# Multi-way Degradation and Process Optimization of Phenol from Simulated Wastewater System

**Original Research Article** 

## 8 10 ABSTRACT

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This research based on the comparative study between microbial, enzymatic and photocatalytic phenol degradation. Different experiments were carried out under three distinct methodologies and seek to examine which method is more feasible between them through various aspects. For the microbial study, E-coli was used for phenol degradation at an optimum condition of E-coli. In an enzymatic study, peroxidase was extracted from soybean seed hulls, and it was purified. The purified peroxidase enzyme was applied in phenolic solution at neutral pH. The  $H_2O_2/UV/TiO_2$  scheme was adopted in the photocatalytic treatment of phenol. Maximum phenol degradation was observed in photocatalysis. From this comparative study, a microbial method was more time consuming and an enzymatic method having more steps to the experiment performed while photocatalysis had less time with a more feasible method.

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Keywords: Comparative study, microbial treatment, enzymatic treatment, photocatalytic
 treatment, first-order reaction kinetics

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#### 16 1. INTRODUCTION

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18 Recently, considerable attention received by biodegradation of aromatic compounds by many researchers due to their toxicity. Among them, phenol and its derivatives are a 19 20 standard compound in wastewater of many industries such as oil refineries [1], coal refining, 21 petroleum, textiles and pharmaceuticals [2]. It is quite known related to the toxicity of phenols towards the whole environment and has been incorporated in the list of pollutants by 22 23 the U.S. Environmental Protection Agency [3]. Many researchers engaged in research on phenol degradation by diverse techniques and methods. The attention is that to investigate 24 25 which technology will be most feasible, eco-friendly, cost-effective and time abstaining and 26 this idea is the primary goal of the present investigation. The present study comprises three parts viz. microbial degradation, enzymatic degradation and photocatalytic degradation. 27 28

Until today, many investigators have been reported numerous types of microorganisms to remove phenol from wastewater. From the literature review, some microorganisms can consume phenol as a sole source of carbon and energy. These bacterial species include Streptococcus enidermis [4]. Escherichia coli. Micrococcus en Brucella en [5]. Bacillus

32 Streptococcus epidermis [4], Escherichia coli, Micrococcus sp., Brucella sp. [5], Bacillus 33 subtilis, Pseudomonas putida, Acinetobacter calcoaceticus, Bacillus subtilis [6-8] and 34 Streptococcus sp. [8].

Besides, enzymes are applied in biodegradation study of the phenol. Enzymes play a vital role in phenol biodegradation reactions as a biocatalyst. These enzymes include Peroxidase,

37 Chloroperoxidase, Lignin peroxidase, Mn-peroxidase [9] and catalase [10] that isolated from

38 specific plants viz. soybean [11], horseradish, radish [12], and their materials such as seeds

[13], leaves [14], stem [15], roots [16]. Tyrosinase and Laccase [9] are obtained fromdifferent fungal species.

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42 In recent years, photocatalysis has been developed in wastewater treatment. In this 43 technique, some photocatalysts and their chemically modified transformations were 44 employed for the photodegradation of toxic compounds. The  $TiO_2$  and ZnO were broadly 45 worked as a photocatalyst in this technique [17-20]. Many researchers increase the 46 efficiency of a catalyst by doping with metals such as Ag, Fe, Pr, Co, V under various 47 illumination systems [21]. Some researchers synthesized bimetallic or trimetallic 48 transformations for degradation study [22].

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Here, we focus on all related aspects or parameters to select a better, efficient, cost-effective and feasible degradation technique. From the overall primary study, we use *E. coli* for the microbial study while peroxidase extracted from soybean seed hulls and selected for the further process of phenol degradation. Alike we introduced single  $TiO_2$  nanoparticles in phenolic wastewater under both UV and Solar light.

## 2. MATERIAL AND METHODS / EXPERIMENTAL DETAILS / METHODOLOGY

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### 58 2.1 Materials

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All analytical grade and HPLC grade chemicals were purchased from Fisher scientific and
 Himedia, Mumbai, India. Milli-Q water used for chemical preparations obtained from Milli-Q
 make of Schimadzu, Japan. E-coli microbial culture gave by my friend. Soybean seeds were
 collected from agricultural fields and washed thoroughly with distilled water.

