalytic materials have
51 been suggested within the last decades [7-12]. Even with such a pool of reported research over more
52 than 30 years, photocatalytic technology is still not convincing enough to be applied extensively in
53 real-world due to many technological and economic issues. Shortly, to convince the scientific
54 community to invest in photocatalytic technology, several issues should be pointed out, including the
55 efficiency of the photocatalytic system compared to existing technologies for a given application, the
56 cost and sustainability of the system. In terms of water purification via the photocatalytic process,
57 many benefits and weaknesses have been raised recently, putting the photocatalytic technology under
58 huge criticisms. Advantages of photocatalysis can involve using solar light as a free energy resource
59 to purify the water on sunny days/regions in a quasi-continuous economic way. However, a correct
60 application should be found to use photocatalysis since this technology is applicable under limited
61 conditions (i.e., less dense wastewaters) because of the use of light to activate the photocatalysts.
62 Based on the reported research, photocatalytic technology has shown low mass transfer (primarily
63 when naked semiconductors are used), generation of by-products, surface deactivation, and
64 complicated or/and expensive synthesis of materials, and so on, as discussed recently by Djellabi et
65 al. [13]. For better technology transfer, several approaches have been suggested to overcome such
66 drawbacks. One of them, the combination of photocatalytic semiconductors with highly adsorptive
67 materials, has been considered as a successful approach. In such a combination, several unique
68 characteristics can be obtained as follows: (i): the coating of small nanoparticles (NPs) of
69 semiconductors on the large particle of adsorbent solves the problem of hard recovery of NPs after
70 the treatment. (ii): The photocatalytic action of NPs on the surface of adsorbent can limit the fast

71 saturation since the oxidative by photogenerated reactive oxygen species (ROSs) reduces the content
72 of pollutants and liberates different adsorptive sites. (iii): Unlike naked semiconductors, the
73 photogenerated by-products can be adsorbed on the surface for further oxidation, avoiding their
74 toxicity in water. (iv): visible light response of semiconductors can be improved when they are
75 hybridized with some adsorptive materials due to the photosensitizing or surface interactions [14, 15].
76 Synergism between the photocatalytic activity and the adsorption effect can be found, wherein a
77 cooperative mechanism, so-called Adsorb and Shuttle process, can take place to remove the pollutants
78 even at higher concentrations [16, 17].

79 Bismuth phosphate (BiPO_4), a relatively newly emerging photocatalyst, which was used for the first
80 time in 2010 [18], has been reported to be very effective towards the oxidation of organic pollutants
81 due to its more positive valance band compared to the common TiO_2 , allowing better oxidation of
82 water molecules into a high yield of hydroxyl radicals. However, BiPO_4 exhibits a large bandgap (in
83 the range of 3.5–4.6 eV), and it requires a strong UV light irradiation to be activated. Loeb *et al.*
84 reported in their critical review [19] that BiPO_4 could be a great alternative photocatalyst for the
85 oxidation and mineralization of organic pollutants. At the same time, creative and yet-to-be-devised
86 approaches to fix its drawbacks are needed to benefit from this promising photocatalyst for its use in
87 photocatalytic wastewater treatment. Several approaches have been already reported to enhance the
88 adsorption ability and visible light response of BiPO_4 such as doping [20-22], heterojunction systems
89 [23-25], oxygen vacancy [26-29], phase junction [30-32], and combination with π -conjugated
90 materials [33-35].

91 In the present work, the enhancement of the ability of BiPO_4 was carried out via the hybridization
92 with two clays (smectite and montmorillonite) to enhance its adsorption capacity and photoactivity
93 under UV and solar light towards the oxidation of diclofenac. To further improve the performance of
94 BiPO_4 -clay based photocatalysts, hexadecyl trimethyl ammonium bromide (HDTMA) was used to
95 modify the clay before the coating of BiPO_4 . The role of HDTMA in enhancing the adsorption and

96 photocatalytic abilities was investigated. Finally, the optimization of BiPO₄ amount coated on the
97 surface HDTMA modified smectite was studied to figure out the ratio of BiPO₄ towards the
98 adsorption and photoactivity.

