

A Sustainable Natural Clam Shell Derived Photocatalyst for the Effective Adsorption and Photodegradation of Organic Dyes

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2 effective adsorption and photodegradation of organic dyes

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Abstract

38 In response to the increasing desire for modern industries to be both green and
39 sustainable, there has been increasing research focus on the reutilization of natural
40 waste materials to effectively remove and degrade toxic wastewater effluents. One
41 interesting food industry waste product is clam shells. Here a new photocatalytic
42 nanomaterial derived from marine clam shells was successfully prepared and
43 characterized. Thereafter the material was applied for the removal of two target dyes
44 from aqueous solution, where the effect of both catalyst dose and initial dye
45 concentration on adsorption and photocatalysis properties was investigated. The
46 maximum adsorption capacities of methylene blue (100 mg/L) and Congo red (500
47 mg/L) were 123.45 mg/g and 679.91 mg/g, respectively, where adsorption followed
48 pseudo second order kinetics predominantly via a chemical adsorption process. The
49 photodegradation removal efficiencies of the two dye solutions under visible light
50 irradiation were 99.6% and 83.3% for MB and CR respectively. These results
51 demonstrated that a clam shell catalyst also exhibited excellent degradation
52 performance in a mixed dye solution with strong degradation capability and low cost,
53 making the material a good candidate for practical field remediation of dye
54 contaminated wastewater.

55

56 **Keywords:**

57 Clam shell; Calcination; Congo red; Methylene blue; Photocatalysis

58

59 1. Introduction

60 While water is essential for all life, the rapid growth in modern human
61 civilizations globally, concurrent with vigorous and rapid industrialization has
62 resulted in significant pollution of water resources through the discharge of waste
63 materials, especially by the dye industry. Water resource contamination by dye
64 wastewaters is becoming increasingly serious (Chequer et al., 2009). Methylene blue
65 (MB), Congo red (CR) and many other organic dyes are widely and routinely used in
66 large quantities by various dye industries. These dye molecules have a complex
67 molecular structure containing many diverse functional groups such as amino,
68 hydroxyl and the benzene ring (Balakrishnan et al., 2016), which while giving the dye
69 strong coloring ability also result in strong toxicity and poor biodegradability (Chen et
70 al., 2018). These latter features are of concern because the efflux of large
71 concentrations of dyes into the natural environment may thus lead to environmental
72 harm. Hence, development of new materials to treat dye wastewater is urgently
73 required to mitigate environmental pollution. While many non-destructive methods,
74 such as adsorption, membrane filtration, and precipitation, have previously been
75 successfully employed to remove dye from wastewater, the disadvantage of non-
76 destructive methods are that do they nor destroy the dye but merely concentrate the
77 dye into either sludge or silt, which still needs further treatment to remove the hazard
78 (Tsai et al., 2020). In comparison, destructive methods, can directly decompose large
79 dyes into non-toxic smaller molecules, can be directly discharged into the
80 environment. Therefore, of the destructive methods, photocatalytic reduction has
81 emerged as a popular destructive method. For example, bismuth oxychloride
82 nanomaterials were synthesised via a hydrothermal method, and used for the
83 degradation rate of Congo red with a 72% photocatalytic removal efficiency (Ismail et
84 al., 2019). A $\text{Fe}_2\text{O}_3\text{-CeO}_2$ nanocomposite prepared by a precipitation and improved sol
85 gel self-ignition method, exhibited a 96% photodegradation removal efficiency for CR
86 (Aboutaleb et al., 2019). Ma et al (2017) also successfully synthesized a MgZnCr-
87 TiO_2 nanocomposite photocatalyst by co-precipitation, which removed 99.8% of CR

88 after 180 min. However, the disadvantage of these artificially synthesized catalysts
89 was that the preparation process was complex and, also required expensive and toxic
90 chemical reagents. The application of nanomaterials for photocatalysis is highly
91 desirable due the nanomaterial's high surface areas and semiconductor heterogeneous
92 structures, which significantly improves dye degradation efficiency. Although many
93 nanoparticles have been reported for effective degradation, there are only a few
94 studies which have considered the preparation of photocatalysts from waste materials.
95 Shariffuddin et al (2018) investigated the feasibility of using CaO derived from cockle
96 shells as catalyst for MB removal and demonstrated good degradation. Likewise, E-
97 CaO nanoparticles derived from waste eggshell were successfully prepared by
98 calcination method for efficient photodegradation of methylene blue (MB) and
99 toluidine blue (TB) (Sree et al., 2020), where preparation of E-CaO was both eco-
100 friendly and sustainable. These two studies clearly indicated that recycling wastes as
101 nanomaterial remediating agents was certainly feasible. Hence, it is expected, that in
102 the ongoing search for more efficient and sustainable degradation of dyes, natural
103 photocatalysts derived from waste biomass waste will provide an economical,
104 sustainable and environmentally friendly method.

105 Clams are widely distributed in both the North and South Seas of China, where
106 they grow rapidly, with a short breeding cycle and exhibit strong adaptability, thus
107 making them an excellent shellfish suitable for artificial high-density cultivation.
108 Clam shell is a significant by-product of aquaculture, where not being fully utilized
109 beneficially, results in large numbers of discarded clam shells being accumulated.
110 These discarded clam shells contain remnants of meat and other organic residuals,
111 which will rot potentially producing toxins after long-term exposure to the
112 environment, which has negative impact on aquaculture, and also causes serious
113 environmental pollution. Therefore, mishandling of by-products from aquaculture has

114 previously led to environmental crisis due to the unwanted release of hazardous
115 substances (Penarodriguez et al., 2010). Thus, an urgent waste issue to be resolved is
116 how effectively and beneficially utilize waste clam shells. Previously oyster shells
117 have been used to effectively adsorb H₂S from wastewater, with a maximum
118 adsorption capacity of 12 mg g⁻¹ (Asaoka et al., 2009). Through calcination and
119 hydration, waste oyster shell has also been pretreated to increase its specific surface
120 area and pore volume, for the removal of SO₂/NO_x (Jung et al., 2007). Oyster shell
121 has also been used as an adsorption and filtration medium for phosphorus removal,
122 where its use decreased eutrophication in wastewater (Park et al., 2008). have used
123 Calcined mussel shells have been used to remove mercury (Hg) from water with good
124 efficiency, reaching 90% in 55 min (Penarodriguez et al., 2010). A mixture of mussel
125 shell calcined ash, sewage sludge and wood ash were used successfully as an
126 adsorbent for several metal ions, with removal efficiencies for Hg, arsenic (As) and
127 chromium (Cr) of is 98–99, 90–96, and 32%, respectively (Secoreigosa et al., 2014).
128 Many different natural materials have also been used to specifically remove
129 anthropogenic dyes. The residual biomass of *Spirulina platensis* removed up to 82.6%
130 of CR (Nautiyal et al., 2016), while carbonized kelp biochar performed even better,
131 removing up to 94% of MB (Zhou et al., 2018). These reports indicate that waste
132 utilization of marine aquaculture by-products has been an important issue. Even today,
133 increasing attention has been directed towards better methods for the utilization of
134 marine aquaculture by-products, as many studies have stressed the importance and
135 necessity of global waste utilization. Thus, there seems to be great potential for the

136 utilization of clam shell for cost-effective environmental pollution control. However,
137 to the best of our knowledge, the potential for clam shell to act as a natural
138 photocatalyst for dye degradation has not been previously studied, but this seems
139 likely given the previous successes outlined above.

140 Thus, in this work, a natural photocatalyst based on calcium oxide derived from
141 clam shell was synthesised via a high temperature calcination method and its
142 physiochemical properties fully characterized by SEM, TEM, BET, XRD, XPS, UV-
143 vis and TGA. Thereafter, the as-prepared photocatalysts were evaluated for their
144 ability to adsorb and photocatalytically degrade both cationic (MB) and anionic (CR)
145 dyes, where the mechanism of adsorption and photocatalytic degradation was
146 explored.

147

148 **2. Materials and Methods**

149 *2.1. Materials*

150 Clam shells were collected from a local market in Zhoushan, Zhejiang Province.
151 Methylene blue (MB), Congo red (CR), sodium hydroxide, hydrochloric acid, p-
152 benzoquinone, isopropanol, hydrogen peroxide and ammonium oxalate were all
153 purchased from Shanghai Guoyao Chemical Reagent Co., Ltd. All chemicals used in
154 this work were analytical grade and were used directly as received without further
155 purification.