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## 65 2.2 Microbial Methodology

E-coli bacterial culture was grown on slants of nutrient agar medium for further microbial 66 67 phenol degradation study and stored at 4°C until further use. Then the minimal salt medium 68 was prepared as Na<sub>2</sub>HPO<sub>4</sub> (33.9 g), KH<sub>2</sub>PO<sub>4</sub> (15 g), NH<sub>4</sub>Cl (5 g), NaCl (2.5 g), 2 ml of 69 MgSO<sub>4</sub> (0.1 M) and 0.1 ml of CaCl<sub>2</sub> (1 M) per liter for actual degrading study [4]. All media and required glassware autoclaved at 121°C and 15 lbs for 15 min. for sterilizing before the 70 commencement of experiments. Four consecutive same interval different concentrations of 71 72 phenolic wastewater were prepared in the range between 250 mg/L to 1000 mg/L in 73 phosphate buffer with pH 7.0. The reaction mixture had contained only MSM media and 74 phenol that was used as a control mixture in a microbial study. Similarly, bacterial inoculum 75 had been added to the control mixture for further phenol degradation study. Experiments 76 were carried out in a 250 ml conical flask containing 50 ml of MSM media with phenol concentration of above-given range. The mixture was incubated at room temperature (37°C 77 78  $\pm$  2) on the shaker (100 rpm). Samples were collected at every 24 h time interval for five 79 davs.

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The samples were centrifuged, and the remaining phenol concentration had been determined quantitatively by direct UV-visible spectrophotometric method [23]. Optical density was measured at  $\lambda_{max} = 269$  nm. Remaining concentration of phenol (%) was calculated as following formula:

% Phenol degradation = 
$$\frac{Absorbance \ of \ sample}{Slope \ phenol \ degradation \ (by \ graph)} \dots (1)$$

- 86 2.3 Enzymatic Methodology
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The experimental procedures of SBP extraction and purification were followed with some modifications reported by Liu et al. 2005. The fresh soybean seed hulls weighed and washed with milli-Q water. These cleaned seeds were soaked in milli-Q water for overnight. The soaked seeds were smashed and blended with 500 ml milli-Q water for 10 to 15 min. Then the homogenized mixture was filtered through cheesecloth and after that filtrate of cheesecloth centrifuged at 10,000 rpm for 20 min at 4°C. The collected supernatant was rich in proteins.

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96 The SBP purification process was performed as reported in Liu et al. 2005. The process 97 included three steps. A first step was acetone-ammonium sulphate cooperation precipitation. 98 It comprised both acetone and ammonium sulphate precipitation simultaneously. The 99 volume of acetone taken 0.3 fold of the original amount and solid ammonium sulphate added 100 to form up to 45% saturation. This combination placed in a refrigerator for 2 h. After that, the 101 mixture was centrifuged for 15 min at 5000 to 7000 rpm. The supernatant and precipitant 102 collected separately. This 45% saturation was continued to 75% saturation by adding solid 103 ammonium sulphate again with 0.3 fold acetone in the supernatant. A mixture was 104 centrifuged for 15 min at 5000 to 7000 rpm. Only one condition followed that the acetone 105 was pre-stored in a refrigerator and that cooled acetone was added under a cold 106 atmosphere in all our experimental sets. The resulted precipitants were dissolved in milli-Q water to get primary purified SBP. The second step consisted acetone precipitation lonely. 107 108 The volume of acetone mixed as 1.4 fold separately into the primary purified SBP. A mixture 109 was centrifuged for 15 min at 5000 to 7000 rpm. The resulted precipitant was dissolved in 110 milli-Q water to get secondary purified SBP. The third step included only zinc sulphate 111 precipitation. Before introducing zinc sulphate into the enzyme solutions, the pH adjusted on 112 eight by HCl or NaOH and then 1.0mol L-1 zinc sulphate solution was mixed to form 0.015 113 mol/l zinc concentration. A mixture was centrifuged for 15 min at 5000 to 7000 rpm. Lastly, 114 the supernatant was collected and denoted as highly purified SBP enzyme solution [24].