99 **2. Materials and methods**

100 **2.1. Synthesis of photocatalysts**

101 The raw montmorillonite used in this study was obtained from Roussel deposit in Maghnia (Algeria).
102 Its cationic exchange capacity is 89.30 mmol/100 g. The raw smectite is gray clay derived from Djebel
103 haidoudi near El hamma, and it exhibits a cation exchange capacity of 64.28 meq/100 g. The
104 fabrication of Montmorillonite-BiPO₄ (M- BiPO₄) was carried out by simple precipitation of BiNO₃
105 and NaH₂PO₄ on the surface of Montmorillonite. For this purpose, 1 g of montmorillonite is spread
106 in 20 mL ultrapure water and sonicated for 20 min. Then, under stirring, 5 mL of BiNO₃ (0.3 M) is
107 added dropwise into montmorillonite mixture and left for 60 min under stirring at 60°C. After that, 5
108 mL of NaH₂PO₄ (0.3 M) is added dropwise into the mixture, and stirred for 2 h. The obtained solid
109 was washed and dried at 120°C for a night. HTDMA-Montmorillonite (HTDMA-M) and HTDMA-
110 smectite (HTDMA-S) were prepared by adding a desired amount of HTDMA to water-
111 montmorillonite or smectite, followed by microwave treatment. The sample was recovered and dried
112 at 120 for a night. HTDMA-M-BiPO₄ and HTDMA-S-BiPO₄ were prepared by precipitation of
113 BiNO₃ and NaH₂PO₄ as described above. Bare BiPO₄ was also prepared for the comparison.

114 **2.2. Characterization**

115 Fourier transform infrared spectroscopy analyses (FT-IR) analysis on the as-prepared samples was
116 carried out using a Bruker Vertex 70 spectrophotometer (Bruker, Billerica, MA, US). XRD spectra
117 of samples were recorded on a PANalytical X'PERT-PRO diffractometer with monochromatic CuK α
118 radiation ($\lambda = 1.54056 \text{ \AA}$). X-ray photoelectron spectra (XPS) were recorded on a XPS PHI Quantum

119 instrument. UV–Vis absorption spectra were recorded in the wavelength range of 300–800 nm on
120 Cary 5000 UV-Vis spectrophotometer (Agilent Technology).

121 **2.3. Adsorption and photocatalytic tests**

122 The efficiency of as-prepared photocatalysts was tested against the photocatalytic oxidation of
123 diclofenac under UV ($\lambda_{\text{max}} = 365 \text{ nm}$, 100 W/m^2) and solar light ($35 \text{ W}\cdot\text{m}^{-2}$, ULTRA VITALUX 300
124 W-OSRAM, OSRAM, München, Germany) irradiations. Dark and photolysis experiments were carried
125 out for the purpose of comparison. At different time intervals, samples were collected and filtered
126 using filter paper of $0.45 \mu\text{m}$ diameter. The analysis of diclofenac was performed using a UV-vis
127 spectrophotometer at a wavelength of 276 nm .

