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Highlights:

- OG worked as an intriguing photosensitizer in aiding the photodynamic inactivation
- The PDI was efficient at eradicating planktonic *P. fluorescens* and its biofilm
- The mode of bactericidal action of OG-mediated PDI was systemically investigated
- Bacterial inactivation by PDI might be due to multi-damage to cellular components
- Nanofibers in combination with PDI have superiorities for salamander preservation

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7	Ultra-efficient photodynamic inactivation system based on blue light and alkyl
8	gallate for synergistic antibacterial and ablation biofilms against Pseudomonas
9	fluorescens and smart application with electrospun nanofibers
10	on Chinese giant salamander preservation
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28 ABSTRACT

29 Pseudomonas fluorescens is a Gram-negative spoilage bacterium and dense biofilm producer, which will cause food spoilage and persistent contamination. Here, we report an ultra-efficient 30 photodynamic inactivation (PDI) system based on blue light (BL) and a multifunctional food 31 additive, octyl gallate (OG) to eradicate bacteria and biofilm of P. fluorescens. OG can rapidly 32 penetrate the cells and produce a high-level toxic reactive oxygen species (ROS) triggered by BL 33 irradiation. Both OG and ROS are critical to destroying the bacterial membranes and rupture cell 34 bodies, causing protein components alterations and DNA fragmentation in bacteria. Moreover, OG 35 plus BL irradiation can efficiently not only prevent the formation of biofilms but also scavenge the 36 existing biofilms. Additionally, the situ photodynamic antibacterial activity of OG/PLA electrospun 37 nanofibers was evaluated during the salamander storge. Our studies prove that the OG-mediated 38 PDI can provide a simple and ultra-effective platform for combating bacteria and eradicating 39 biofilm. 40

41

Keywords: Photodynamic inactivation; alkyl gallates; antimicrobial mechanism; reactive oxidative
 species; electrospun nanofibers; *Pseudomonas fluorescens*

45 **1. Introduction**

46 *Pseudomonas fluorescence* is a food spoilage bacterium largely responsible for the deterioration of aquatic products and dairy products, which can grow well at low temperatures and facilitate the 47 spoilage of aquatic products in cold-chain transportation as compared with other spoilage (Caldera 48 et al., 2016). Worse still, P. fluorescence cells can attach to food contact surfaces and form biofilms 49 readily, making it more difficult to be eliminated. Meanwhile, the biofilm protects organisms 50 against desiccation, biocides, some antibiotics and metallic cations, ultraviolet radiation (Flemming 51 & Wingender, 2010). To avoid these problems, antibiotics are often used to reduce their threat, but 52 indiscriminate use of antibiotics often might facilitate the emergence of drug resistance. Therefore, 53 it is urgent to develop efficient strategies to kill *P. fluorescence* and eradicate its biofilm. 54

55

As distinguished from traditional thermal-based technologies used for foods decontamination, some 56 non-thermal procedures including ultrasound, cold plasma, high hydrostatic pressure, pulsed 57 electric field, and pulsed light processing have been developed to inhibit the growth of 58 microorganisms and preserve nutritional quality and sensory acceptability of food (Ortega-Rivas & 59 Salmerón-Ochoa, 2014). Despite such significant potentials, some limitations, such as low 60 compatibility for broader food applications, higher processing requirements and costs, as well as the 61 emergence of microbial tolerance, limit the wide application of them (Cebrian et al., 2016). 62 Recently, a novel emerging non-thermal processing by light called photodynamic inactivation (PDI) 63 is gaining focus and are being applied for microbial growth control in the food industry (Ferrario et 64 al., 2015). PDI is an athermal photochemical reaction based on the combination of the simultaneous 65 presence of light, and photosensitizers (PSs) and oxygen (Luksiene & Zukauskas, 2009). Bacteria 66 containing PSs have the ability to absorb light at specific wavelengths. Once the light is absorbed, 67 the PS gets excited to a higher energy state under the presence of oxygen. On their way back to the 68 ground state, they collide with oxygen in the cytoplasm, transferring energy and subsequent 69 producing high reactive oxygen species (ROS). The ROS would interact with adjacent intracellular 70 components, such as lipids, proteins and nucleic acids, leading to bacterial death (Nakamura et al., 71 2012). Although UV light has been widely used to decontaminate foods and inactivate various 72 foodborne pathogens and biofilm cells, it has restricted application in the food industry due to its 73 low penetration ability for solids or opaque liquids and causing serious eye and skin damage to food 74 operators, as well as food sensory degradation (Kim et al., 2016). In contrast, LEDs can also 75 effectively inactivate pathogens and preserve food in postharvest stages and avoid the mentioned 76 issues related to UV radiation (D'Souza Yuk, Khoo, & Zhou, 2015). LED technology being 77 low-hazard (no mercury), low energy consumption, safe and high durability, as well as broader and 78

higher antibacterial effect on microorganisms, LED-based PDI have recently been explored more
and more as a novel preservation technology in food processing (Kim et al., 2017; Luksiene &
Zukauskas, 2009).