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116 Enzyme assay and protein content were examined after each purification step by the 117 procedures in Kolhe et al. 2015 [13]. The RZ values were assayed after each purification 118 steps. The purified SBP stored at 4°C till the further use of an enzyme. Different 119 concentrations of phenolic wastewater were prepared in the range between 250 mg/L to 120 1000 mg/L in phosphate buffer with pH 7.0. The reaction mixture contained 50 ml phenolic 121 wastewater, 30 per cent  $H_2O_2$  and enzyme solution. The sample was collected as a control 122 before kept for reaction and analyzed it. This combination kept on a rotary shaker for 10 h, 123 and aliquots were collected at every 1h time interval.

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125 The remaining phenol concentration of each sample had determined quantitatively by the 126 direct UV-visible spectrophotometric method at phenol  $\lambda_{max}$ . The remaining concentration of 127 phenol (%) was calculated by formula 1.

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### 129 2.4 Photocatalytic Methodology

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131 The third methodology opted as photocatalytic degradation of phenol. In this study, TiO<sub>2</sub> 132 nanoparticles were used as a photocatalyst while 11 watts of UV lamp was used as 133 illumination for energy. Various concentrations of phenolic wastewater were prepared in the 134 range between 250 mg/L to 1000 mg/L. The pH range kept as 2, 4, 6, 8 and 10 and adjusted 135 with 0.1 M HCl and 0.1 M NaOH solutions. The retention time was 10 h, but samples were 136 collected at every 1h time interval. The reaction mixture contained 50 ml phenolic solution, 137 30%  $H_2O_2$  and TiO<sub>2</sub> nanoparticles. The sample was taken as a control before kept on a 138 magnetic stirrer for reaction and analyzed it.

140 The remaining phenol concentration of each sample had determined quantitatively by the 141 direct UV-visible spectrophotometric method at phenol  $\lambda_{max}$ . The residual concentration of 142 phenol (%) was calculated by formula 1. The first and second order kinetics study were 143 evaluated from graphs of log concentration versus irradiation time [25].

145 3. RESULTS AND DISCUSSION

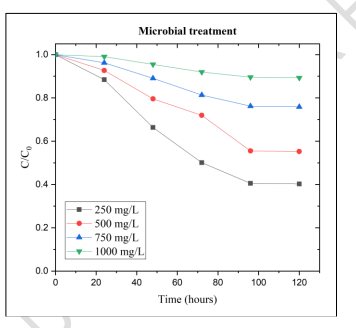
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# 147 3.1 Microbial Treatment148

149 The phenol degradation performance of E-coli strain was examined for different phenol 150 concentrations viz. 250 mg/L, 500 mg/L, 750 mg/L and 1000 mg/L at various time intervals. 151 The per cent phenol degradation was derived based on residual phenol concentration. 152 Figure 1 effect of phenol concentration shows that 60.07% phenol degradation observed at 153 250 mg/L phenolic concentration at neutral pH after 96 h, as the phenolic concentration 154 increases the phenol degradation decreases. Hence, only 11.75% phenol degradation 155 observed in 1000 mg/L phenolic concentration at neutral pH after 96 h. Reshma et al. also gave a treatment of E-coli on phenolic wastewater. They obtained 100% phenol degradation 156 157 for 10 mg/L phenolic solution. We had only 60.07% phenol degradation because 250 mg/L 158 concentration was much more than 10 mg/L concentration. Some bacterial strain may have 159 died at a more phenolic concentration; hence, the E-coli bacterial strain had not achieved 100% phenol degradation. 160

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164 Fig. 1. Phenol degradation by microbial treatment

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## 166 **3.2 Enzymatic Treatment**

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The SBP was extracted from soybean seed hulls by blending it for 10 to 15 min. During the blending of soybean seed hulls, the blended material was lightly warmed, but this thing is not essential because the SBP activity persisted up to 75°C [11]. A volume of the original enzyme solution was recorded as 530 ml. Table 1 shows the enzyme purification steps and their characteristics. A product of the last purification step having 71.01% recovery and 1.12 RZ value which is near about 1.32 RZ value reported in Liu et al. [24]. This enzyme

purification method is more comfortable and cost-effective than other purification methods
because it is merely based on only precipitation technique. Total volume, total activity,%
recovery, protein content, specific activity, fold purification and RZ value for each step were
showed in table 1.