156 *2.2. Preparation of clam shell catalyst*

157 Clam shells were initially washed with tap water to remove any obvious

158 contamination from the surface of the clam shell. Thereafter, clam shells were
159 immersed in a solution of 1 M sodium hydroxide and heated to 85 °C in a water bath
160 remove any residual meat from the surface of the clam shells, then, the samples were
161 rinsed with fresh water (pH = 7), and evaporated until dryness at 80 °C on a stove.
162 The clam shells were soaked in 1 M hydrochloric acid solution in a clam pot
163 overnight, washed with deionized water several times and dried. Finally, the clam
164 shell was calcined in a muffle furnace for 2 h at 800, 900, and 1000 °C respectively.
165 After calcination, the white powder obtained was ground, sieved (100 mesh) and
166 stored in a desiccator for later use.

167 *2.3. Characterization*

168 Specific surface area and pore size distribution of samples, was calculated using
169 the BET method from N₂ adsorption/desorption isotherms obtained using a static
170 volumetric adsorption analyzer (Micromeritics ASAP 2010, Shanghai, China).
171 Morphology and microstructure of the as-prepared samples was determined via a
172 scanning electron microscope (SEM; S4800, Hitachi, Japan) and high-resolution
173 transmission electron microscope (HR-TEM; Joel-2100, JEOL, Japan). Crystal
174 structures were analyzed by X-ray diffraction (XRD, RIGAKU miniflex/600, Japan)
175 in the range between 20 ° – 80 °. Thermogravimetric analysis was conducted on a
176 TGA Q5000 (American TA instrument Q Series) to better understand phase
177 transformations during calcination. Optical properties of samples, were examined
178 using a Cary 500 UV-vis NIR spectrophotometer in the range 200 – 800 nm. The
179 chemical states of elements at the near surface of samples were also studied using X-

180 ray photoelectron spectroscopy (XPS, Phi 5000C ESCA system, USA).

181

182 *2.4. Photocatalytic experiment*

183 As-prepared photocatalyst (20 mg) was added into an aqueous solution of either
184 MB or CR (40 mL) and the solution was magnetically stirred in the dark. for 2 hours
185 for the reaction to achieve adsorption equilibrium. During this time aliquots of the
186 supernatant solution were removed at regular intervals (i.e., 0, 5, 10, 20, 30, 60, 90
187 and 120 min) and analyzed for the residual dye concentration. To assess adsorption
188 performance, residual dye concentration were determined photometrically.

189 After 2 h dark reaction, a light source (xenon lamp, HSX-F300) was used to
190 initiate photocatalytic experiments. The vertical distance between the light source and
191 the reactor was 10 cm, and supernatant samples were again taken at regularly intervals
192 to assess photocatalytic performance. In the photocatalytic system, a cooling reflux
193 device was also connected to the reaction vessel to eliminate the influence of heat
194 generated by light on the experiment.

195 The effects of different dye concentrations (MB: 50, 75, and 100 ppm; CR: 400,
196 450, and 500 ppm), and different catalyst doses (20, 40, 60, and 80 mg) were also
197 explored.

198 *2.5. Determination of dye concentration*

199 After the reaction single dye solution samples were initially centrifuged at 4000 r
200 min⁻¹ for 10 min, and the absorbance of the supernatant subsequently measured using
201 an UV-vis spectrophotometer. The wavelengths of absorbance maxima for the two

202 dyes were λ_{MB} : 664, and λ_{CR} : 497 nm. The removal efficiency (R%) of adsorbed dye
 203 was calculated by Eq (1), where the amount of dye adsorbed on the catalyst q_e (mg/g)
 204 was calculated by Eq (2).

$$205 \quad R(\%) = \frac{C_0 - C_e}{C_0} \times 100\% \quad (1)$$

$$206 \quad q_e = \frac{(C_0 - C_e)V}{m} \quad (2)$$

$$207 \quad -\ln\left(\frac{C_t}{C_0}\right) = kt \quad (3)$$

208

209 where C_0 and C_e (mg L^{-1}) were respectively, the initial and equilibrium dye
 210 concentrations in solution, m (g) was the mass of catalyst and V (L) was the volume
 211 of dye solution. In order to study reaction kinetics, the rate constant k was obtained by
 212 curve fitting data to Eq (3) t , where C_0 is the initial dye concentration after dark
 213 reaction and C_t is the dye concentration at light irradiation time t .

214

215

2.6. Adsorption kinetics under dark reaction

216 Adsorption kinetics of the single dye system was further studied by adding catalyst
 217 to the dye solution under dark conditions with constant magnetic stirring and measuring
 218 the residual dye concentration in the solution at fixed time periods. The kinetic data so
 219 obtained was fit to both pseudo first order (PFO Eq 4) and second order (PSO Eq 5)
 220 kinetics to evaluate the kinetics of adsorption.

$$221 \quad \ln(q_e - q_t) = \ln q_e - k_1 t \quad (4)$$

$$222 \quad \frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \frac{t}{q_e} \quad (5)$$

223 where q_t (mg g^{-1}) is the dye adsorption capacity at time (t); and k_1 (min^{-1}) and k_2 (g
224 $\text{mg}^{-1} \text{min}^{-1}$) are the rate constants for PFO and PSO models, respectively.

225

226 **3. Results and discussion**

227 *3.1. Characterization of catalyst*

228 *3.1.1. Morphology*

229 The changes in clam shell powder morphology following calcination at three
230 different temperatures were monitored using SEM and TEM (Fig. 1). Before
231 calcination, clam shell powder had a spongy porous structure (Fig. a,b), due to
232 presence of organic matter and a calcium carbonate skeleton after soaking in acid.
233 However, after calcination at 800 °C, the clam shell surface became fragmented and
234 bulky, due to the consumption of organic matter and growth of crystal particles (Fig.
235 1c,d). As calcination temperature further increased (900 and 1000 °C), clam shell
236 powder morphology and surface structure became more compact. The increase in
237 particle size was attributed particle sintering, resulting in Ostwald ripening, which
238 causes growth of larger particles at the expense of smaller ones. This was consistent
239 with previous results which had shown an increase in average surface grain size with
240 calcination temperature (Peralta et al., 2017).

241 More detailed morphological characteristics of clam shell calcined at 1000 °C
242 were obtained by HR-TEM (Fig. 1 i-k), and showed that after high temperature
243 calcination the clam shell catalyst (CSC) had a layered structure where the distance
244 between two adjacent lattices at the interface was 0.286 nm, which was consistent
245 with XRD results (section 3.1.3) corresponding to the CaO (111) crystal face. In

246 addition, the observed heterojunctions between dislocations in the crystal structure
247 may be due to the rearrangement of additional atomic layers produced by the presence
248 of trace metals in the CSC during calcination (Wei et al., 2018). This may enable
249 greater electron migration to the conductive band of CaO, since the energy band gap
250 is reduced and charge carrier separation is increased, therefore improving
251 photocatalytic performance.

252 < Insert Fig. 1 here >

253 3.1.2. BET analysis

254 All samples exhibited Type IV isotherms (Fig. S1 a and b) with hysteresis loops.
255 The presence of small hysteresis loops at $P/P_0 > 0.4$ indicated the existence of both
256 microporous and mesoporous structures (Yao et al., 2020). After calcination, specific
257 surface area, pore volume and average pore diameter of the catalyst all decreased
258 (Table S1). This was because as a result of an increase in calcination temperature, the
259 inner part of the particles shrunk more, leading to an overall decrease in the pore size
260 of the material. However, specific surface area of the catalyst did increase slightly
261 between 800 and 1000°C, which was attributed to the formation of micro pores
262 following decomposition of calcium carbonate. Adherence to the Type IV isotherm
263 model clearly indicated the formation of multilayers on the catalyst surface during
264 high temperature reactions, which was also clearly observed in TEM imagery (Fig 1
265 i).