82

83 Lots of studies in PDI focus on the synthesis or discovery of more effective PSs. They can be divided into endogenous and exogenous PSs from the source. Porphyrins are the most well-known 84 natural endogenous PSs which are found in many bacterial and fungal cells (Rapacka-Zdonczyk et 85 al., 2019). When the magnitude of inactivation with endogenous PSs is lower than desired, it is 86 essential to use an exogenous PS to enhance it. At present, many artificially synthesized exogenous 87 PSs, such as chlorine, phthalocyanines and phenothiazinium dyes, etc, show good photoactivity. 88 However, safety considerations, organoleptic changes and consumer perceptions associated with the 89 use of an exogenous PS are also crucial to the application of PDI in food processing. Thus, natural 90 exogenous PSs such as hypericin, Vitamin K3 (Sheng et al., 2020), and curcumin are good 91 candidates for food application, given that they have no toxic or genotoxic effects. Recently, 92 phenolic compounds, especially phenolic acids, have been extensively studied in the food industry 93 due to their various bioactive properties, especially antimicrobial activities. Limited studies have 94 directly utilized them including gallic acid (GA), caffeic acid (CA), chlorogenic acid as PSs to 95 generate ROS including H₂O₂ and •OH radicals under the exposure of blue light in the presence of 96 dissolved oxygen, effectively inactivating bacteria (Nakamura et al., 2012; 2015; 2017). Electrons 97 are transferred from photo-oxidized polyphenols to dissolved oxygen to produce H₂O₂, which is 98 then photolyzed by blue light to produce •OH radicals (Nakamura et al., 2013), as the main 99 contributor to the bactericidal activity of such PDI. Besides, PDI technology based on 100 photo-oxidation of CA has been developed to eliminate S. mutans biofilm (Nakamura et al., 2017). 101 102 On the other hand, our research group has long been engaged in the study of the antibacterial activities and mechanism of a variety of phenolic acids and their ester derivatives (Shi et al., 2018; 103 2020; 2021). Some of them have been found to exhibit stronger antibacterial and anti-biofilm 104 activities against foodborne pathogens, compared to the corresponding phenolic acids. Among them, 105 octyl gallate (OG) showed the superior interaction/affinity with membranes and antibacterial 106 activity against E. coli and S. aureus (Shi et al., 2020), as well as P. fluorescence (Zhang et al., 107 2021). Surprisingly, the potential of OG with low concentration as a novel PS for PDI has been 108 found in this work, which endows OG-mediated PDI with intriguing bactericidal efficacy to achieve 109 rapid eradication of pathogens and biofilms in a relatively short time due to both photodynamic and 110 intrinsic antibacterial properties of OG itself. Besides, since OG has been permitted for use as an 111 antioxidant additive in food (FDA, 2001a, 2001b), it is supposed to be safe for humans and can be 112

considered as an alternative exogenous PS of PDI. Additionally, electrospinning is one of the 113 promising encapsulation methods in which a variety of active substances are encapsulated in the 114 nanofiber matrix (Wen et al., 2017). Electrospun nanofibers with a large surface area to mass ratio 115 have been proposed for stabilizing or controlling the release of the active compounds such as 116 117 antibacterial agents in food processing and packaging (Kayaci & Uyar, 2012), showing longer lasting antibacterial activity. Recently, we successfully utilized OG as a multi-functionalized food 118 additive combined with the advantages of electrospinning nanofibers for the preservation of Taihu 119 icefish in China (Shi et al., 2021). Therefore, inspired by the combination of these ideas, we chose 120 OG as a photosensitizer combined with blue light to inactivate P. fluorescence, and further 121 developed an ideal antibacterial strategy based on the combination of PDI and electrospinning 122 nanofibers to protect sea foods from spoilage bacteria, avoid their quality degradation and flavor 123 loss and extend the shelf life during storage. 124

125

Herein, the aim of the present study was to examine the synergistic effect of octyl gallate (OG) and blue light (BL) to inhibit the growth of *P. fluorescence* planktonic bacteria as well as eradicate its biofilm, and further elucidate the mode of action. Besides, we investigated electrospun nanofibers embedded with the photosensitizer OG as a PDI-based packaging material and evaluated its antibacterial activity in the storage and preservation of Chinese giant salamander. The dual role of OG in improving the production of ROS as a novel PS on one hand and the enhanced antimicrobial activity as an effective antibacterial on the other, are highlighted for the first time.

133 **2. Materials and methods**

134 2.1. Materials and light source

Octyl gallate (OG) was prepared by us and the structure was characterized by ¹H NMR and ¹³C 135 NMR. β -cyclodextrin (β CD) was purchased from Aladdin and acetic acid (99%, analytical reagent 136 grade) was obtained from Macklin, Shanghai. All other reagents used were of analytical grade. 137 Light-emitting diode arrays (420 nm, 10×10 cm, 90 W×2; Xuzhou Aijia Electronic Technology Co., 138 Ltd, China) with an emission maximum of 420 nm as a light source were used for PDI. The blue 139 LEDs were enclosed by deep photo accessories to avoid the entry of light from outside and the LED 140 system was equipped with a cooling fan and a heat sink to dissipate the heat. For irradiation, the 141 light sources were placed at 1.0 cm from the samples. The illumination energy of blue LED (212 142 143 mW/cm²) was determined by using an energy meter console (PM100D, Thorlabs, New Jersey, USA) attached with a photodiode power sensor (S120VC, THORLABS, Newton, USA). The dosage (in 144 J/cm^2) received by each bacterial suspension was calculated by multiplying the intensity (in W/cm^2) 145 by the irradiation time (in seconds). 146

147

148 2.2. *Microorganisms*

P. fluorescens isolated from Russian sturgeon (*Acipenser gueldenstaedti*) by us (Zhang et al., 2021) and *S. aureus* ATCC 6538 (CMCC, Beijing, China) were used in this study. The pre-cultured bacterial cells were inoculated into the fresh Luria-Bertani (LB) medium (Hangzhou Microbial Reagent, Co. Ltd, China), and grown to an exponential phase at 30 °C and 37 °C, respectively, with the agitation of 180 rpm. They were used as indicator strains for all experiments to evaluate the inhibitory activity of OG-mediated PDI.

155

156 2.3. Antimicrobial activity of PDI on the planktonic P. fluorescens in vitro

The OG+BL treatments were performed according to the methods (Nakumura et al., 2012) described before. The overnight bacterial culture of *P. fluorescens* was diluted in sterilized 0.1, 0.2 and 0.4 mM OG (or GA) solution prepared in normal saline to reach a final concentration of approximately 6 Log CFU/mL. Then, 5 mL of the bacterial suspension was transferred to a glass tube and followed immediately by BL exposure applied for 0 ~ 30 min (Irradiation intensity: 212 mW/cm², distance:1 cm). Bacterial suspension incubated in the dark with OG (or GA) for the same

duration was used as a control. Next, the treated solution was serially diluted in 0.9% (w/v) saline, 163 after which 100 µL of the dilution was seeded on LB agar plates. The plates were incubated at 30 °C 164 for 24 h before enumeration, and the reductions of bacteria were determined. To confirm whether 165 the bactericidal effect of PDI might be associated with ROS, ROS scavengers CAT (1200 U/mL), 166 167 DMSO (2.8 M) and TEMPOL (16 mM) were added simultaneously with OG. The cultured bacterial with ~6 Log CFU/mL P. fluorescens was treated in the absence or presence of BL light for 15 min. 168 The control without ROS inhibitor was incubated in the dark for the same duration. Subsequently, 169 the standard plate counting method described in 2.3. was performed on all samples. 170