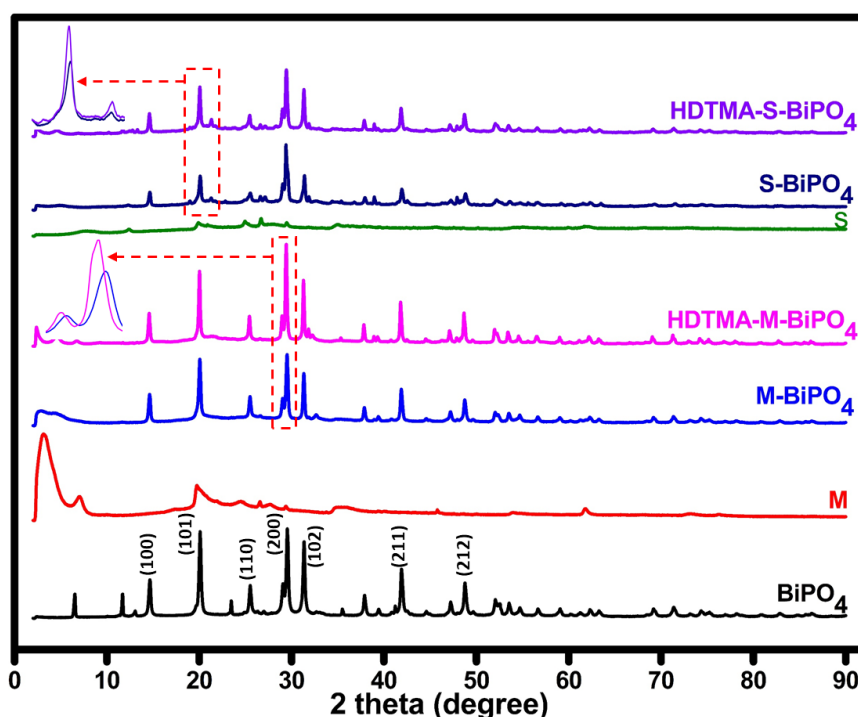
128 **3. Results and discussion**

129 **3.1. Characterization of materials**

130 XRD patterns of BiPO_4 , M, M- BiPO_4 , HDTMA-M- BiPO_4 , S, S- BiPO_4 and HDTMA-M- BiPO_4
131 samples are shown in **Figure 1**. In BiPO_4 and all BiPO_4 modified clays, diffraction peaks were
132 appeared at around 14.7 , 20.1 , 25.5 , 29.5 , 31.3 , 41.9 , and 48.7° which fit with the crystal orientations
133 of the hexagonal phase of BiPO_4 (space group: P3121(152), JCPDS, card no. 15-0766) [36].
134 Montmorillonite's XRD pattern shows a strong d(100) basal spacing reflection at around $2\theta = 5.87^\circ$
135 [37]. However, after the coating of BiPO_4 on the surface of M, this peak was reduced significantly,
136 confirming the formation of BiPO_4 nanoparticles between the montmorillonite layers. The bare
137 smectite pattern does not show a strong basal spacing diffraction peak compared to the
138 montmorillonite sample. The coating of BiPO_4 on HDTMA modified clays resulted in stronger
139 intensities of BiPO_4 diffraction peaks. This may be due to the better crystallization and distribution
140 of BiPO_4 nanoparticles on the surface of HDTMA-clays. The HDTMA organic fragments can interact
141 between the layers of the clay, and also it may react with the components of the clay on the external
142 surface. It may be deduced that the clays with larger basal spacing can receive HDTMA organic
143 fragments compared to those with lower basal spacing. Therefore, the introduction of BiPO_4 on

144 HDTMA modified clays having different basal spacing could have different manners. It is expected
145 that the BiPO_4 could be accumulated in the basal spacing of montmorillonite, while BiPO_4 might be
146 mostly coated on the external surface of smectite because of its low basal spacing.

147 **Figure 2** shows FTIR curves of different samples. In bare BiPO_4 , the large band at $750\text{-}1200\text{ cm}^{-1}$ is
148 assigned to the characteristic asymmetric stretching modes of P–O bonds of PO_4^{3-} [38]. Peaks at the
149 range $520\text{-}600\text{ cm}^{-1}$ are due to the bending vibrations of O–P–O linkage. Characteristic peaks of
150 BiPO_4 were not very obvious in BiPO_4 modified clay samples, which could be overlapped with the
151 original peaks since both montmorillonite and smectite clays have absorption peaks at the same area,
152 i.e., the band at around 1000 cm^{-1} and around 600 cm^{-1} is due to Si–O stretching in-plane and Al–O–
153 Si deformation, respectively [37, 39]. FTIR confirmed the fixation of HTDMA in HDTMA-M- BiPO_4 ,
154 S, S- BiPO_4 , and HDTMA-M- BiPO_4 samples curves, wherein two peaks at around 2936 and 2871 cm^{-1}
155 ¹ were appeared due to the presence of CH_2 and CH_3 , respectively [40].

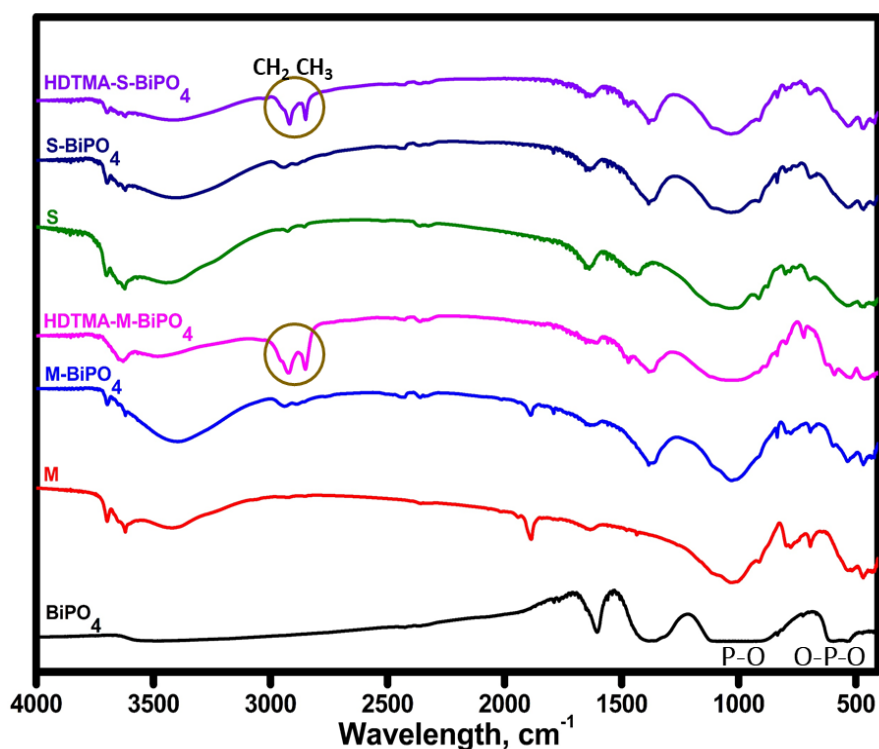


156

157 **Figure 1.** XRD patterns of BiPO_4 , M, M- BiPO_4 , HDTMA-M- BiPO_4 , S, S- BiPO_4 and HDTMA-S-

158

BiPO_4 samples.