266

267 3.1.3. X-ray diffraction analysis

268 The XRD patterns of samples calcined at different temperatures (Fig. 2a),
269 showed that the main component of untreated clam shell was CaCO_3 , where
270 diffraction peaks at $2\theta = 26.21^\circ$, 27.21° , 45.85° and 50.22° correspond to calcium
271 carbonate on (111), (021), (221) and (132) crystal faces, respectively (PDF 70-1849).
272 However, the XRD pattern of samples calcined at 800°C for 2 h did not match well
273 with either CaCO_3 or CaO . This was because during heat treatment, calcium
274 carbonate, the main component of the clam shell, would have experienced major
275 structural changes from orthorhombic to trigonal-rhombohedral (Sree et al., 2020).
276 However, 800°C is not enough for complete conversion, so the calcined material
277 contains a large portion of intermediate phase/material between calcium carbonate and
278 calcium oxide. For samples calcined for 2 h at either 900 or 1000°C the diffraction
279 peaks at 32.20° , 37.35° , 53.85° , 64.15° and 67.37° corresponded to calcium oxide on
280 (111), (200), (220), (311) and (222) crystallographic planes, respectively (Jiang et al.,
281 2018). In addition, closer comparison of the XRD patterns of the samples calcined at
282 900°C and 1000°C , showed that several small peaks present at 900°C disappeared at
283 1000°C . This was attributed to either further mass loss of trace metals or the
284 complete conversion of intermediate substance to CaO . During calcination, calcium
285 carbonate, the main component of clam shell, undergoes an initial phase change from
286 aragonite to calcite, and then subsequently from calcite to CaO .

287 < Insert Fig. 2 here >

288 3.1.4. Thermogravimetric analysis

289 Thermogravimetric analysis provides further information on the transformation

290 of clam shell powder during calcination. The initial mass lost from clam shell was
291 only 0.47% due to the loss of water between 30 - 200 °C (Fig. 2b). Thereafter, the
292 curve decreased slightly between 200 - 500 °C, due to the decomposition and removal
293 of organic components of the clam shell, whereas weight loss between 500 – 610 °C
294 was attributed to a phase transformation from aragonite to calcite. Thereafter, the
295 mass of clam shell decreased significantly between 610 and 800 °C, due to further
296 conversion of calcite to CaO and CO₂, during calcination (Shariffuddin et al., 2018).
297 At 800 - 900 °C, further mass loss was caused by removal of any residual
298 intermediate products from incomplete conversion of CaCO₃ to CaO, and thereafter
299 the slight decrease observed at 900 - 1000 °C, may be due to the presence of a small
300 amount of unconverted CaCO₃ and further mass loss of trace metals. All these results
301 were in good agreement with both XRD (Section 3.1.3) and XPS (Section 3.1.6)
302 analysis.

303

304 *3.1.5. Ultraviolet visible spectrum analysis*

305 Diffuse reflectance spectroscopy in the UV-vis range was employed to
306 investigate the optical properties of uncalcined clam shell powder and clam shell
307 calcined at three different temperatures (Fig. 2c), as well as the light absorption
308 characteristics and band gap energy of the CSCs (Fig. 2d). The wavelength absorption
309 maxima of uncalcined, 800, 900 and 1000 °C samples were 277, 271, 276 and 273
310 nm, respectively (Fig. 2c), corresponding to the optical band gap obtained from Tauc
311 diagram (Fig. 2d). It was found that the optical band gap of the catalysts decreased

312 with an increase in calcination temperature, and thus the lowest band gap (2.64 eV)
313 was obtained at 1000 °C. As the calcination temperature increased, the weight of clam
314 shell decreased, and the specific gravity of trace metal elements increased. Therefore,
315 the decrease in optical band gap with increased calcination temperature was attributed
316 to the appearance of localised energy states in the CaO band gap and the existence of
317 a higher content of lattice defects, i.e., increases in trace metals and oxygen vacancies
318 (Shajahan et al., 2020). High resolution TEM (Section 3.1.1) had also confirmed the
319 existence of a large number of lattice defects.

320

321 *3.1.6. XPS analysis*

322 In order to analyze the valence state and content of surface elements in clam
323 shell powder before and after high temperature (1000 °C) calcination, XPS was
324 performed and compared on clam shell treated at 800 and 1000 °C (Fig. 3). Survey
325 scans (Fig. 3a), indicated that the main components of clam shells at 800 and 1000 °C
326 were C, O, Ca and a small amount of Na. It is likely that traces of other metals such as
327 magnesium (Mg) are also present but these would be at levels below the detection
328 limit of XPS (i.e., < 0.01 %).

329 For clam shell powder calcined at 800 °C, the XPS spectra (Fig. 3b) exhibited
330 many superimposed C1s peaks (C1-C3), where C1 with a binding energy of 285 eV,
331 was attributed to amorphous carbon/adventitious carbon, a small standard C2 peak at
332 286.1 eV, corresponded to the C-O, and the C3 peak at 289.4 eV corresponded to a
333 binding energy commonly associated with metal carbonate MCO_3 , and was thus

334 attributed to CaCO_3 . In comparison, for clam shell calcined at 1000 °C (Fig. 3e), the
335 binding energy of the adventitious carbon peak C1 occurred at 285 eV, the C2 peak
336 for organic carbon (C-O bond) occurred at 286 eV, the C3 peak attributable to COOR
337 at 288 eV, and the C4 peak corresponding to MCO_3 at 289.4 eV. However, compared
338 with the sample calcined at 800 °C, the sample calcined at 1000 °C, exhibited two
339 extra weak peaks (C5 and C6) at 292 and 295 eV, respectively, which were attributed
340 to compounds formed by carbon and/or halogen. The observed peak (O1) at 530.8 eV
341 for the 800 °C calcined sample in the O1s XPS spectra was attributed to CO_3^{2-} of
342 CaCO_3 (Fig. 3c); whereas the binding energy of the peak at 531.6 eV (peak O2) for
343 the clam shell calcined at 1000 °C (Fig. 3f). corresponded to CaO.

344 Calcium typically exhibits two characteristic binding energies, i.e., $\text{Ca}2p_{3/2}$ (346
345 eV) and $\text{Ca}2p_{1/2}$ (349.6 eV) (Tangboriboon et al., 2012). Here high-energy resolution
346 spectra of Ca2p peaks showed a complex pattern that could be deconvoluted into three
347 (Ca1, CA2 and Ca3) constituent peaks (Fig. 3d and g). For sample calcined at 800 °C,
348 peak Ca1 (346 eV), corresponding to CaCO_3 and peak Ca2 (347.1 eV) was associated
349 with CaO. In comparison, for the 1000 °C calcined sample the area of the CaO peak
350 (Ca2) increased, indicating that the content of CaO was relatively more when the
351 sample was pyrolyzed at the higher temperature.

352 < Insert Fig. 3 here >

353

354 3.2. Adsorption-photodegradation performance

355 3.2.1. Effect of different catalysts

356 While the total adsorption-photodegradation removal efficiency of MB and CR
357 for calcined clam shells at 1000 °C reached 99.7 and 90.8%, respectively, under dark
358 conditions, the adsorption only removal efficiencies of MB and CR reached only 25.9
359 and 70.5%, respectively (Fig. 4). In comparison the total adsorption-photodegradation
360 removal efficiency of untreated clam shells (MB: 15.9%, CR: 10.4%), and those
361 calcined at 800 °C (MB: 53.9%, CR: 25.9%), or 900 °C, (MB: 85.7%, CR: 81.1%)
362 was lower. This indicated that the adsorption and/or photocatalytic properties of
363 uncalcined clam shell or clam shell calcined at temperatures < 1000 °C were poorer
364 than clam shell calcined at 1000 °C. It can also be clearly seen (Fig. 4) that the
365 adsorption performance (dark environment) of clam shell calcined at different
366 temperatures was consistently better than that of non-calcined clam shell. This
367 suggested that decomposition of residual organic matter in the original clam shell via
368 heat treatment during calcination, results in significantly more pore structure. Where
369 the increase in adsorption performance with calcination temperature, could be
370 attributed to increased pore forming effects from CO₂ generated by thermal
371 decomposition of CaCO₃, the main component of clam shell. The photocatalytic
372 activity (under light) of the clam shell powder was also enhanced by an increased
373 calcination temperature. This may be due to an increase in the relative content of
374 calcium oxide due to heating and the interaction of trace transition metals in the clam
375 shell. HR-TEM images confirmed the presences of a significant number of
376 dislocations and defects in the CaO crystal structure. Overall, the results from the
377 adsorption-photodegradation study indicated that the best photocatalytic performance

378 was obtained when clam shell powder was calcined at 1000 °C. This was attributed to
379 synergetic effects of enhanced absorption and increased heterojunctions among some
380 dislocations in the crystal structures, which also improved photodegradation
381 efficiency.