171

172 2.4. Mechanism studies of the synergistic antibacterial treatment

173 2.4.1. Association of OG or GA with P. fluorescens

The uptake of OG (or GA) in P. fluorescens cells with or without BL irradiation was measured by 174 using diphenylboric acid 2-aminoethyl ester (DPBA) (Shi et al., 2021). A volume of 1 mL 0.9% 175 (w/v) saline containing 8 Log CFU/mL P. fluorescens was mixed with or without GA or OG (0.4 176 mM). Then, 1 mL of the bacterial suspension was added to a 24-well plate (Costar 3599, Corning, 177 USA) and exposed to BL for 20 min as described previously. Controls were treated in the same 178 manner, but in the absence of phenolics or BL irradiation. Then, each sample was transferred back 179 to the tube and centrifuged at 10,000×g for 5 min to remove the supernatant. The pellet was washed 180 twice using DI water followed by vortexing and then DPBA solution (450 µL, 0.2%, w/v, in DMSO) 181 was added to the pellet and then incubated for 5 min. The final suspension (200 µL) was transferred 182 to a 96-well plate and the fluorescence intensity was measured at excitation/emission wavelength of 183 405/465 nm using a microplate reader (Synergy H1, BioTek, Winooski, VT, USA). The 184 fluorescence intensity (I) ratio was corrected using the following equation: 185

186

$$I_{corrected} = I_{sample} - I_{control}$$

187 where I_{sample} stands for the fluorescence intensity of the sample exposed to the treatment and $I_{control}$ 188 stands for the fluorescence intensity of the control.

189

Quantification of phenolics (OG or GA) adsorption was also observed through confocal laser
 scanning microscopy according to the method reported by Wang et al. (2017) with some
 modifications. Detailed descriptions are given in SI (Section 1.1)

2',7'-dichlorofluorescein diacetate (DCFH-DA) was used to assess the generation of ROS in the 195 bacterial cells (Shi et al., 2021). The overnight test culture (8 Log CFU/mL) was treated by OG (0.4 196 mM) with or without ROS scavengers including catalase (CAT), dimethyl sulfoxide (DMSO) or 197 TEMPOL. The CAT, DMSO or TEMPOL solution was added into a sample solution to achieve a 198 final concentration of 1200 U/mL, 2.8 M and 4 mM, respectively. The suspensions were incubated 199 in the dark with OG for the same duration were used as positive controls. After BL irradiation for 20 200 min, the excess OG in samples was removed by centrifugation (10000×g, 4 °C, 5 min) and the 201 bacteria were resuspended with cold phosphate-buffered saline (0.1 mM PBS, pH 7.2-7.4). 10 µM 202 DCFH-DA was mixed with samples and treated in dark at 30 °C for 30 min. The incubated solution 203 was washed twice using PBS to remove excess dye. Finally, the fluorescence spectrum of solutions 204 was measured at excitation wavelengths of 485 nm with a wavelength of 525 nm with a microplate 205 reader (Synergy H1 Multi-Mode Reader, BioTek, Winooski, VT, USA). Bacterial suspension with 206 water treated in the dark was considered as a negative control, and its fluorescence intensity (F_0) 207 was used as the reference to calculate the relative fluorescence unit (RFU) for other treatments 208 using the equation: 209

210

Relative fluorescence unit (RFU) = F_s/F_0

211 where F_s was the fluorescence intensity of the sample with treatments.

212

Finally, to visualize the ROS generated by PDI treatment, the confocal laser scanning microscope (CLSM, Leica TCS SP8, Germany) using ×63 oil immersion objective lens was used to observe the cells at excitation/emission wavelength of 484/525 nm.

216

217 2.4.3. Hydroxyl radical experiments using flow cytometry.

Flow cytometry was used to detect hydroxyl radical formation. Hydroxyphenyl-fluorescein (HPF) is an anthracene derivative of fluorescein which becomes fluorescent when it was activated by hydroxyl radical (Shi et al., 2021). Detailed descriptions of the method are given in SI (Section 1.2).

222 2.4.4. Investigation of the cell membrane damage

To evaluate the cell membrane damage induced by OG+BL treatment, the uptake of propidium iodide (PI) and scanning electron microscope (SEM) were performed according to our previously published papers (Shi et al., 2020; 2021). Detailed descriptions of the methods are given in SI
(Section 1.3).

227

228 2.4.5. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE)

SDS-PAGE analysis was performed to determine the alternations of bacterial proteins before and
 after the treatment of OG and BL irradiation. Detailed descriptions of the method are given in SI
 (Section 1.4).

232

233 2.4.6. Genome integrity determination

The damage of DNA caused by OG+BL was investigated by the agarose gel electrophoresis.
Detailed procedures were offered in SI (Section 1.5).

236

237 2.5. Inhibition assay for biofilm formation

The assay method of Damiano et al. (2017) was carried out in 24-well flat-bottom polystyrene 238 plates (Costar 3599, Corning, USA). Briefly, 100 µL of P. fluorescens (~7 Log CFU/mL) were 239 diluted 1:10 into 900 µL of LB broth (1.0% tryptone, 0.5% yeast extract, 1.0% NaCl) in 24-well 240 plates to obtain the final concentration of ~6 Log CFU/mL and then cultivated with for 0~96 h 241 without agitation. After incubation, bacterial suspension of each well was gently decanted, the wells 242 were washed 3 times with sterile PBS (0.1 M, pH 7.2) to remove planktonic bacteria and biofilms 243 were stained for 30 min using crystal violet (0.1%, w/v). The stained biofilms were rinsed 3 times 244 with distilled water and extracted with 1 mL of 95% ethanol. Biofilms were quantified at 600 nm by 245 reading the microplates (Synergy H1 Multi-Mode Reader, BioTek, Winooski, VT, USA). 246

247

For determination of the effect of different treatments on biofilm formation, 100 μ L of *P. fluorescens* (the original suspension was diluted to ~8 Log CFU/mL) were diluted into 900 μ L of LB 24-well flat-bottom polystyrene plates containing (0.15 mM or 0.1 mM) OG (below the minimum inhibitory concentration), with ethanol serving as the control. After cultivation at 30 °C for 15 min in the absence or presence of BL irradiation (94.8 mW/cm², distance: 2 cm) (subinhibitory external condition). Biofilms were quantified by the crystal violet assay above.