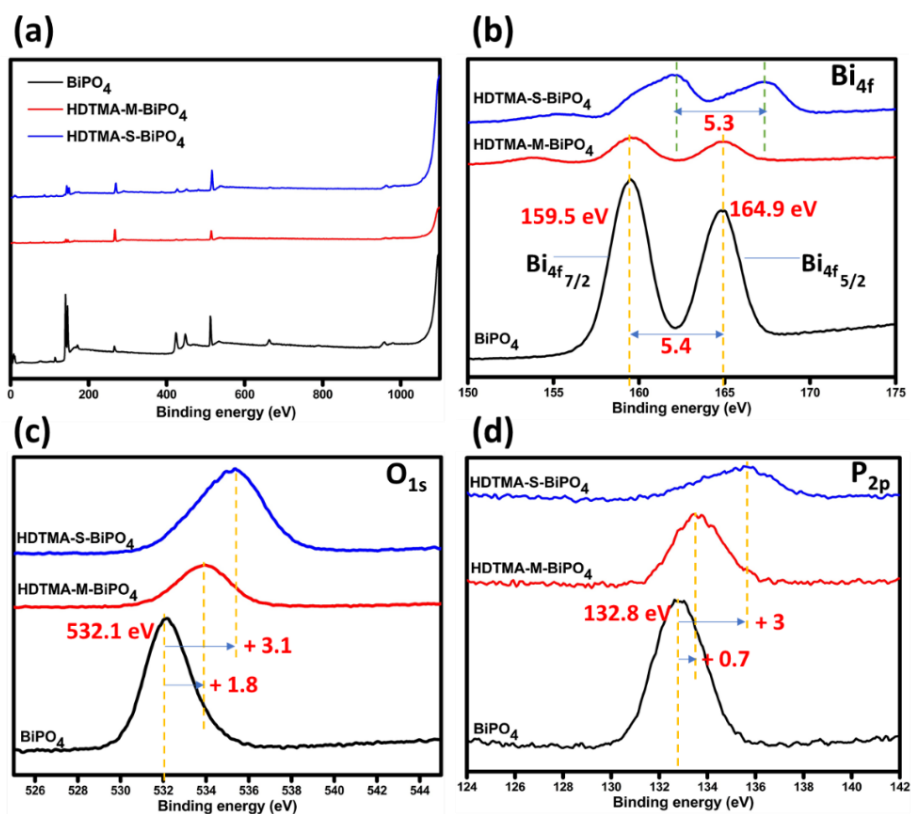
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160 **Figure 2.** FTIR spectra of BiPO₄, M, M-BiPO₄, HDTMA-M-BiPO₄, S, S-BiPO₄ and HDTMA-S-
 161 BiPO₄ samples.

162 The results of XPS analysis carried out on BiPO₄, HDTMA-M-BiPO₄, and HDTMA-S-BiPO₄
 163 samples are shown in **Figure 3**. In terms of Bi_{4f} high resolution (**Figure 3,b**), BiPO₄ profile shows
 164 two intense peaks at 159.5 and 164.9 eV, which are due to Bi_{4f7/2} and Bi_{4f5/2}, respectively, showing
 165 the trivalent oxidation state of Bi species [41]. The distance between Bi_{4f7/2} and Bi_{4f5/2} was calculated
 166 to be 5.3 eV. The position of these peaks was not affected in the case of HDTMA-M-BiPO₄, but a
 167 significant change in the intensity is observed because of the low quantity of BiPO₄. In the case of
 168 HDTMA-S-BiPO₄, a huge change in Bi_{4f} peaks was recorded. The shape of peaks was deformed
 169 along with a significant shift towards higher binding energies. The peaks shifting was about 1.3 eV,
 170 while the distance between Bi_{4f7/2} and Bi_{4f5/2} was reduced to 5.3 eV. Such a change proves the strong
 171 interactions between the atoms of BiPO₄ and the components of HDTMA-S. **Figure 3,c** shows O_{1s}
 172 profiles in BiPO₄, HDTMA-M-BiPO₄ and HDTMA-S-BiPO₄. Bare BiPO₄ shows a single peak at
 173 532.2 eV assigned to crystalline oxygen [42]. O_{1s} spectra in HDTMA-M-BiPO₄ and HDTMA-S-