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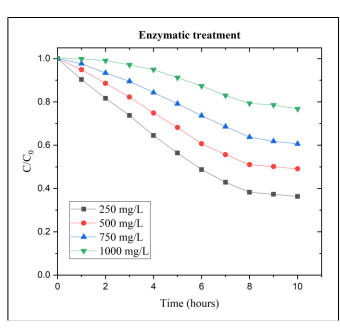
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#### Table 1. Purification steps and their characterization of SBP

Steps	Total Volume (ml)	Total Activity (U/ml)	Recovery (%)	Protein Content (mg/ml)	Specific Activity (U/mg)	Fold Purification	RZ value
Original enzyme solution Acetone-	530	6.091	100	2.325	2.62	1	0.19
ammonium sulphate cooperation precipitation	100	5.451	89.49	0.847	6.44	2.46	0.47
Acetone precipitation Zinc	10	4.847	79.58	0.461	10.51	4.01	0.83
sulphate precipitation	10	4.325	71.01	0.257	16.83	6.42	1.12

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This purified SBP was introduced in various phenol concentrations viz. 250 mg/L, 500 mg/L, 182 750 mg/L and 1000 mg/L at various time intervals to evaluate the phenol degradation. The 183 per cent phenol degradation was determined based on residual phenol concentration. Figure 184 2 effects of phenol concentration shows that 62.31% phenol degradation obtained in 250 185 mg/L phenolic concentration at neutral pH after 8 h. As in microbial treatment, here also 186 observed that as phenol concentration increases the phenol degradation decreases. Hence, 187 188 only 21.82% phenol degradation observed in 1000 mg/L phenolic concentration at neutral 189 pH after 8 h but this 21.82% phenol degradation is more as compared to microbial treatment. Pradeep et al. also gave a treatment of SBP on phenolic wastewater. They obtained 72% 190 phenol degradation of 100 mg/L phenolic solution. We had 62.31% phenol degradation in 191 250 mg/L concentration, which was more. 192



194 195

Fig. 2. Phenol degradation by enzymatic treatment

#### 3.3 Photocatalytic Treatment

### 200 3.3.1 Effect of pH condition

201 202 Some properties of photocatalysts are highly pH dependent. Hence phenol degradation at 203 different pH carried out under UV light. In this treatment, TiO<sub>2</sub> nanoparticles were used as a 204 photocatalyst. These nanoparticles introduced at different pH (2-10) conditions to examine 205 the phenol degradation. It is clearly seen that in figure 3, the basic conditions are unfavorable while acidic conditions are favorable for the photocatalytic degradation of 206 207 phenol. In acidic medium, from pH 2 to pH 6 phenol degradation increases and after pH 6 it was decreased. The higher phenol degradation was observed with 63.08% at pH 6. The 208 209 optimal pH condition was found acidic.

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211 Phenol has a pKa value of 9.95 and can be charged positively or negatively under the pH 212 range studied; i.e., the attraction and interaction between both photocatalyst and phenol will 213 be diverse with the solution pH. Moreover, as the pKa value of phenol is 9.95, it has negative 214 charge above pH 9.95 ≈ 10 and referred as phenolate anions but the conversion of 215 phenolate anions is commencing when solution pH in between 6 to 8 [26]. Conversely, in 216 highly acidic condition phenol gets a positive charge while in weak acidic and neutral 217 condition phenol molecules exist primarily in their non-ionic form. Additionally, the maximum 218 OH<sup>•</sup> radicals are produced in the pH range of 6 to 7 [27], due to this reason rate of phenol 219 degradation is higher in this pH range. These hydroxyl radicals, which are formed from some 220 photocatalytic oxidative and reductive reactions. They have a capacity to directly break down 221 of an aromatic ring of phenol molecule and transmute them into the final products are CO<sub>2</sub> 222 and H<sub>2</sub>O through various intermediates, because they are extremely strong, non-selective 223 oxidants [28].