To count planktonic bacteria, the suspension cultures (100 μ L) were centrifuged at 5000 rpm at 4 °C to collect the pellets, which were resuspended in 1 mL of 0.9% NaCl and 10-fold serially diluted for enumeration.

258

259 2.6. Per-formed biofilm eradication assay

P. fluorescens biofilms were incubated as described above. In brief, 100 µL of P. fluorescens were 260 added to 900 µL of LB to obtain the final concentration of ~6 Log CFU/mL (as described before) 261 and then cultivated at 30 °C for 60 h without agitation. The bacterial suspension was discarded and 262 wells were rinsed 3 times with sterile PBS to remove the loosely attached bacteria. To evaluate the 263 effect of OG on the biofilm, OG concentrations in fresh LB medium were added to the pre-formed 264 biofilm in the wells, with ethanol serving as an extra negative control. The plates were cultivated at 265 30 °C for 30, 60 and 90 min without shaking in the absence or presence of BL irradiation (212 266 mW/cm²). Biofilms were fixed with methanol for 15 min and staining was performed as before. To 267 quantify cell viability in biofilm, the treated biofilms were washed with PBS three times and 268 swabbed using a sterile cotton swab (Nakumura et al., 2017). The collected bacteria were suspended 269 in 1 mL of saline, and the mixture was serially diluted 10-fold with saline, and 100 µL of the 270 dilution was plated on LB agar at 30 °C overnight. On the other hand, for viable colony counts of 271 planktonic cells, the suspension cultures after treatments were diluted 10-fold with saline, and the 272 dilution (100 µL) was plated on LB agar for enumeration of planktonic cells. 273

274

275 2.7. Confocal laser scanning microscopy (CLSM)

To observe the formation of the biofilm after various treatments, samples were captured by CLSM refer to Seo and Kang (2020) with some modifications. Biofilm was stained using a LIVE/DEAD *Bac*light Bacterial Viability Kits (Molecular Probes, Invitrogen, USA) according to the manufacturer's instructions. The biofilm was washed with PBS and visualized using a CLSM. More details were given in SI (Section 1.6).

281

282 2.8. Preparation of nanofibers (NFs)

The NFs were prepared using the method reported by us (Shi et al., 2020) with some modifications. More detailed descriptions of the method are given in SI (Section 1.7). The morphology of The collected fibers was observed via SEM (Hitachi T-1000, Hitachi High-Technologies Corporation, Tokyo, Japan) (see Fig. S1) and the preservation of giant salamander was shown in Fig. S4.

287

288 2.9. In situ antibacterial activity of OG/PLA NFs

To assess the synergistic effect of OG/PLA NFs and PDI on the preservation of Chinese giant salamander, the changes in the total viable count and flavor of samples during storage were studied. The salamander meat cubes were inoculated by immersion in a bacterial suspension of *P. fluorescens* (3 Log CFU/mL) for 30 seconds. The treated salamander meats were singly sealed with OG/PLA NFs and followed immediately by BL exposure for 30 min at a distance of 1 cm. Meanwhile, to evaluate the quality of the giant salamander with different treatments during the storage, the electronic nose was applied. More details were given in SI (Section 1.8 and 1.9).

297 2.10. Statistical analysis

Results were expressed as mean \pm standard deviation. T Statistical significance between different treatments was determined using t-test; *P*-values ≤ 0.05 were used to determine significant differences.

301

302 3. Results and discussion

303 3.1. Bactericidal effect of the OG-mediated PDI on P. fluorescens cells suspension

We evaluated the use of BL illumination in combination with OG as an alternative approach to 304 achieve P. fluorescens inactivation (Fig. 1). P. fluorescens was killed synergistically by BL 305 combined with OG in both BL and OG dose-dependent manners in planktonic solutions. As shown 306 in Fig. 1A-(a), viable bacterial counts for P. fluorescens without any treatment were ~6.5 Log 307 CFU/mL, while a significant reduction was observed in the sample treated by BL irradiance (190.8 308 J/cm²) for 15 min. OG inhibited the growth of *P. fluorescens* in an OG concentration-dependent 309 manner, and the OG-mediated PDI led to higher antibacterial potency as compared with those 310 equivalents in the absence of BL irradiation (P < 0.01). Under 190.8 J/cm² irradiation of BL, the P. 311 fluorescens cells were decreased from 3.8 to 1.8 Log CFU/mL when the OG concentration was 312 elevated from 0.1 to 0.2 mM. Moreover, bacterial cells could not be detectable from the samples 313 treated by BL irradiation with 0.4 mM OG. In Fig. 1A-(b), compared to the positive controls with 314 only BL irradiation, significant decreases of bacterial cells were observed in samples with the 315

OG-mediated PDI (P<0.01). Also, the BL irradiation time (dosage) significantly influenced the activity of *P. fluorescens*. After 10 min irradiation with OG (0.1, 0.2 and 0.4 mM), the cells were decreased to 4.5, 3.5 and 2.2 Log CFU/mL (P<0.05), respectively. However, an obvious and continuous decrease in the bacterial cells was achieved when PDI irradiation time was extended from 10 to 15 min. Thus, these findings ensure that the efficacy of OG-mediated PDI-induced inactivation against *P. fluorescens* was typically dependent on the photosensitizer concentration and irradiation dosage.

323

In Fig. 1B, when the Gram-negative P. fluorescens and Gram-positive S. aureus were treated with 324 the OG-mediated PDI, viable counts decreased in a time-dependent manner. Moreover, S. aureus 325 tended to show higher susceptibility to the OG+BL treatment than P. fluorescens. It may be due to 326 that the S. aureus was more susceptible to OG than P. fluorescens. Also, the MIC and MBC values 327 of OG against S. aureus are lower than those against P. fluorescens (Table S1). Although treatment 328 with OG or BL irradiation performed in this present study could cause a reduction in the remaining 329 bacterial count, BL irradiation of the suspension containing 0.4 mM OG could significantly kill the 330 bacteria within 15 min. A stronger synergistic effect on bacteria in vitro was found when OG was 331 combined with BL illumination. 332