382 Photodegradation kinetics of both MB and CR by CSCs varied with the
383 calcination temperature of the catalyst (Fig. 4b and d). The kinetic data for the
384 degradation of both MB and CR best fit the pseudo first order kinetic model (R^2
385 approaching 1), where the rate constant (k) increased with calcination temperature
386 (Table 1), and the higher the k value, the better the photocatalytic performance and the
387 greater the photodegradation amount. Thus, in all subsequent experiments, only clam
388 shell calcined at 1000 °C were investigated.

389 The photodegradation of both MB and CR by 1000 °C calcined CSC occurs in
390 the visible light region of the absorption spectrum (Fig. 4e and f). As reaction time
391 increased, peaks of MB and CR both decreased sharply, and some peaks disappeared
392 or were slightly shifted in wavelength, which indicated that both MB and CR
393 molecules were being destroyed, and that new substances (degradation products) were
394 also formed (Shariffuddin et al., 2018).

395 < Insert Fig. 4 here >

396 < Insert Table 1 here >

397

398 3.2.2. Adsorption performances

399 The adsorption performance of 1000 °C calcined clam shell powder varied with

400 the initial concentrations of MB and CR within 2 h. (Fig. S2) Over 2 h the
401 concentration of both MB or CR decreased gradually after exposure to the CSC until
402 the adsorption equilibrium was reached. When the initial dye concentration was low,
403 equilibrium was reached relatively quickly, within 30 min). For example, for MB a
404 sharp decrease in MB concentration was observed in the first 10 min followed by a
405 slower adsorption until the MB concentration stabilized. This is because when the
406 initial dye concentration is low, the adsorption reaction can quickly reach equilibrium
407 because there are a large number of sites available on the surface of CSC for
408 adsorption relative to the number of MB molecules. As the initial concentration of dye
409 increases, a more competitive adsorption for MB molecules for surface sites is
410 induced. Thus, as more active sites become occupied, it becomes increasingly more
411 difficult to adsorb dye due to a weaker attraction between the surface of CSC and the
412 MB molecules, which leads to slowing of dye adsorption rate in the latter stages of the
413 reaction.

414 In order to further investigate the adsorption mechanism of the MB dye on to the
415 CSC, both pseudo first order (PFO) (Eq 4) and pseudo second order (PSO) (Eq 5)
416 kinetic models were used for data analysis. The kinetic data for MB adsorption on to
417 CSC fit both the PFO (Fig S2(c)) and PSO (Fig. S2(d)) models well with good linear
418 behavior. However, the goodness of fit (R^2) for the PSO model was slightly higher
419 than that for the PFO model, and the adsorption capacity at equilibrium obtained from
420 PSO fitting was also closer to the experimental value (Table S2). The corresponding
421 adsorption capacities for initial MB concentrations of 50, 75 and 100 ppm were

422 experimentally is 60.52, 96.72 and 123.45 mg g⁻¹, respectively. Therefore, the PSO
423 model was more suitable for describing the adsorption system operating for the CSC.
424 Based on the underlying assumptions of the PSO equation model this indicates that
425 MB participated in chemical adsorption, involving the sharing or exchange of
426 electrons between the hydrophilic edge sites of the CSC and the dye cations
427 (Hosseinzadeh et al., 2015). As the initial MB concentration increased, the goodness
428 of fit of adsorption capacity (q_e) also increased, and became very close to the
429 experimentally derived q_e value, but the adsorption rate (k_2 value) decreased. This
430 decrease in rate constant is due to the intense adsorption competition between
431 relatively limited surface-active sites and a large number of MB molecules in a higher
432 concentration dye solution, resulting in a relatively low-rate constant (Liu et al.,
433 2014).

434 Similarly, the initial adsorption capacities of CR at 400, 455 and 500 ppm were
435 592.76, 653.26 and 679.91 mg g⁻¹, respectively and fitting of the kinetic data to the
436 same two models (Fig. S2 (g) and (h)) showed that the PSO model better described
437 the adsorption system of CR on the CSC, and calculated adsorption amount at
438 equilibrium was also closer to the experimental value (Table S2). As for MB, since the
439 PSO model assumes chemical adsorption as the dominant adsorption process
440 (Ardejani et al., 2008; Parvin et al., 2019), which involves the sharing or exchange of
441 electrons between the hydrophilic edge sites of the CSC and the dye ions, this result
442 also suggests that negatively charged CR ions were removed from solution via a
443 chemical adsorption process. This adsorption may be caused by a combination of π - π

444 stacking, hydrogen bonding, and van der Waals forces. In addition, the k_1 values for
445 CR were much lower than those calculated for MB, which indicated that the CSC had
446 a higher affinity for the negatively charged dye (Dai et al., 2018).

447 3.2.3. *Effect of catalyst dose*

448 The dose of catalyst is an important practical consideration and also plays an
449 important role in the efficiency of photocatalytic reaction. An increase in catalyst dose
450 will increase the overall concentration of active substance and hence increase the
451 number of active sites in the reaction system. Thus, the probability of contact of dye
452 molecules with the active sites increases, as does the reaction probability, and rate of
453 reaction, leading to improved dye removal efficiency. For photodegradation removal
454 processes there will generally be an increase in removal efficiency with increased
455 dose until at a certain concentration the increase in catalyst amount reduces the light
456 transmittance of the suspension (Nguyen et al., 2018) and hence the removal
457 efficiency.

458 For MB and CR (Fig. 5) as the amount of CSC increased from 20, 40, 60 to 80
459 mg, the removal efficiencies of MB were 99.7, 99.8, 99.9 and 99.9%, respectively,
460 and 83.3, 88.6, 96.7% and 97.0% for CR, respectively. Thus, for both dyes' removal
461 efficiency increases with catalyst dose, being almost a negligible for MB and only
462 modest for CR. This indicated that even the lowest dose used here had an adsorption
463 capacity well above the amount of dye in solution. Even though the increase in MB
464 removal efficiency with dose was not very significant, the time required to achieve
465 this removal became less as dose increased (Fig. 5b). Since even a 20 mg catalyst

466 dose could achieve a strong catalytic effect, in 40 mL dye solutions a catalyst dose of
467 0.5 g L^{-1} was selected for all subsequent experiments. According to the experimental
468 data, enough binding sites can be reached at 20 mg. Although the removal efficiency
469 further increases with the increase of the amount of catalyst, it will cause the waste of
470 catalyst, that is, some more binding sites are not used.

471 < Insert Fig. 5 here >

472 < Insert Table 2 here >

473 The best fit kinetic parameters (Table 2) for MB and CR degradation under
474 different catalyst doses showed that the degradation of both MB and CR followed
475 pseudo first order kinetics, with good correlation (R^2 close to 1).

476 The k value for MB increased slightly in the range of 20 - 40 mg, and decreased
477 between 40 and 80 mg. This was potentially because the increase of catalyst above 40
478 mg reduced the light transmittance of the suspension, making it more difficult for
479 photons to reach the CSC surface, and the amount of $\bullet\text{OH}$ free radical formed
480 between dye molecules and catalyst surface became relatively small. However, for
481 CR, the k value increased significantly between 20 and 60 mg, and slightly increased
482 from 60 to 80 mg.

483 3.2.4. *Effect of initial dye concentration*

484 The removal efficiencies of MB dye at initial dye concentration of 50, 75 and
485 100 ppm were 99.8%, 99.7% and 99.6%, respectively (Fig. 6). Although the removal
486 efficiency did not significantly change with initial dye concentration, the time
487 required for the photodegradation become longer (Fig. 6). Likewise, after 120 min of

488 illumination, the overall removal efficiency of CR dye decreased with an increase in
489 the initial dye concentration, being 98.2, 90.8 and 83.3%, for CR concentration of
490 400, 450 and 500 ppm respectively. This is because the number of •OH radicals
491 formed on the catalyst surface and the number of interactions with dye molecules
492 determine the overall efficiency of photocatalytic degradation. With an increase in the
493 number of dye molecules in solution, the number of dye molecules contacting the
494 catalyst surface also increase, causing active sites on the catalyst surface covered by
495 them, these will affect the arrival of photons on the surface of the catalyst, which will
496 reduce the catalytic efficiency, resulting in less •OH radical generation on the catalyst
497 surface. Therefore, increased dye concentration can lead to a reduction in both
498 degradation efficiency and reaction rate (Zhu et al., 2000).