174 BiPO₄ show a significant shifting towards higher binding energies by around 1.8 and 3.1 eV,
175 respectively, which might be due to the adsorbed oxygen and the chemical interactions between
176 crystalline O²⁻ of BiPO₄ and HDTMA-clay supports. The P_{2p} of PO₄³⁻ state was appeared at a binding
177 energy of 132.8 eV in bare BiPO₄ [43], while a notable shift towards higher energies by 0.7 and 3 eV
178 was detected in HDTMA-M-BiPO₄ and HDTMA-S-BiPO₄ samples, respectively. The observed peaks
179 shifting in HDTMA-M-BiPO₄ and HDTMA-S-BiPO₄ reflects the migration of electron density of
180 BiPO₄ to HDTMA-clay components and the formation of a built-in electric field at the interface. It is
181 important to point out that the XPS peak shifting in HDTMA-S-BiPO₄ is more evident than HDTMA-
182 M-BiPO₄, suggesting that the characteristics of clays play an important role in forming the surface
183 interactions. In fact, from XRD patterns, it can be seen that M samples exhibit a strong interlayer d-
184 spacing ($2\theta = 5.87^\circ$), wherein, BiPO₄ particles can be introduced and fixed inside the interlayers, as
185 confirmed by the XRD pattern of M-BiPO₄ (significant reduction $2\theta = 5.87^\circ$ in peak). However, in
186 terms of the S sample, BiPO₄ might be fixed on the external surface due to the low interlayer spacing,
187 which allows the interaction with the added HDTMA to form more interactions and heterostructure.
188 The optical properties of samples were estimated by UV-Vis diffuse reflectance (Figure 4). Respect
189 to bare BiPO₄, all samples showed a red-shift in the absorption edge. The band gaps of BiPO₄,
190 HDTMA-S-BiPO₄ and HDTMA-M-BiPO₄ were estimated about 3.87, 2.75 and 2.60 eV, respectively.
191 Interestingly, samples with HDTMA showed as well an ulterior peak centrated at around 380 nm
192 which proves the formation of heterojunction system that would boost further the absorption of light
193 and the photosensitization effect.

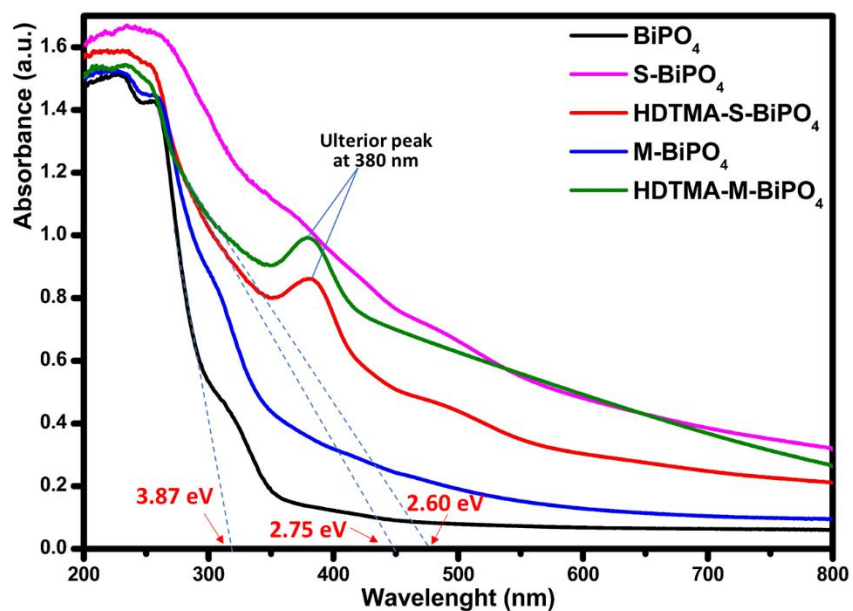
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196 **Figure 3.** (a): XPS survey spectra of BiPO_4 , HDTMA-M- BiPO_4 , and HDTMA-S- BiPO_4 samples.

197 (b): $\text{Bi}4f$ high-resolution profiles. (c): $\text{O}1s$ high-resolution profiles. (d): $\text{P}2p$ high-resolution profiles.