333

Cossu et al. (2016) also reported that a synergistic interaction between 10 mM gallic acid (GA) and 334 UV-A (365 nm) light could inactivate E. coli O157:H7. Nakamura et al. (2012) reported 335 that >5-Log CFU/mL reduction was obtained when the suspension of S. aureus was co-incubated 336 with 4 mM gallic acid (GA) for 15 min when exposed to LED light (400 nm; 80 mW/cm²). By 337 contrast, in the present study, neither GA alone nor the combined treatment with BL irradiation 338 339 showed a substantial bactericidal effect (<1-Log). Moreover, under the dark condition, OG showed higher bactericidal activity than GA (Fig. 1B). Also, it should be noted that although GA has 340 antimicrobial activity, OG exerted stronger inhibitory capacity than GA because Gram-negative P. 341 fluorescens and Gram-positive S. aureus were effectively killed by OG in LB media, at very low 342 concentration, presenting an MBC of 3.2 mM for P. fluorescens and 0.1 mM for S. aureus (Table 343 S1), respectively. Wang et al. (2017) reported that the effect of propyl gallate (10 mM)+UV-A on 344 the inactivation of E. coli O157:H7 was stronger than that treated by GA (10 mM)+UV-A. These 345 findings suggest that (1) the antimicrobial potency of OG depends largely on the hydrophobic 346 portion (the alkyl group) of the molecule, which is also in agreement with the findings reported by 347 Kubo et al. (2004) and us (2021). (2) As for the OG-mediated PDI, OG alone or BL irradiation does 348 not solely contribute to the remarkable reduction in the CFU of bacteria, reliably suggesting the 349

occurrence of interaction between OG and BL.

- 351
- 352

[Fig. 1.]

353 3.2. Antimicrobial mechanism of the OG-mediated PDI against P. fluorescens

354 *3.2.1. Interaction between OG and bacterial cells*

The remarkable photosensitizing efficiency of OG suggests that it could be exploited to obtain 355 efficient bacterial photoinactivation. To gain further insight into the mechanisms underlying the 356 antibacterial effects of OG-mediated PDI, we firstly analyzed the binding affinity of OG toward 357 bacteria. In the present case, DPBA has been employed to detect the cellular uptake of OG or GA in 358 359 P. fluorescens. It is a specific dye that becomes fluorescent when it was combined with flavonoid compounds (Shi et al., 2021). Fig. 2A shows that P. fluorescens treated with OG+BL has a higher 360 fluorescent intensity (3179 ± 294.0) than that incubated with OG alone (1147 ± 41.9) , indicating 361 that the BL irradiation increased the level of OG uptake in *P. fluorescens*. Also, the extent of uptake 362 of OG is significantly stronger ($P \le 0.05$) than that of GA regardless of the absence or presence of 363 BL irradiation, indicating that OG shows higher binding affinity capacity to bacteria compared to 364 GA. Moreover, the either OG or GA uptake results happen to coincide with their inactivation results 365 in Fig. 1B, indicating that the cellular uptake of OG was a vital element for OG-mediated PDI and 366 the higher uptake of OG under BL irradiation enhanced the antimicrobial effect. Their discrepancy 367 in the affinity to bacteria could be explained by their different hydrophobicity. The Log P value of 368 OG is 4.63 while it is 0.4 for GA (Table S1). Both of them possess the same hydrophilic portion, the 369 pyrogallol group, thus distinguishing the role of the hydrophobic alkyl portion of OG, as discussed 370 371 in our previous studies (Shi et al., 2018, 2019, 2021). The enhanced association of OG toward bacteria was further determined by using confocal microscopy. In Fig. S2, the amount of OG 372 internalized by or bound tightly with bacterial cells was remarkably more than that of GA during the 373 same period, further corroborating that the affinity to bacteria varied with the alkyl chain length. 374 Compared to GA, amphiphilic OG could more effectively bind to bacteria, leading to increased 375 intracellular uptake. These results illustrated that OG as an alternative antimicrobial 376 photosensitizing agent induced a more effective photokilling of pathogenic microbial cells than GA. 377 378

379 *3.2.2. OG and BL induced ROS generation*

380 It had been reported that ROS generation, especially interior hydroxyl radicals (•OH), is a critical

step for the bactericidal effect of BL irradiated polyphenols (caffeic acid, chlorogenic acid, gallic 381 acid, and proanthocyanidin) against a broad range of pathogens in the presence of dissolved oxygen 382 (Nakamura et al., 2015). Moreover, the reaction began with the photo-oxidation of the polyphenolic 383 hydroxyl group. As such, upon reaching the cells and following light activation, it is reasonable to 384 385 speculate that OG may be also excited by BL to generate ROS. Thus, to evaluate the generation of ROS in P. fluorescens induced by photoirradiated OG, DCFH-DA was used as a fluorescent probe 386 as it can be oxidized by cytosolic ROS to green fluorescent 2',7'-dichlorofluorescein (DCF). As 387 shown in Fig. 2B, bacteria treated by the BL irritation had significantly higher fluorescent intensity 388 than the corresponding samples without BL irradiation. Moreover, cells treated by OG+BL showed 389 the strongest fluorescent intensity, followed by bacteria treated with OG alone in the dark, 390 representing the efficient intracellular ROS production during the combination OG with the 391 exposure of BL. Even though P. fluorescens treated by GA had significantly lower fluorescent 392 intensity than bacteria treated by OG regardless of BL irradiation, GA with BL irradiation for 30 393 min could cause a higher fluorescent intensity as compared to samples treated by GA in dark. It is 394 noteworthy that these findings are also in consistency with their antibacterial capacities in Fig. 1B. 395 It indicated that ROS production plays a crucial role in OG-mediated photokilling and OG is a 396 superior photosensitizer as compared to GA. To further visualize the generation of ROS induced by 397 OG+BL treatment, P. fluorescens cells exposed to the selected treatments were observed using a 398 confocal laser scanning microscope (CLSM). In Fig. 2C, the combination of OG with BL irradiation 399 caused a higher level of ROS than both controls and OG alone, indicating that BL irradiation played 400 an important role in the production of ROS. To further ascertain the specific ROS generation in 401 OG-mediated PDI, the bacteria were incubated in the presence of several radical scavengers. We 402 exposed OG-treated *P. fluorescens* to the H_2O_2 scavenger (CAT), •OH scavenger (DMSO), and • O_2^- 403 404 scavenger (TEMPOL), respectively, in the absence or presence of BL irradiation. In Fig. 2D-(a), the addition of these scavengers could significantly reduce the relative fluorescence unit as compared to 405 406 the positive controls treated with OG alone. Moreover, as for BL irradiated samples, CAT and DMSO led to a higher level decline in the relative fluorescence unit than TEMPOL. Besides, the 407 importance of intracellular ROS formation in the bactericidal activity of the duo was established by 408 the ability of either CAT or DMSO (Fig. 2D-(b)). These findings indicated that the OG and BL 409 interaction promoted the production of ROS (mainly, H₂O₂ and •OH) which is essential for 410 OG-mediated PDI. Additionally, the intracellular •OH formation in the bactericidal action of 411 OG-mediated PDI was also confirmed by the flow cytometry with the dye HPF, which was 412 specifically oxidized by •OH radicals. The histogram distribution in Fig. 2E showed the contents of 413 intracellular •OH radicals of different groups. Compare to the samples exposed to OG or BL alone, 414

a higher hydroxyl radical content was observed in bacteria treated with OG+BL. Moreover, we
found DMSO significantly reduced the level of •OH induced by the duo. Together, these results
imply that •OH radicals formation in *P. fluorescens* is directly related to OG-mediated PDI.