499 The variation in best fit degradation kinetic parameters for both MB and CR with
500 different concentrations are shown in Table 3. The k values of both MB and CR
501 increased as the initial dye concentration decreased. The maximum k value of 0.058
502 min⁻¹ was observed for MB, which demonstrated that this catalyst had the highest
503 photocatalytic degradation rate for MB. This was also much higher than result
504 obtained using other semiconductors such as TiO₂/MoS₂ with k = 0.040 min⁻¹ (Ibukun
505 et al., 2019), g-C₃N₄/MnV₂O₆ (1:1), k = 0.022 min⁻¹ (Nithya et al., 2019);
506 CaTiO₃:0.5%Eu³⁺, k = 0.005 min⁻¹ (Chen et al., 2020b); and Bi₂CrO₆, k = 0.006 min⁻¹
507 (Li et al., 2020).

508 < Insert Fig. 6 here >

509 < Insert Table 3 here >

510 3.2.5. Influence of free radical scavengers on catalyst performance

511 Since the photodegradation efficiency of the catalyst is likely to depend on the
512 concentration of free radicals at the surface, free radical scavengers can affect
513 photodegradation performance. Here, in order to examine the factors affecting the
514 performance of the catalyst, isopropanol (IP) as $\bullet\text{OH}$ radical scavenger, p-
515 benzoquinone (BQ) as an $\bullet\text{O}_2^-$ radical scavenger and ammonium oxalate (AO) as h^+
516 scavenger were added to the reaction system (Wen et al., 2016; Chen et al., 2020a).
517 Under illumination from a xenon lamp, the removal efficiencies of MB with BQ, AO
518 and IP were 88.6%, 93.9% and 95.0%, respectively (Fig. 7) compared to a MB
519 removal efficiency of 99.9 with no added scavengers. Similarly, the removal
520 efficiencies for CR after adding BQ, AO and IP were 64.9, 83.5 and 88.4%
521 respectively, compared to 90.8% without any added scavenger.

522 The presence of BQ, significantly inhibited the removal efficiencies of both MB
523 and CR. This was because BQ leads a decrease in the amount of superoxide radicals
524 present, which leads to a significant decrease in the photodegradation rate (Table 4).
525 In comparison, AO and IP induced only slight decreases in the removal efficiencies of
526 MB and CR. These experimental results imply that while the formation of $\bullet\text{O}_2^-$ and h^+
527 in the photocatalytic reaction are both essential for a high removal efficiency, $\bullet\text{O}_2^-$ is
528 the dominant reactive species in this reaction.

529 These observations allow a removal mechanism to be proposed. When the CSC
530 is irradiated by a xenon lamp, h^+ electron holes are generated in the collision gap.
531 Oxygen molecules in the mixed solution form $\bullet\text{O}_2^-$ superoxide radicals by electron

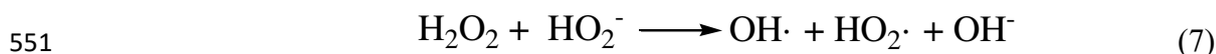
532 removal, and holes (h^+) react with water to form $\bullet OH$ radical (Baliarsingh et al.,
533 2014).

534 < Insert Fig. 7 here >

535 < Insert Table 4 here >

536 The addition of hydrogen peroxide (H_2O_2) is often used as an amendment to
537 enhance photocatalytic degradation of organic materials by enhancing free radical
538 generation. Here, under H_2O_2 assisted photocatalysis, the removal efficiencies of MB
539 and CR were 99.9% and 97.1%, respectively, indicating that compared to the blank
540 samples (99.6% and 90.8%), the addition of H_2O_2 significantly improved the
541 photodegradation removal efficiency. The reaction rate constants of the two dyes were
542 also close to each other, indicating that these two dyes were removed via similar
543 mechanisms.

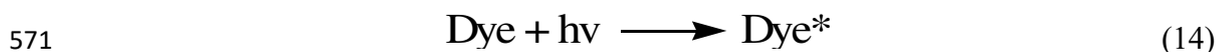
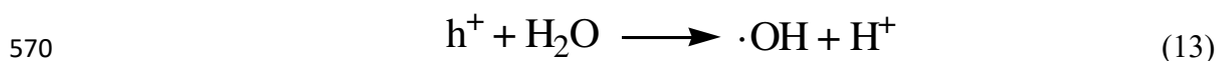
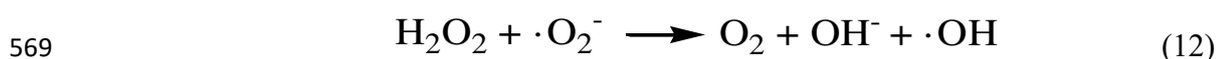
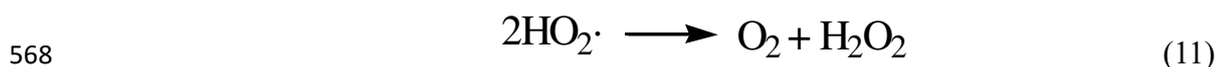
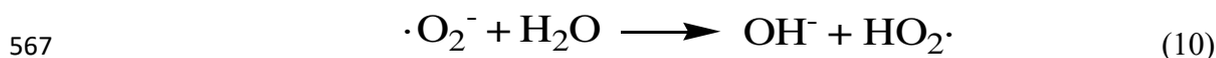
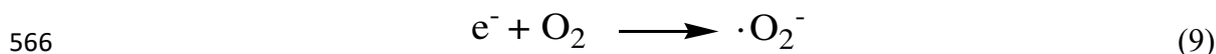
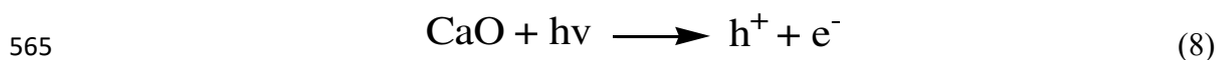
544 H_2O_2 assisted photocatalysis works through the formation of hydrogen peroxide
545 ion (HO_2^-) via reaction with H_2O , accelerating the hydrolysis of H_2O_2 forming OH
546 radical, and is accompanied by the formation of hydrogen hydrate ion (H_3O^+)
547 (Kausley et al., 2017), and the further reaction of HO_2^- and H_2O_2 forms OH radical,
548 which is the decisive step of the degradation rate of organic pollutants in aqueous
549 solution (Li et al., 2018). The reaction mechanism is summarized as follows:



552

553 In the process of photocatalytic degradation of dyes, the electrons generated by
554 the catalyst can react with O_2 to form $\bullet O_2^-$, and then the products can react with water

555 to form hydrogenated oxygen radicals (HO₂•) and hydroxyl radicals (OH⁻). The
 556 generated HO₂• rearranges to form oxygen (O₂) and peroxide hydrogen (H₂O₂)
 557 (Kumar et al., 2015), and H₂O₂ reacts with superoxide radical to form hydroxyl
 558 radical (OH⁻). The generated hole (h⁺) interacts with water to form a highly reactive
 559 hydroxyl radical, the hole itself can also attack dye molecules to produce by-products
 560 as well. After the formation of free radicals, the reactive substance (•O₂⁻) attacks dye
 561 molecules until the end of decolourisation and ring opening reaction, effectively
 562 degrading the dye into smaller intermediates and the final products (CO₂ and H₂O)
 563 (Kaviyarasu et al., 2017; Magdalane et al., 2016; Chen et al., 2008). The possible
 564 photocatalytic reactions are as follows:



573

574 3.3. Proposed photocatalytic mechanism

575 Based on the above experimental studies, a possible mechanism for the
 576 photocatalytic degradation of dyes by the CSC produced here was proposed (Fig S3).
 577 Under light irradiation, the electrons gain sufficient energy to become excited and

578 transition from the valence band to the conduction band, and form excitons (electron-
579 hole pairs). These excitons are known to act in four potential ways: (1) intralattice
580 recombination, (2) recombination at surface active sites, (3) hole (h^+) oxidizes
581 external dyes, and (4) electron (e^-) reduces external dyes. High photocatalytic activity
582 can be achieved through the minimum photocatalytic system of electron hole
583 recombination, and the combination of effective absorption and enhanced electron
584 transition from heterojunctions among some dislocations in the crystal structures (Xu
585 et al., 2018; Park et al., 2014). It was previously reported that photocatalytic
586 degradation efficiency can also be further improved via effective size reduction of
587 CaO nanoparticles and the innate nano properties of the initial raw materials
588 (Anantharaman et al., 2016). Hence, calcium oxide nanoparticles synthesized by
589 calcination of marine biomass, such as shells or other biomass materials containing
590 metal oxides and trace transition metals, which can act as excellent natural
591 photocatalysts for the degradation of dyes from the textile industries. In contrast,
592 while the photodegradation efficiency of 50 mg CaO derived from eggshell biomass
593 catalyst can reach 96.2% at an initial MB dye concentration of 20 mg L⁻¹ through
594 parameter optimization (Sree et al, 2020), here, the photodegradation efficiency using
595 a 20 mg CSC derived from marine biomass reached 99.6% for an initial MB dye
596 solution concentration of 100 mg L⁻¹.