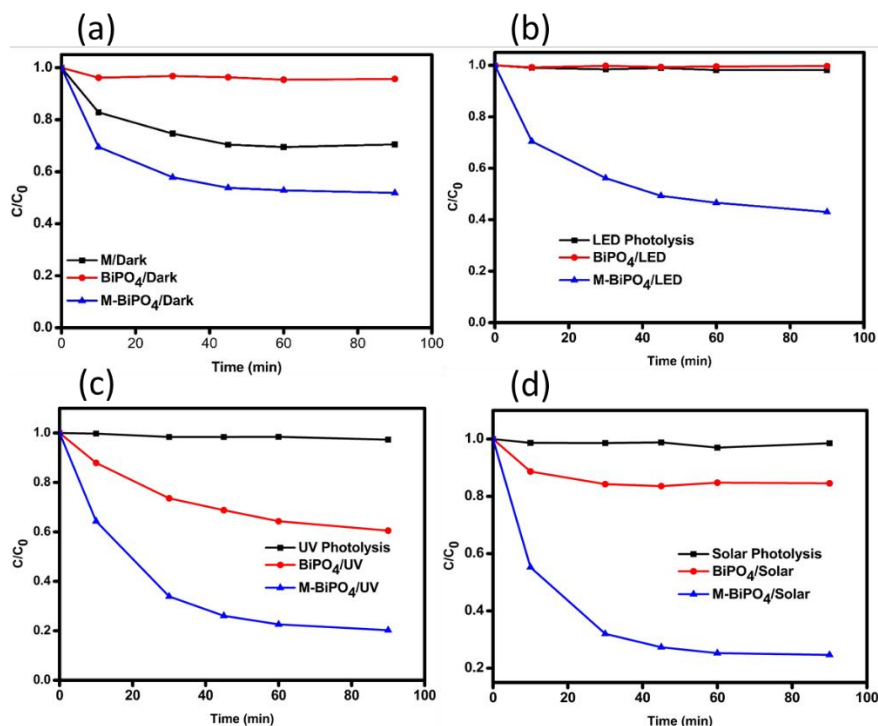


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199 **Figure 4.** UV-Vis diffuse reflectance spectra of bare BiPO_4 , S- BiPO_4 , HDTMA-S- BiPO_4 , M- BiPO_4
 200 and HDTMA-M- BiPO_4 .

201 3.2. Photocatalytic activity

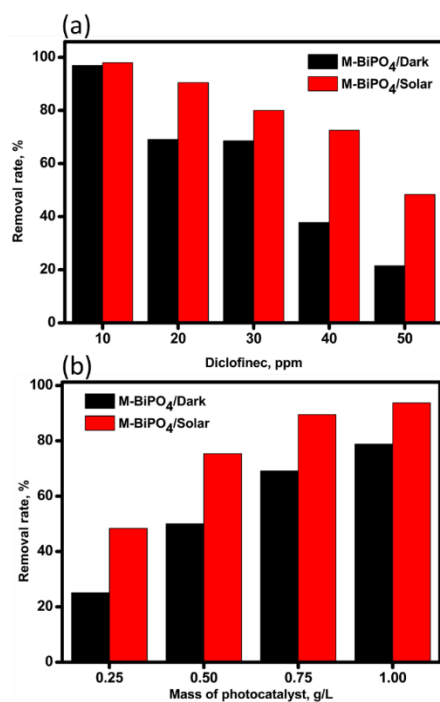
202 The adsorption and photoactivity of bare BiPO₄ and M-BiPO₄ under LED, UV, and solar light were
203 tested comparatively, and the results are shown in **Figure 5**. From **Figure 5.a**, it can be noticed that
204 the coating of BiPO₄ on the M surface led to enhanced adsorption ability compared to bare M and
205 BiPO₄, which could be due to the formation of novel surface adsorption sites. Under LED light
206 irradiation (**Figure 5.b**), no photocatalytic activity was observed using bare BiPO₄, while it can be
207 seen that M- BiPO₄ has a slight photocatalytic activity under light. It is expected that bare BiPO₄ is
208 not photoactive under visible light due to its large bandgap. However, its hybridization with the
209 montmorillonite by in situ precipitation might lead to form chemical interactions with the
210 components, as shown in XPS spectra. Under UV light (**Figure 5. c**), bare BiPO₄ shows a modest
211 photocatalytic activity, wherein less than 40 % diclofenac was removed. On the other hand, M-BiPO₄
212 shows much better photocatalytic oxidation under UV light of diclofenac, more than 80 % within 90
213 min. The reasons behind the enhanced removal of diclofenac under UV is undoubtedly due to the
214 combination of adsorption and photocatalytic activity in the same platform, wherein the adsorb & and
215 shuttle process can accelerate the photooxidation of diclofenac. The adsorptive domain (M) acts to
216 concentrate the diclofenac molecules on the surface nearby the photoactive BiPO₄, resulting in fast
217 oxidation by photogenerated ROSs [16]. Indeed, the bridge between the surface of M and BiPO₄
218 might help to enhance the generation of ROSs due to the better e⁻/h⁺ charges separation. Remarkably,
219 M- BiPO₄ showed excellent photoactivity under solar light, which could be due to the self-doping of
220 BiPO₄ or structure modification during the precipitation of BiPO₄ on the surface of M. Bonds between
221 BiPO₄ and M might lead to decrease the bandgap and promotes the charge generation and separation.