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Akagawa et al.(2003) reported that catechins and other polyphenolic compounds could induce H₂O₂ 419 generation in solutions under neutral or alkaline conditions. Also, it was found that H₂O₂ synthesis 420 by (-)-epicatechin gallate is attributed to its bactericidal activity, which increases at higher pH 421 values (Arakawa, Maeda, Okubo, & Shimamura, 2004) and a possible mechanism for H₂O₂ 422 generation in a catechin solution was proposed by Mochizuki et al. (2002). In this case, blue light 423 might trigger H₂O₂ photolysis to form •OH, further bolstering the bactericidal activity of the duo 424 because •OH is the most destructive of all ROS to bacteria (Nakamura et al., 2015). And it has been 425 reported that 500 mM H₂O₂ can kill S. aurous only with 1-Log reduction (Kanno et al., 2012). Thus, 426 it is reasonable to assume that H_2O_2 would probably act as a source of •OH in the photolysis 427 reaction, rather than be a major contributor to the bactericidal action in a short time (15-30 min), 428 which is in agreement with the results reported by Nakumura et al. (2012). On the other hand, it has 429 been proved that the internalized OG may interfere with the activity of ETC on the cytoplasmic 430 membrane to produce toxic •OH, further leading to cellular damage and death (Shi et al., 2021; 431 Kubo et al., 2004). Therefore, we inferred that the ROS observed in the present study may be 432 produced directly by OG upon oxidation by BL in bacteria, and by indirectly mediating ROS 433 formation through activating a variety of intracellular metabolic pathways. A possible scheme for 434 potential pathways linking intracellular ROS generation induced by OG-mediated PDI is illustrated 435 in scheme 1. The internalized OG may interfere with the activity of ETC on the cytoplasmic 436 membrane, generating toxic ROS (Shi et al., 2021). More importantly, OG can be oxidized to 437 438 quinone or hydroxyl related derivatives with the production of H₂O₂ using BL as the oxidizing source and oxygen as an oxidation catalyst, owing to the tendency of the three aromatic hydroxyl 439 groups of OG to undergo autooxidation or oxidation (Wang et al., 2019). Then, H₂O₂ would be 440 turned into •OH through the photolysis reaction by BL (Nakamura et al., 2015). 441

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443 3.2.3. OG and BL induced alterations in the bacterial cell membrane, proteins and DNA

OG with a high affinity to the cell membrane may permeate the outer membrane to induce oxidative stress in bacterial cells. Then, the •OH generated around the phospholipid membrane will further have an effect on the structure and function of membranes. Some phenolic compounds had been observed to disintegrate the bacterial outer membrane (Shi et al., 2018, 2019). Thus, the membrane

damage of *P. fluorescens* was evaluated using PI as a fluorescent staining probe, and the increasing 448 fluorescence signal represented the severe damage to the cytoplasmic membrane of cells (Stiefel, 449 Schmidt-Emrich, Maniura-Weber, & Ren, 2015). In Fig. 2F, the presence of BL irradiation caused a 450 significant increase (P < 0.05) in the fluorescence intensity for all the treatments, compared to the 451 452 corresponding samples in the dark. Although the fluorescence intensity of cells exposed to 0.4 mM GA+BL or BL irradiation alone (NS) was significantly higher than their correspondings, no 453 bactericidal effect could be observed and logarithmic reduction of CFU/mL in these cases was 454 <1-Log (Fig. 1B). Bacteria treated by OG+BL showed the strongest fluorescence intensity of PI, 455 followed by the samples incubated with OG in the dark. However, only OG+BL treatment caused 456 substantial inactivation (>5-Log) in the reduction of P. fluorescens within 15 min and S. aureus 457 within only 10 min (Fig. 1B). It is interesting to note that the tendency in PI uptake happened to 458 coincide with the findings of cellular uptake of phenolics in Fig. 2A, indicating cellular uptake of 459 phenolics correlated positively with the damage of the bacterial membrane. SEM was used to 460 further observe alternations in the surface morphology of bacteria treated by OG+BL. Samples 461 treated by 0.4 mM OG without BL irradiation exhibit uneven and shriveled appearance (Fig. 462 2G-(b)), with a comparison of regular rod shape of controls (Fig. 2G-(a)). Serious damage to the 463 cell walls of *P. fluorescens* occurred during the treatment of OG+BL with cell collapsing and the 464 leakage of large cytoplasmic components (Fig. 2G-(c)), indicating that membrane damage was 465 strongly associated with the bactericidal effect of OG+BL treatment. 466