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### 225 3.3.2 Effect of catalyst load

227 To examine the effect of TiO<sub>2</sub> nanocatalyst dosing on the phenol degradation, several 228 experiments carried out at catalyst loading from 1 to 4 g/L with 250 mg/L pollutant 229 concentration. Figure 3 indicates that the increase in the amount of nanocatalyst loading also increases the rate of phenol degradation up to a particular catalyst dose of 3 g/L. This 230 231 increased rate of degradation may be due to the higher surface area. Nevertheless, after 3 232 g/L amount of catalyst loading the degradation rate starts declining. As the catalyst load 233 increases, the experimental solution becomes turbid and resulting in UV rays getting 234 scattered resulting in a decrease in reaction rate [29]. The maximum phenol degradation at 3 235 g/L of catalysts doses considered as an optimum condition for further study.

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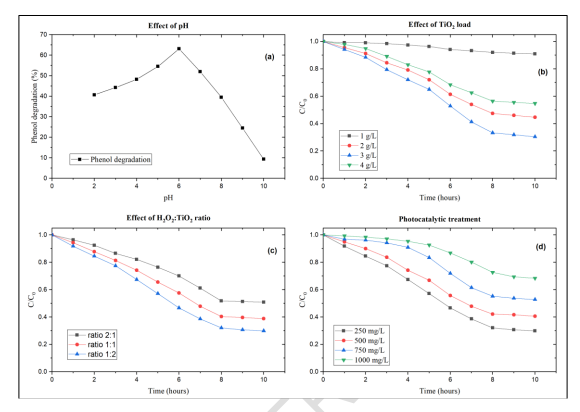
# 237 3.3.3 Effect of H<sub>2</sub>O<sub>2</sub> and TiO<sub>2</sub> ratio 238

3.3.4 Effect of phenol concentration

239 An oxidizing agent is another aspect of the photocatalytic oxidation process. Other 240 experimental sets were performed for the study of the impact of various rates between  $H_2O_2$ 241 and catalyst load as 2:1, 1:1 and 1:2. Figure 4 shows that a maximum phenol degradation 242 recorded at 1:2 ratio. It happens obviously because half the quantity of H<sub>2</sub>O<sub>2</sub> as on catalyst 243 dose was enough for phenol degradation. The H<sub>2</sub>O<sub>2</sub> used only an oxidizing agent in a 244 reaction medium. There is no use of a double quantity of  $H_2O_2$  in a reaction mixture. 245 Because in an excess amount of H<sub>2</sub>O<sub>2</sub> reacts with those hydroxyl radicals which are 246 responsible for degrading the pollutant molecule [30]. While the same quantities of H<sub>2</sub>O<sub>2</sub> and 247 catalyst load, also not well for the degradation because there is no sufficient amount of 248 catalyst in a mixture. This phenomenon also reported as earlier in 2001 by Ghaly et al. 249

#### 250 251

252 TiO<sub>2</sub> nanoparticles applied in various phenol concentrations viz. 250 mg/L, 500 mg/L, 750 253 mg/L and 1000 mg/L at various time intervals to evaluate the phenol degradation. The per 254 cent phenol degradation was determined based on residual phenol concentration. As initial 255 phenol concentration increases, the rate of phenol degradation decreases from 250 mg/L to 256 1000 mg/L. This happens due to the competitive adsorption on the active sites of 257 photocatalyst between the hydroxide radicals and phenol molecules [31]. Figure 4 effect of 258 phenol concentration shows that 68.39% phenol degradation obtained in 250 mg/L phenolic 259 concentration at neutral pH after 8 h. As in microbial treatment, here also seen that the 260 phenolic concentration increases the phenol degradation decreases. Hence, only 28.46 % 261 phenol degradation observed in 1000 mg/L phenolic concentration at neutral pH after 8 h, 262 but this 28.46% phenol degradation is more than in microbial treatment. Pradeep et al. also 263 gave a treatment of SBP on phenolic wastewater. They obtained 72% phenol degradation of 264 100 mg/L phenolic solution. We had 68.39% phenol degradation in 250 mg/L concentration, which was more. 265



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Fig. 3. (a) Phenol degradation at various pH conditions, (b) Effect of TiO<sub>2</sub> nanoparticles loading on phenol degradation, (c) Effect of H<sub>2</sub>O<sub>2</sub>:TiO<sub>2</sub> nanoparticle ratio on phenol degradation and (d) Effect of different phenolic concentration on phenol degradation under UV light

#### 274 3.3.5 Degradation rate kinetics

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The kinetic study of photodegradation of phenol was investigated for  $UV/H_2O_2/TiO_2$  system. A model with a higher value of correlation coefficient (R<sup>2</sup>) considered as more applicable. The equation for first and second order kinetics shown below.