597 *3.4. Feasibility of practical application*

598 In the textile industry, dye wastewater typically contains many kinds of different
599 dyes which need to be simultaneously removed. Hence here, the feasibility of

600 applying a CSC for practical wastewater treatment was evaluated in a mixed dye
601 solution. The results indicated that 20 mg of CSC could easily degrade both a mixed
602 dark colored binary dye solution (100 ppm MB + 100 ppm CR) and a ternary dye
603 system (75 ppm MB + 75 ppm CR + 75 ppm rhodamine B (Rh B)) into a transparent
604 solution (Fig S4). Analysis of the absorption spectra of the mixed dye solution system
605 under visible-light irradiation, showed that the initial peaks of the mixed dye solution
606 decreased sharply as reaction time increases, with some peaks shifting slightly or
607 completely disappearing, which strongly suggested that mixtures of dye molecules
608 were destroyed or degraded, and new substances were formed.

609 **4. Conclusion**

610 A novel photocatalyst nanomaterial derived from natural clam shell was
611 successfully prepared via a facile calcination process. The as prepared material was
612 then evaluated for its potential to treat dye contaminated wastewater. Under xenon
613 lamp illumination, the overall absorbance- photocatalytic removal efficiencies for 100
614 mg L⁻¹ MB and 500 mg L⁻¹ CR reached 99.6 and 83.3 %, respectively. Increasing the
615 calcination temperature enhanced the catalytic performance of the catalyst. This was
616 attributed to synergetic effects of increased absorption together with increased
617 heterojunctions among some dislocations in the crystal structures. Increased numbers
618 of heterojunction can greatly reduce the energy required for electron transition,
619 making it much easier to form holes and excited electrons. Consequently, the
620 inexpensive clam shell photocatalyst, synthesized in this work from a marine
621 biowaste, may have great potential for future practical large-scale industrial

622 wastewater treatment applications.

623

624 **Declaration of Competing Interest**

625 The authors declare that they have no known competing financial interests or
626 personal relationships that could have appeared to influence the work reported in this
627 paper.

628

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636

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783 **Figure captions:**

784 **Figure 1.** SEM images of clam shell samples: (a), (b) uncalcined; (c), (d) treated at 800 °C; (e), (f) treated
785 at 900 °C; (g), (h) treated at 1000 °C; and TEM image (i), (j) and (k) treated at 1000 °C.

786 **Figure 2.** (a) XRD spectra of uncalcined, treated at 800, 900, and 1000 °C clam shell samples. (b)
787 Thermogravimetric analysis of clam shell during calcination process. (c) Absorption spectrum and (d)
788 optical band gap of different clam shell samples.

789 **Figure 3.** (a) XPS survey scan of CSCs at 800 °C and 1000 °C; XPS spectra of clam shell samples at:
790 800 °C (b) Ca2p, (c) C1s, (d) O1s, and 1000 °C (e) Ca2p, (f) C1s, (g) O1s of CSCs.

791 **Figure 4.** Photocatalytic experiments of MB (a, b,) and CR (c, d,) with calcined catalysts at different
792 temperatures: (a), (c) curve of removal rate with time, (b), (d) plot of log of concentration with time.
793 Photodegradation absorption spectra of (e) MB and (f) CR solutions in the presence of clam shell
794 powder calcined at 1000 °C.

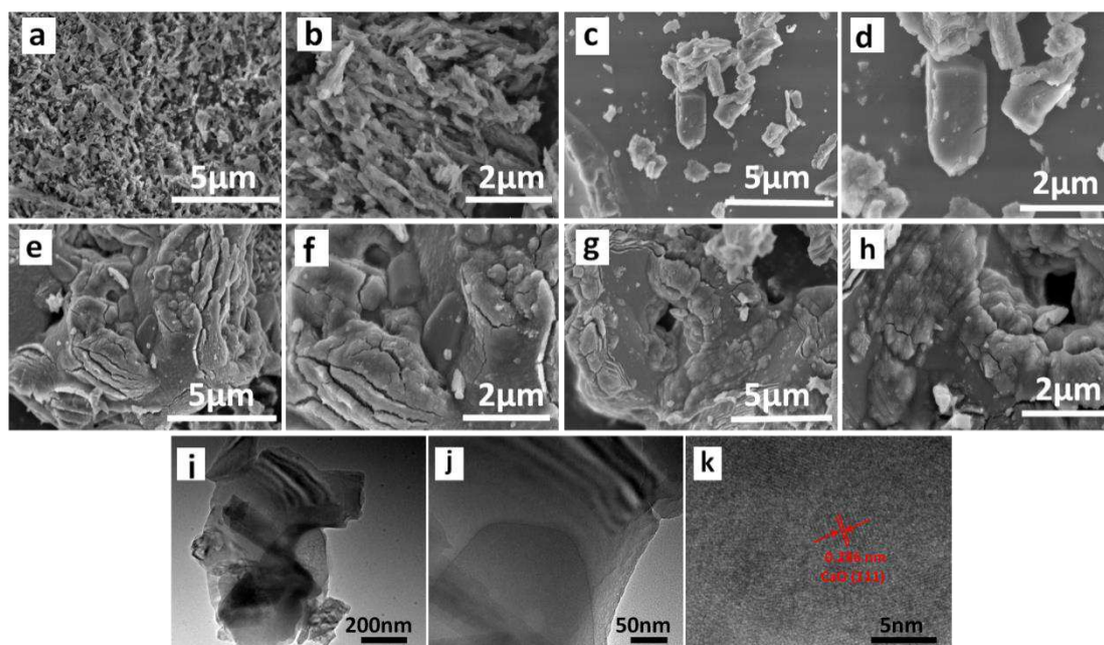
795 **Figure 5.** Photocatalytic experiment of MB (a, b, c) and CR (d, e, f) with different doses of clam shell at
796 1000 °C ((a), (d) curve of absorbance with time, (b), (e) curve of removal rate with time, (c), (f) plot of
797 log of concentration with time.

798 **Figure 6.** Photocatalytic experiment of clam shell at 1000 °C for MB (a, b, c) and CR (d, e, f); Curves
799 of absorbance with time (a) and (d); Curves of removal rate with time (b), (e); Plots of log of
800 concentration with time (c), (f).

801 **Figure 7.** Effects of scavenger and superoxide on MB (a, b, c) and CR (d, e, f) degradation.