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The mentioned ROS may induce oxidative damage to the lipids, proteins and DNA of cells and 468 ultimately led to bacterial death (Park et al., 2009). The alterations of membrane proteins of P. 469 fluorescens upon OG-mediated PDI treatment were evaluated by using SDS-PAGE. Fig. 2H showed 470 471 the similar band number and intensity of proteins from P. fluorescens without (Lane 1) or with BL irradiation (Land 2), implying that such dosage of BL irradiation had no effect on the membrane 472 proteins integrity of *P. fluorescens*. OG at 0.4 mM only caused a slight decrease in the intensity of 473 protein bands (Lane 3). In contrast, a significant decrease of band intensity (especially, for ~63 kDa 474 and ~45 kDa protein) was observed from the sample treated with OG+BL irradiation and a new 475 band (~35 kDa) was formed (Lane 4), highly suggesting that the OG-mediated PDI induced the 476 great damage of membrane proteins and the exogenous photosensitizer (OG) might play a dominant 477 role in eradicating P. fluorescens. DNA is the basis of normal physiological activities and the 478 reproduction of bacteria. Except for the damage to the cell membrane and protein, the effect of 479 OG-mediated PDI on the damage of genomic DNA of *P. fluorescens* is measured and shown in Fig. 480 2I. No significant changes in the band intensity of genomic DNA were observed in the sample 481

treated by BL irradiation alone compared to the control, indicating individual BL irradiation didn't 482 display sufficient damage towards the genomic DNA of P. fluorescens. The band intensity of the 483 group treated with OG or OG+BL was significantly decreased and the treatment of OG+BL almost 484 completely wreck the integrity of genomic DNA of P. fluorescens, which is also consistent with 485 486 both the generation of ROS in Fig. 2B and the antibacterial results in Fig. 1. These discrepancies of damage towards the genomic DNA could be reasonably explained by the difference in the 487 generation of intracellular ROS induced by OG alone and OG+BL. These results are also consistent 488 with the previous study of curcumin-mediated PDI on the genomic DNA of L. monocytogenes 489 (Huang et al., 2020). 490

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[Fig. 2.]

494 3.3. Proposed mechanism of OG-mediated PDI

A scheme of possible bactericidal action mechanisms of OG-mediated PDI is illustrated in Scheme 495 1. OG can interact with the bacteria cell membrane due to its high affinity to the membrane in terms 496 of its hydrophobic portion to exert itself an excellent antimicrobial ability. Then, OG can induce 497 membrane rupture and the leakage of cellular constituent materials, leading to the death of P. 498 fluorescens (I). The internalized OG can further interfere with the activity of the electron transport 499 chain (ETC) on the cytoplasmic membrane, causing increased production of toxic ROS including 500 •OH (II) (Shi et al., 2021). On the other hand, the polyphenolic hydroxyl group of OG would be 501 photo-oxidized by BL. Regarding oxidation, H₂O₂ could be produced through electron transfer from 502 photo-oxidized OG to dissolved oxygen. The H_2O_2 is subsequently photolyzed by BL to generate 503 •OH (III). Consequently, •OH from these two possible pathways would cause lethal oxidative 504 damage to lipid, protein and DNA peroxidation, which further contribute to cellular damage and 505 death. Moreover, OG-mediated PDI can interact simultaneously with multiple targets in the 506 bacterial cells, like the cell membranes, lipids, proteins and DNA, making it difficult for bacteria to 507 the development of antimicrobial resistance (Almeida, Faustino, & Tome, 2015). Thus, the 508 combination of OG and BL irradiation could be considered a low-risk treatment for the 509 development of bacterial resistance or tolerance due to their multitargeted modes of action. 510

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[Scheme 1.]

514 3.4. Inhibitory effect of OG-mediated PDI on P. fluorescens of biofilm formation

P. fluorescens is high prevalence in the fish-processing industry and causes a serious problem due to 515 adhesion to surfaces and biofilm mode of survival under severe conditions (Nilsson, Ross, & 516 Bowman, 2011). Inspired by the highly bactericidal efficient and synergistic effect of OG+BL 517 irradiation, the OG-mediated PDI was also expected to effectively inhibit biofilm formation of P. 518 fluorescens. Before the PDI treatment, the biofilm formation of the P. fluoresens at 30 °C was 519 determined by the crystal violet staining assay, and the results showed that 60 h was the optimal 520 incubation time for the formation of *P. fluoresens* biofilm (Fig. S3). The inhibitory effect of OG+BL 521 irradiation on *P. fluorescens* biofilm formation was presented in Fig. 3. The experiments in which 522 planktonic P. fluorescens were first treated either with (0.05 and 0.1 mM) OG or BL irradiation 523 (94.8 mW/cm²) only, as well as OG+BL for 15 min and then left for 60 h to allow the recovery to 524 form a biofilm, verified that the efficacy of OG+BL irradiation for the inhibition of biofilm 525 formation. In Fig. 3A, when used alone, OG (0.05 mM) or BL irradiation treatment could not cause 526 a significant reduction in the viability of planktonic cells compared to that of the negative control. 527 Cell survival determination indicated that 0.05 mM OG coupled with BL irradiation had a slight 528 effect on the viability of planktonic cells, though, at 0.1 mM, planktonic cells were reduced by only 529 12%. These results indicated that all treatments are performed under sub-lethal conditions have no 530 or minor effect on the activity of P. fluorescens. In Fig. 3B, when samples were treated with BL 531 irradiation alone, no significant change in biofilm formation was observed. Moreover, treatment 532 with OG alone at 0.05 and 0.1 mM could induce a significant reduction in the formation of biofilms 533 by 25% and 57%, respectively. Furthermore, at concentrations of 0.05 and 0.1 mM, OG reduced 534 biofilm formation by approximately 59% and 84%, respectively, with BL irradiation for 15 min. 535 These results confirmed that BL irradiation significantly enhanced the inhibitory effect of OG on P. 536 fluorescens biofilm formation. 537

538 3D confocal scanning microscopy (CLSM) experiments using Live/Dead Baclight assay kit was performed to further elucidate the synergistic antibiofilm effect of OG and BL irradiation. The 539 green-fluorescence nucleic acid stain SYTO 9 could enter into all (live and dead) bacterial cells in 540 biofilms by penetrating. However, PI (red fluorescence) penetrates only dead bacteria with damaged 541 membranes, generally causing a reduction in the SYTO 9 stain fluorescence (Jung, Wen, & Sun, 542 2019). Therefore, living bacteria with an intact membrane stain fluorescence green, whereas dead 543 bacteria stain red when observed via CLSM. As shown in Fig. 3(C), compared with negative control 544 (Fig. 3C-(a)) and positive controls (BL or OG alone) (Fig. 3C-(b), (c)), biofilms treated with 545 546 OG+BL presented a large amount of red fluorescent spots (Fig. 3(C)-(d)-(f)), due to the highly active synergistic effect of OG and BL irradiation. Also, the inhibitory effect of 0.1 mM OG on 547

biofilm formation was similar to those treated by 0.05 mM OG coupled with BL irradiation (Fig. 3C-(d), (e)). These results were in good agreement with the findings in Fig. 3B.

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[Fig. 3.]