279 First order reaction kinetics: 
$$\log(qe - qt) = \log(qe) - \left(\frac{Kf}{2.303}\right)t$$
 (2)

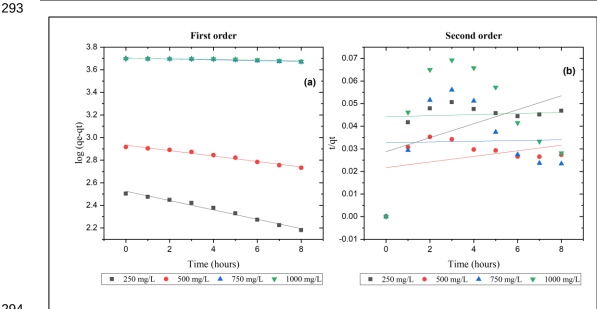
280 Second order reaction kinetics: 
$$\left(\frac{t}{qt}\right) = \left(\frac{1}{Ksqe^2}\right) + \left(\frac{1}{qe}\right)t$$
 (3)

Where  $q_e$  and  $q_t$  are the amounts of phenol degradation (mg g<sup>-1</sup>) at equilibrium time and at 281 time t (min), respectively. K<sub>f</sub> is the rate constant of first-order reaction (min<sup>-1</sup>) which can be 282 283 obtained from the slope of log (qe-qt) versus time plot. Also, a rate constant of pseudo-284 second-order  $K_s$  reaction (g mg<sup>-1</sup> min) can be obtained from t/qt versus t plot. For the phenol, 285 first-order reaction kinetic was fitted than second-order reaction kinetics first order having a maximum value of R<sup>2</sup>. Besides the apparent first-order rate constants decreased with the 286 increase of initial phenol concentrations [32]. Hence, kinetic constant based on phenol 287 288 degradation by UV calculated for a first-order reaction. Table no. 2 shows a description of first-order reaction kinetics. 289

290

#### 291 Table 2. Description of first-order reaction kinetics

Substrate	Concentration (mg/L)	K (min <sup>-1</sup> )	R <sup>2</sup>
Phenol	250	0.0953	0.9838
	500	0.0555	0.9793
	750	0.0088	0.8960
	1000	0.0067	0.8546



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Fig. 4. Phenol degradation corresponds to the (a) first-order and (b) second-order model for 250, mg/L, 500 mg/L, 750 mg/L and 1000 mg/L

### 299 4. CONCLUSION

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301 This study adopted three methodologies such as microbial, enzymatic and photocatalytic 302 treatment of phenol for the degradation. Microbial treatment gives 60.07%, enzymatic 303 treatment gives 62.31%, and photocatalytic treatment gives 68.39% phenol degradation in 304 250 g/L phenolic concentration. All treatments give approximately the same phenol 305 degradation, but each treatment has some advantages as well as some disadvantages. 306 About 60.07% phenol degradation achieved under 96 h in microbial treatment whereas 307 62.31% and 68.39% phenol degradation takes place under 8 h in enzymatic and 308 photocatalytic treatment. Based on the time parameter, microbial treatment is a very time-309 consuming method for phenol degradation while remaining both methods are less time-310 consuming.

311

In enzymatic treatment, additional one-step required for phenol degradation. That step was enzyme purification. Enzyme purification method was adopted in this study, and that the purified enzyme used as a catalyst. An enzymatic treatment did not show significant phenol degradation even after purified enzyme introduced in a reaction mixture. In phenol degradation follow another one-step and degrade the phenol which is not much more. Therefore, this enzymatic treatment is not a feasible method for phenol degradation.

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A remaining method is a photocatalytic degradation. It requires less time, no need for extra steps. The maximum phenol degradation achieved in this photocatalytic method, i.e. 68.39%. A whole photocatalytic study performed under acidic condition, this is one thing
noticeable. However, there is no need of extra handling of that acidic medium. Overall, from
the comparative study of all methods reported in this study, the photocatalytic process is
useful for phenol degradation than others.

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## 327 **COMPETING INTERESTS**

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Authors have declared that no competing interests exist.