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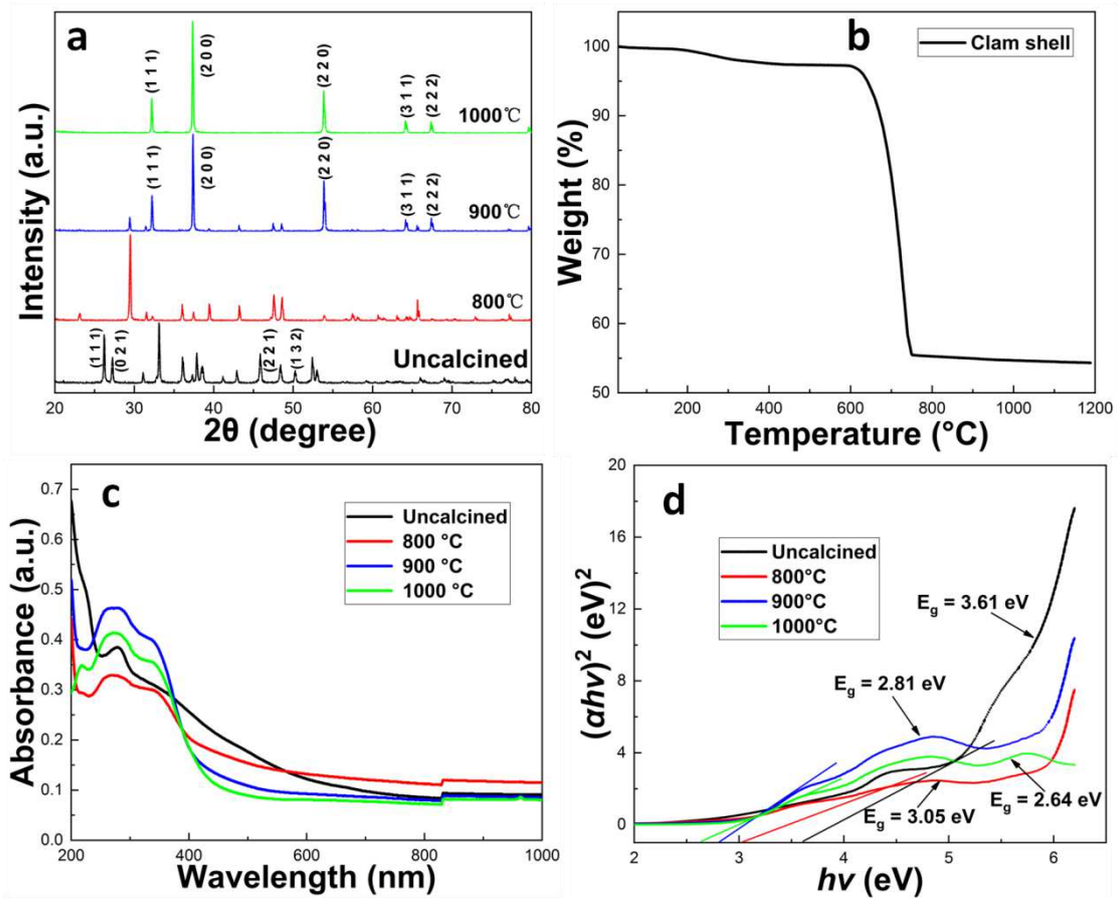
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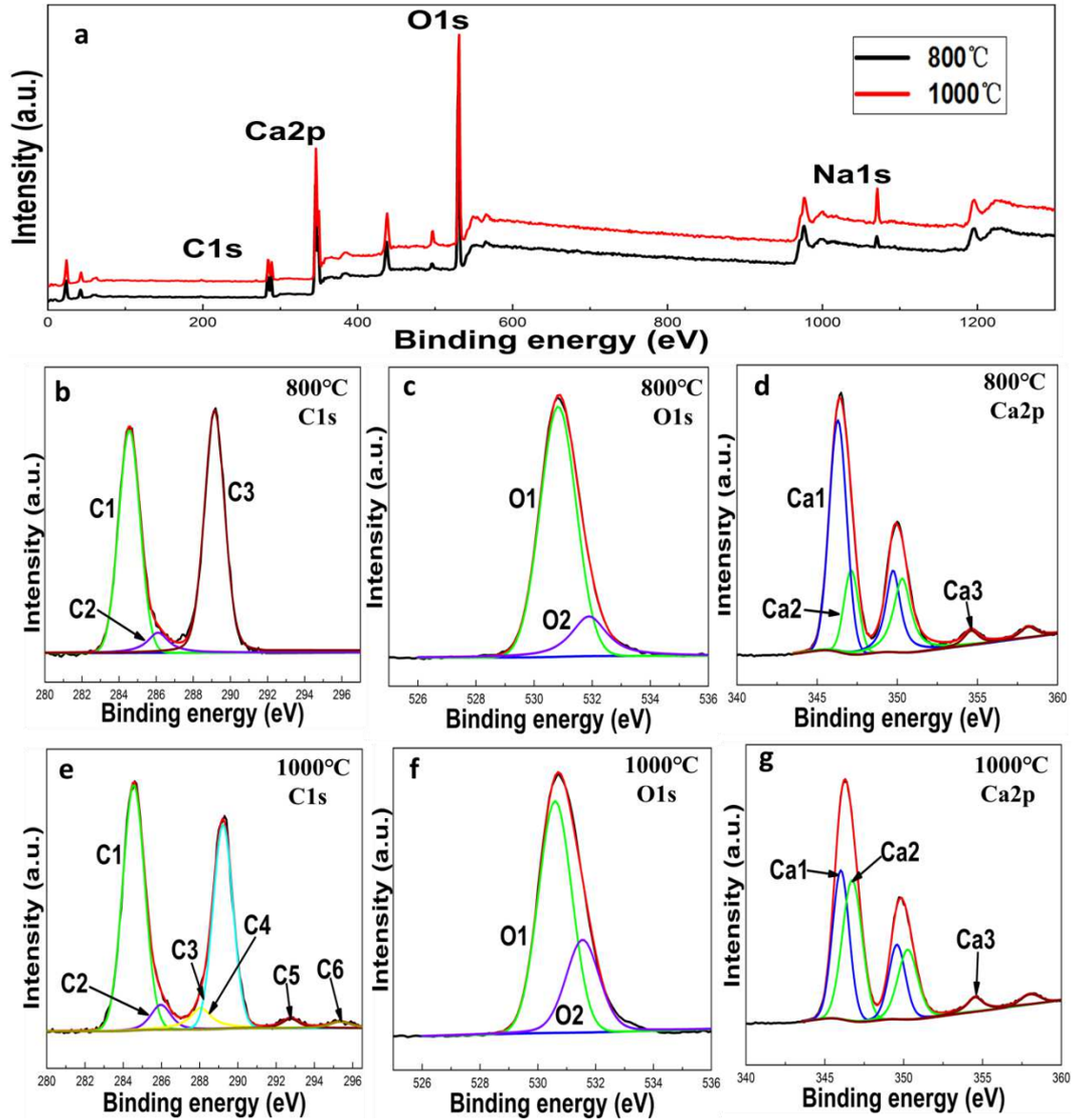
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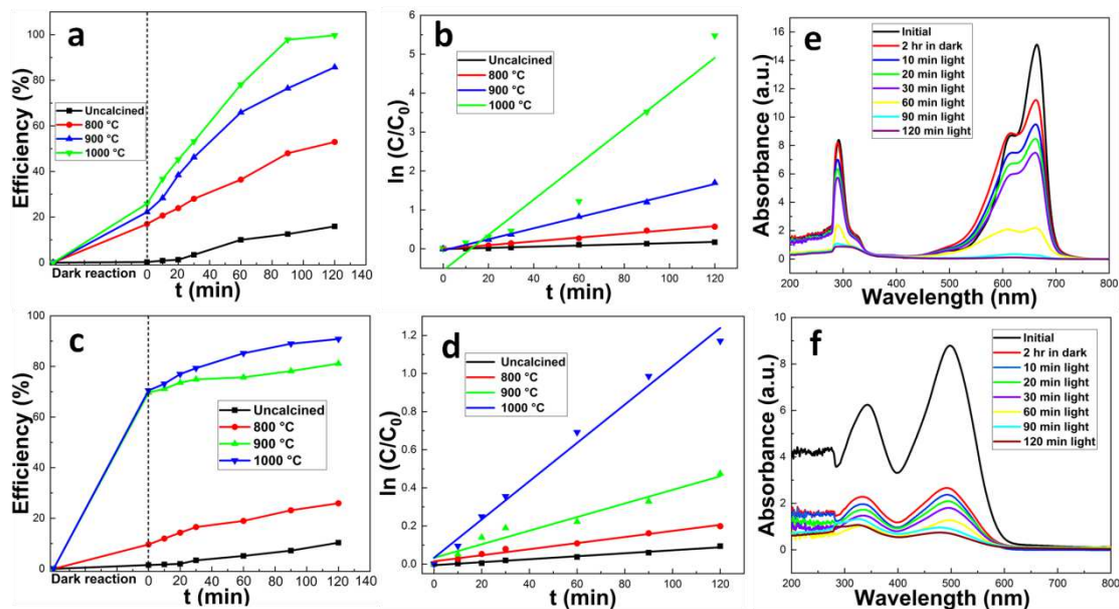
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837 Photodegradation absorption spectra of (e) MB and (f) CR solutions in the presence of clam shell

838 powder calcined at 1000 °C.