222

223 **Figure 5.** (a): Adsorption of diclofenac by bare M, BiPO₄, and M- BiPO₄, (b): photolysis and
 224 photocatalytic oxidation of diclofenac under LED irradiation by BiPO₄ and M- BiPO₄, (c): photolysis
 225 and photocatalytic oxidation of diclofenac under UV irradiation by BiPO₄ and M- BiPO₄, (d):
 226 photolysis and photocatalytic oxidation of diclofenac under solar irradiation by BiPO₄ and M- BiPO₄.
 227 [Photocatalyst]: 0.5 g/L, [Diclofenac]: 20 ppm.

228 To further evaluate the adsorption/photocatalytic synergism in M-BiPO₄ photocatalyst, adsorption
 229 and solar photocatalytic experiments were carried out with different diclofenac concentrations and
 230 different M-BiPO₄ masses. As depicted in **Figure 6.a**, at lower concentrations, the adsorption process
 231 is predominant compared to the photocatalytic oxidation. The more the diclofenac concentration is,
 232 the more photocatalytic oxidation is evident. At concentrations of 40 and 50 ppm, the removal of
 233 diclofenac by photocatalytic effect is pronounced. The behavior of the M-BiPO₄ at different masses
 234 (**Figure 6.b**) shows that the more the mass of photocatalyst is, the more adsorption of diclofenac is
 235 obtained. It is important to mention that the photocatalytic effect, in such a type of materials, has
 236 another advantage compared to simple adsorption, which is the self-cleaning of the surface for further
 237 adsorption, which in turn retards the fast saturation of the material.

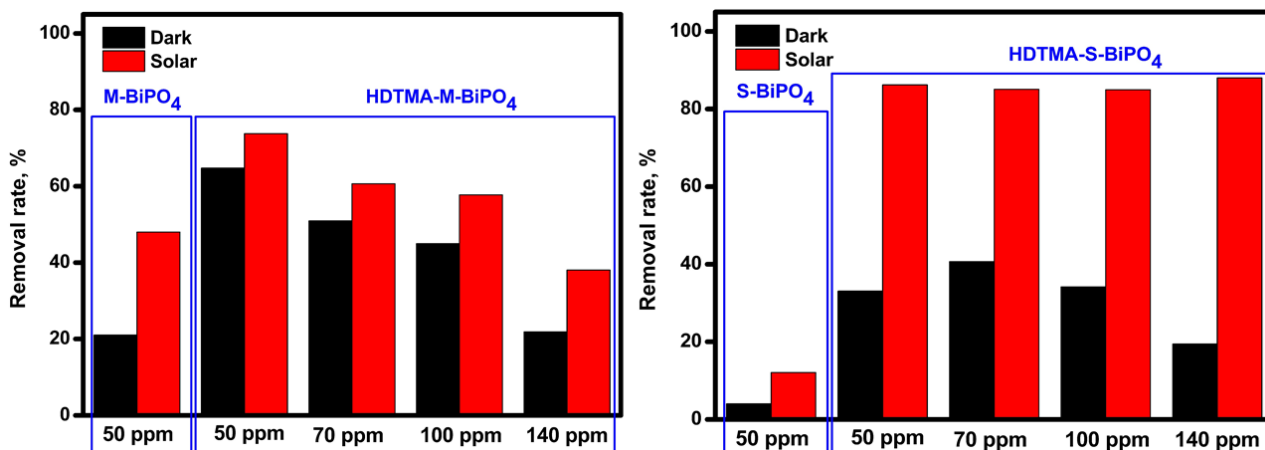


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239 **Figure 6.** (a): Effect of diclofenac concentration on the adsorption and photocatalytic activity under
 240 solar light. [Photocatalyst]: 0.75 g/L, contact time 90 min. (b): Effect of photocatalyst mass on the
 241 removal of diclofenac by adsorption and photocatalytic activity under solar light, [diclofenac]: 20
 242 ppm, contact time 90 min.