- 331332 REFERENCES
- 333

334 1. Hilly M, Adams ML, Nelson SC. A study of digit fusion in the mouse embryo. Clin Exp
 335 Allergy. 2002; 32(4): 489-98.

336

337 1. Sukan A, Sargin S. Enzymatic removal of phenol from industrial wastewaters. Journal of
 338 Biomaterials and Nanobiotechnology. 2013; 4: 300-07.

2. Kolhe PM, Ingle ST, Wagh ND. Degradation of phenol containing wastewater by advance
catalysis system – a review. Annual research and review in biology. 2015; 8(3): 1-15.
(doi:10.9734/ARRB/2015/19936)

342 3. Environmental Protection Agency (EPA). Sampling and Analysis Procedure for Screening
 343 of Industrial Effluents for Priority Pollutants; EPA: Cincinnati, OH, USA, 1977.

4. Mohite BV, Jalgaonwala RE, Pawar S, Morankar A. Isolation and characterization of
phenol degrading bacteria from oil contaminated soil. Innovative Romanian Food
Biotechnology. 2010; 7: 61-5.

347 5. Reshma JK, Thanga VSG, Mathew A. Phenol degradation by bacteria isolated from coir
348 retting beds of south Kerala coast. Journal of Global Biosciences. 2014; 3(6): 935-940.

6. Liu Z, Xie W, Li D, Peng Y, Li Z, Liu S. Biodegradation of phenol by bacteria strain
Acinetobacter Calcoaceticus PA isolated from phenolic wastewater. Int. J. Environ. Res.
Public Health. 2016; 13: 1-8. (doi:10.3390/ijerph13030300)

352 7. Kafilzadeh F, Mokhtari S. Isolation and identification of phenol degrading bacteria from
353 mangrove sediments in the Persian Gulf (Asaluyeh) and their growth kinetics assay.
354 Biomedical and Pharmacology Journal. 2013; 6(2): 189-96.

8. Parihar R, Dubey SK. Identification and characterization of potential phenol degrading
bacterial strains isolated from municipal sewage, Bilaspur, Chhattisgarh. Int. J. Appl. Sci.
Biotechnol. 2016; 4(3): 288-93.

358 9. Rao MA, Scelza R, Scotti R, Gianfreda L. Role of enzymes in the remediation of polluted
asil sci. plant nutr. 2010; 10(3): 333-53.

10. Tayefi-Nasrabadi H, Dehghan G, Daeihassani B, Movafegi A, Samadi A. Some
biochemical properties of catalase from safflower (Carthamus tinctorius L. cv. M-CC-1 90).
African J. Agri. Res. 2011; 6(23): 5225-226.