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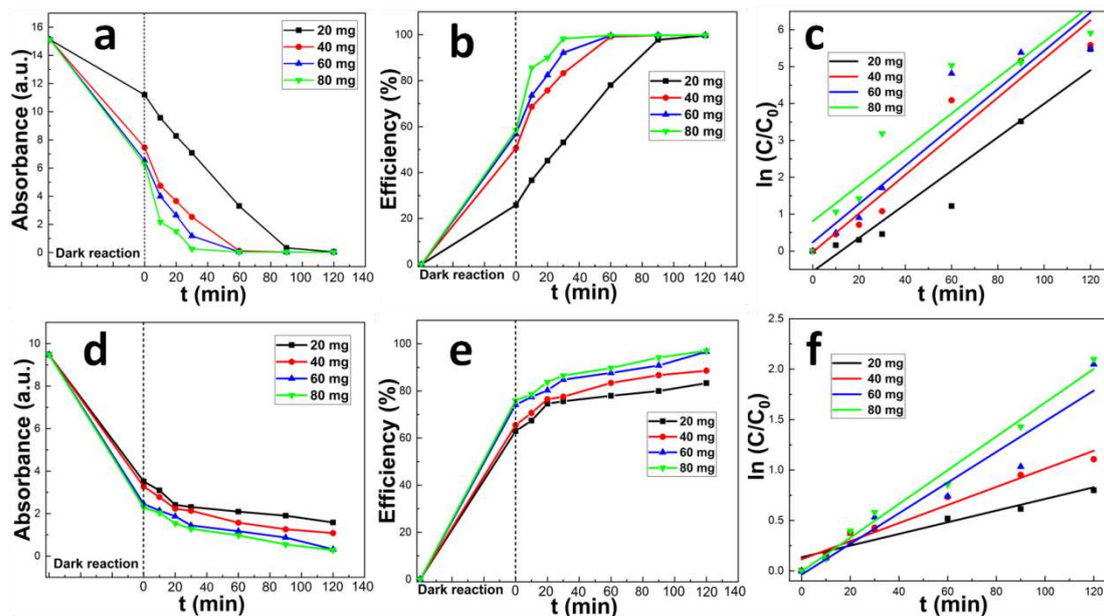
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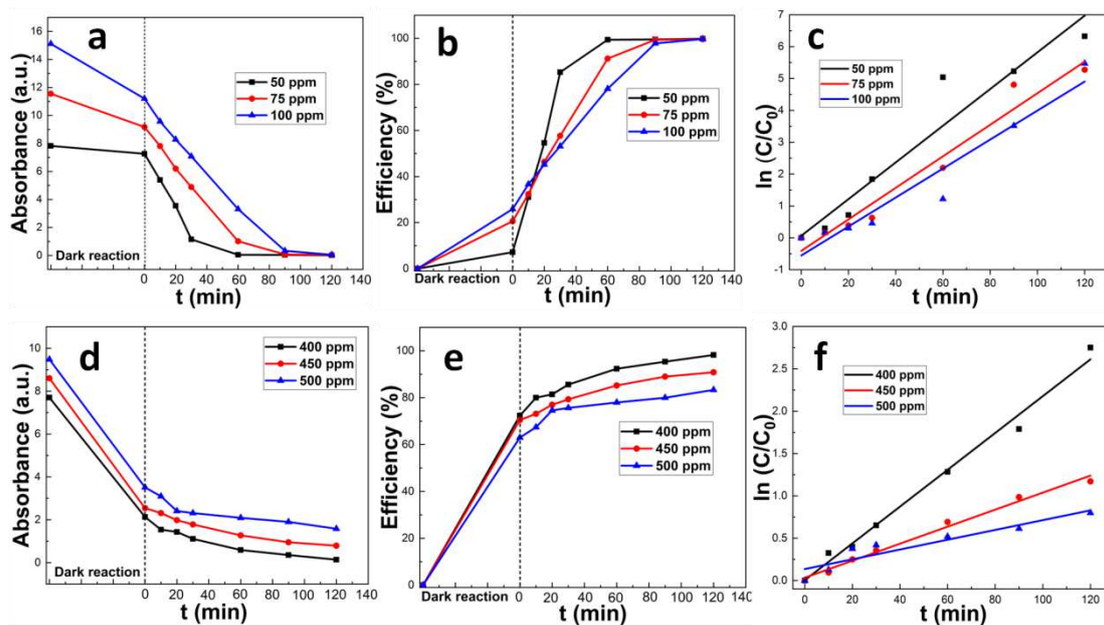
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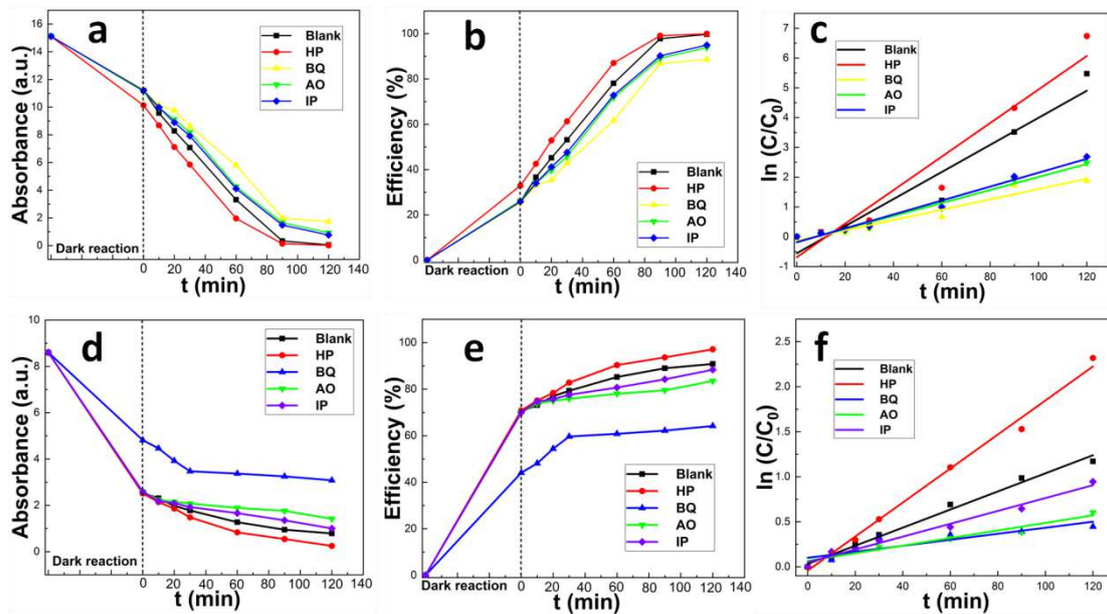
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884 **Figure 7.** Effects of scavenger and superoxide on MB (a, b, c) and CR (d, e, f) degradation.

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899 **Table 1.** Best fit photodegradation kinetic parameters for MB and CR with the four clam shells studied.

Dye	Sample	Uncalcined	800 °C	900 °C	1000 °C
MB	R ²	0.9733	0.9926	0.9973	0.9211
	k	0.0015	0.0049	0.0142	0.0455
CR	R ²	0.9779	0.9762	0.9509	0.9869
	k	0.0008	0.0016	0.0036	0.0101

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908 **Table 2.** Variation in best fit degradation kinetic parameters for MB and CR with the CSC dose.

Dye	Catalyst concentration	20 mg	40 mg	60 mg	80 mg
MB	R ²	0.9211	0.9332	0.8797	0.8541
	k	0.0455	0.0524	0.0519	0.0487
CR	R ²	0.9562	0.9531	0.9228	0.9839
	k	0.0058	0.0090	0.0152	0.0167

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917 **Table 3.** Variation in best fit degradation kinetics parameters for MB and CR with initial dye

918 concentration.

Dye	Dye concentration	50 ppm	75 ppm	100 ppm
MB	R ²	0.912	0.953	0.921
	k	0.058	0.049	0.045
	Dye concentration	400 ppm	450 ppm	500 ppm
CR	R ²	0.987	0.986	0.856
	k	0.022	0.010	0.006

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925 **Table 4.** Variation in best fit degradation kinetic parameters of MB and CR in the presence

926 different scavengers and superoxide.

Dye	Reagent	Blank	HP	BQ	AO	IP
MB	R ²	0.921	0.931	0.937	0.976	0.976
	k	0.045	0.056	0.017	0.022	0.023
CR	R ²	0.986	0.991	0.884	0.933	0.978
	k	0.010	0.019	0.003	0.004	0.007

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Supplementary Files

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