243 To investigate the role of HDTMA and the type of clay support, HDTMA-M-BiPO₄ and HDTMA-
 244 S-BiPO₄ were fabricated, and their photocatalytic activities were evaluated at different diclofenac
 245 concentrations ranging from 50 ppm to 140 ppm (**Figure 7**). Firstly, the photocatalytic tests showed
 246 that M-BiPO₄ exhibits higher photocatalytic activity under solar light compared to S-BiPO₄ at a
 247 concentration of 50 ppm. In this case, M-BiPO₄ might have better porosity and photoactivity as
 248 compared to the S-BiPO₄ sample. The fixation of BiPO₄ on the montmorillonite, which has an
 249 excellent basal spacing, allows a better distribution of BiPO₄ particles. It can be seen in XRD patterns
 250 (**Figure 1**) that the peaks in M-BiPO₄ are more intense than those in S-BiPO₄. In addition, since M-
 251 BiPO₄ has better adsorption ability, a higher photoactivity is expected by adsorb and shuttle means
 252 [44, 45]. However, the introduction of HDTMA to the surface of montmorillonite and smectite before
 253 the coating of BiPO₄ has changed the manner completely. In terms of HDTMA-M-BiPO₄, the

254 adsorption ability of the composite was significantly enhanced, leading to faster removal of
255 diclofenac. It can also be seen that the adsorption is predominant in the photocatalytic action in
256 HDTMA-M-BiPO₄ system, especially at a concentration ranging from 50 to 100 ppm. The removal
257 rates decrease as a function of diclofenac concentration. It might be deduced the enhancement in the
258 effectiveness of HDTMA-M-BiPO₄ is more due to the adsorb and shuttle process that can be appeared
259 in the platform of this material, which improves the concentration of diclofenac molecules on the
260 surface near the photoactive BiPO₄. On top of that, the photocatalytic activity of BiPO₄ fixed on the
261 surface of montmorillonite might also be improved through chemical interactions with HDTMA. In
262 the case of HDTMA-S-BiPO₄, another surprising manner was found. HDTMA has notably enhanced
263 the adsorption ability of the composite, but it has a major role for photocatalytic activity under solar
264 light. The removal of diclofenac under solar light was high even at higher concentrations, by
265 photocatalytic means as a predominant action, unlike HDTMA-S-BiPO₄. Based on these results and
266 XPS analysis, it might be deduced that the photocatalytic activity in HDTMA-S-BiPO₄ under solar
267 light is assigned to the strong interface interactions between BiPO₄ and HDTMA, forming a
268 heterostructure platform. A better visible light response and an enhanced photoproduced charges
269 separation can be found in the HDTMA-S-BiPO₄ surface because of the interface junction. As a
270 result, a higher yield of reactive oxygen species can be produced in the medium. BiPO₄ is an excellent
271 producer of hydroxyl radicals, compared to common photocatalysts (i.e., TiO₂), because of its more
272 positive valance band. However, this photocatalyst requires surface modification to decrease its
273 bandgap and improve the separation of e⁻/h⁺ charge carriers. Interestingly, the hybridization of BiPO₄
274 and HDTMA in our conditions had solved perfectly such structure drawbacks.



275

276 **Figure 7.** Adsorption and photocatalytic oxidation of diclofenac by BiPO₄ modified clays and BiPO₄
 277 modified HDTMA-clays, at different diclofenac concentrations. [Photocatalyst]: 0.375 g/L, contact
 278 time 90 min.

279 4. Conclusions

280 The results of this investigation show the significant role of the co-hybridization of BiPO₄ and
 281 HDTMA on the surface of clays, namely, montmorillonite and smectite. In the first study, it was
 282 found out that the coating of BiPO₄ on the surface of montmorillonite leads enhanced adsorption and
 283 photocatalytic activity for the oxidation of diclofenac under LED, UV, and solar light irradiations.
 284 Compared to M- BiPO₄, S-BiPO₄ was less efficient in adsorption and photocatalytic activity under
 285 solar light. Such a difference in the effectiveness might be due to the different basal spacing of the
 286 original clay material, having the montmorillonite an extra-large basal spacing compared to smectite.
 287 The co-modification of clays by BiPO₄ and HDTMA has shown interesting results in terms of
 288 interface interactions. HDTMA in HDTMA-M-BiPO₄ had a positive effect of improving further the
 289 adsorption ability and the photoactivity, wherein diclofenac was removed mostly by physical
 290 adsorption. However, in HDTMA-S-BiPO₄ composite, HDTMA has enhanced the adsorption ability
 291 from one side. Still, mostly HDTMA has improved the photoactivity of BiPO₄ dramatically by
 292 forming an interfacial bridge, confirmed by XPS analysis, which leads to excellent photoexcitation

293 and separation of charges carries. This study showed promising results in enhancing the adsorption
294 and photoactivity abilities of clay-BiPO₄ by sustainable and straightforward surface modification.

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