- 363 11. Ghaemmaghami F, Alemzadeh I, Motamed S. Seed coat soybean peroxidase: extraction
  364 and biocatalytic properties determination. Iranian J. Chem. Eng. 2010; 7(2): 28-38.
- 365 12. Pradeep NV, Anupama, Hampannvar US. Comparison of horseradish peroxidase,
  366 soybean peroxidase and radish peroxidase for the polymerization of phenol. J. Res. Bio.
  367 2011; 7: 503-07.
- 368 13. Kolhe PM, Ingle ST, Wagh ND. Removal of phenol at higher concentration from aqueous
  369 medium using crude peroxidase catalyzed reaction coupled with hydrogen peroxide. Int. J.
  370 Adv. Sci. Tech. Res. 2015; 3(5): 70-8.
- 14. Anbuselvi S, Balamurugan Kumar S. Purification and characteristics of peroxidase from
  two varieties of Tulsi and Neem. Res. J. Pharmaceut. Biol. Chem. Sci. 2013; 4(1): 648-54.
- 373 15. Yadav P, Singh VK, Yadav M, Singh SK, Yadav S, Yadav KDS. Purification and
  374 characterization of Mn-peroxidase from Musa paradisiaca (Banana) stem juice. Ind. J.
  375 Biochem. Biophy. 2012; 49: 42-8.
- 376 16. Mamatha J, Vedamurthy AB, Shruthi SD. Degradation of phenol by turnip root enzyme
  377 extract. J. Microbiol. Biotech. Res. 2012; 2(3): 426-30.
- 17. Laoufi NA, Tassalit D, Bentahar F. The degradation of phenol in water solution by TiO2
  photocatalysis in a helical reactor. Global NEST J. 2008; 10(3): 404-18.
- 18. Bamuza-Pemu EE, Chirwa EMN. Profile of aromatic intermediates of titanium dioxide
  mediated degradation of phenol. Chem. Eng. Transac. 2013; 35: 1333-338.
  (doi:10.3303/CET1335222)
- 19. Parida KM, Parija S. Photocatalytic degradation of phenol under solar radiation using
  microwave irradiated zinc oxide. Solar Energy. 2006; 80: 1048-054.
  (doi:10.1016/j.solener.2005.04.025)
- 20. Malekshoar G, Pal K, He Q, Yu A, Ray AK. Enhanced solar photocatalytic degradation of
  phenol with coupled graphene-based titanium dioxide and zinc oxide. Ind. Eng. Chem. Res.
  2014; 53: 18824-8832. (doi.org/10.1021/ie501673v)
- 21. Chiou CH, Juang RS. Photocatalytic degradation of phenol in aqueous solutions by Prdoped TiO<sub>2</sub> nanoparticles. J. Haz. Mate. 2007; 149: 1–7.
  (doi:10.1016/j.jhazmat.2007.03.035)
- 392 22. Jethave GN, Fegade UA, Attarde SB, Ingle ST. Facile synthesis of lead doped zinc393 aluminum oxide nanoparticles (LD-ZAO-NPs) for efficient adsorption of anionic dye: kinetic,
  394 isotherm and thermodynamic behaviours. J. Ind. Eng. Chem. 2017. 53: 294-306.
  395 (doi.org/10.1016/j.jiec.2017.04.038)
- 396 23. Nicell JA, Saadi KW, Buchanan ID. Phenol polymerization and precipitation by
   397 horseradish peroxidase enzyme and an additive. Biore. Tech. 1995; 54: 5-16.
- 24. Liu J, Zhang Y, Qui L, Ye L, Xia Y, Su Z. A novel process of purifying soybean hull
  peroxidase. Chem. Biochem. Eng. Q. 2005; 19: 199-205.

400 25. Akbal F, Nur Onar A. Photocatalytic degradation of phenol. Env. Moni. Asses. 2003; 83:401 295-302.

402 26. Abbassian K, Kargari A, Kaghazchi T. Phenol removal from aqueous solutions by a
403 novel industrial solvent. Chemical Engineering Communications. 2015; 202(3): 408–13.
404 (doi.org/10.1080/00986445.2013.848804)

27. Nakabayashi Y, Nosaka Y. The pH dependence of OH radical formation in photoelectrochemical water oxidation with rutile TiO<sub>2</sub> single crystals. Physical Chemistry Chemical
Physics. 2015; 17(45): 30570–0576. (https://doi.org/10.1039/C5CP04531B)

28. Zhang Y, Selvaraj R, Sillanpaa M, Kim Y, Tai CW. The influence of operating parameters
on heterogeneous photocatalytic mineralization of phenol over BiPO<sub>4</sub>. Chemical Engineering
Journal. 2014; 245: 117–23. (https://doi.org/10.1016/j.cej.2014.02.028)

29. Dixit A, Mungray AK, Chakraborty M. Photochemical oxidation of phenol and
chlorophenol by UV/H<sub>2</sub>O<sub>2</sub>/TiO<sub>2</sub> process: a kinetic study. Int. J. Chem. Engi. Appl. 2010; 1(3):
247-50.

30. Ghaly MY, Hartel G, Mayer R, Haseneder R. Photochemical oxidation of p-chlorophenol
by UV/H<sub>2</sub>O<sub>2</sub> and photo-phenton process: a comparative study. Waste Management. 2001;
21: 41-7.

417 31. Ahmed S, Rasul MG, Brown R, Hashib MA. Influence of parameters on the 418 heterogeneous photocatalytic degradation of pesticides and phenolic contaminants in 419 wastewater: a short review. Journal of Environmental Management. 2011; 92(3): 311–30.

420 32. Matthews RW. Kinetics of photocatalytic oxidation of organic solutes over titanium dioxide. Journal of Catalysis. 1988; 111(2): 264–72.