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553 3.5. Ablation effect of OG-mediated PDI on preformed biofilms of P. fluorescens

Apart from the excellent capability of inhibiting biofilm formation, the combination of OG and BL 554 irradiation was capable of efficiently eradicating the mature biofilms. In Fig. 4A, approximately 8% 555 and 4% (P<0.05) of biofilms were removed after treatment with 0.4 mM OG alone or BL irradiation, 556 respectively, in comparison to the negative control. However, the performed biofilms submitted to 557 OG-mediated PDI had a significant reduction, which ranged from 32% to 55%, depending on the 558 dosage of BL irradiation. 3D CLSM (Fig. 4B) images confirmed that in the presence of BL 559 irradiation, the eradication effect of OG was significantly enhanced. OG with BL irradiation for 90 560 min showed the highest biofilm-eradicating efficacy. More specifically, the biofilms of the control 561 sample preserved the integrity of biofilm three-dimensional (3D) architectures and extracellular 562 matrices (Fig. 4B-(a)). When evaluating BL irradiation alone and the OG in dark (Fig. 4B-(b), (c)), 563 although a large number of colonies sustained green, some few were stained in red. However, 564 following treatment with OG+BL, many bacterial cells disappeared throughout the biofilms (Fig. 565 4B-(d), (e), and (f)), indicating that the biofilms were very susceptible to OG-mediated PDI 566 treatment. This qualitative assessment was consistent with quantitative data showing the 567 biovolumes of biofilms in Fig. 6(A). There are numerous dead bacterial cells (red fluorescence cells) 568 569 in biofilms treated by the combination of OG and 90 min BL irradiation (Fig. 4B-(f)), confirming that this treatment is substantially effective in eradicating mature bacterial biofilm together, these 570 obtained results clearly indicate the excellent capability of OG+BL irradiation to eradicate existing 571 biofilms and, therefore, leading to cell death. Cell viability in the treated biofilms further confirmed 572 the enhanced effect, as mentioned above. In comparison with the samples treated with either OG or 573 574 BL alone, the significant reduction of the number of viable bacteria in both the supernatant and biofilm caused by the synergistic action of OG and BL irradiation could be observed (Fig. 4C and 575 D). Moreover, the viable count of bacteria was also reduced in an irradiation dosage-dependent 576 manner and OG-mediated PDI for 90 min resulted in 68% and 79% reduction in supernatant and 577 biofilm cells, respectively. Rocha et al. (2020) also found that PDI can reduce the number of 578 colonies in the biofilm of E. faecalis. Our results revealed that OG-mediated PDI not only exhibited 579 a robust inhibition of biofilm production and formation but also had a better effect on eradicating 580

the existing biofilms.

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[Fig. 4.]

584 3.6. Effect of a combination of OG/PLA NFs based packaging and PDI on giant salamander
585 preservation

Electrospun nanofibers loaded with antimicrobial agents are excellent candidates for food 586 packaging materials to actively prevent food contamination during food storage. Following the 587 successful preparation of OG/PLA NFs, we then evaluated its PDI activity against P. fluorescens. 588 The results in Fig. 5A show that the combination of OG/PLA NFs and BL exposure could lead to 589 remarkable reductions in bacterial counts. The situ antibacterial activity of OG/PLA NFs based 590 packaging was evaluated during the Chinese giant salamander storge. Fig. 5B illuminates the total 591 viable count (TVC) of salamander meat during storage after selected treatments. The TVC of 592 control rose rapidly to 9.3 Log CFU/g after 15 days of storage. Conversely, samples packed with 593 OG/PLA NFs with or without BL irradiation have a lower initial bacterial count as described above, 594 and the number of colonies maintained at a low level (<2 Log CFU/g) during the storage especially 595 for the group treated with irradiation. The contaminated salamander meat was packed with OG/PLA 596 NFs under the BL exposure for 30 min, the bacterial count was reduced by ~99% and a great 597 reduction (80%) of viable bacterial counts could be observed as compared to the control after 15 598 days of storage. These findings further confirm that OG plays a dual role as an antibacterial agent 599 and photosensitizer to effectively mitigate foodborne pathogens. Moreover, The PCA region of 600 salamander meat packaged with OG/PLA NFs was the closest to the fresh sample than other ones 601 (Fig. 5C), suggesting that the OG/PLA NFs with BL exposure can exhibit robust antimicrobial 602 efficacy to effectively maintain the meat quality of salamander. These results indicated that 603 OG/PLA NFs as active food packaging material can prevent the contamination of foodborne 604 pathogens and efficiently improve the shelf life of fresh perishable food during storage. 605

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[Fig. 5.]

609 **4. Conclusion**

In summary, this study demonstrated for the first time that the combination of OG at the lower concentration and BL irradiation exerts notable synergistic bactericidal and biofilm ablation effects

against *P. fluorescens*. The dual roles of OG as a promising PS as well as an effective antibacterial 612 agent has been highlighted in the OG-mediated PDI. We also elaborated the mechanism of 613 synergistic bactericidal action by the simultaneous treatment with OG and BL irradiation that might 614 be related to oxidative stress. And the process of ROS generation induced by OG-mediated PDI is 615 616 more complex than originally thought, likely involving a specific interaction between OG and BL as well as between OG and ETC, especially exterior and interior •OH radicals. Furthermore, the 617 electrospun nanofibers of OG/PLA were fabricated with excellent photodynamic antibacterial 618 activity and also effectively reduced bacteria on the surface of the Chinese giant salamander. 619 Therefore, our obtained excellent results also provide insights into the design and fabrication of 620 future alternative antimicrobial and removing/reducing biofilm approaches that could be used in 621 various situations associated with food sanitary. 622

623

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630 Abbreviation

- 631 Photodynamic inactivation, PDI
- 632 Light-emitting diode, LED
- 633 Blue light, BL
- 634 Photosensitizer, PS
- 635 Reactive oxygen species, ROS
- 636 Octyl gallate, OG
- 637 Gallic acid, GA
- 638 Hydrogen peroxide, H₂O₂
- 639 Hydroxyl radicals, •OH
- 640 Superoxide anion $\cdot O_2^-$
- 641 Electron transport chain, ETC
- 642 Total viable count, TVC
- 643 Pseudomonas fluorescens, P. fluorescens
- 644 Staphylococcus aureus, S. aureus

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- 769



Scheme 1. Proposed mechanism of the OG-mediated photodynamic inactivation against *P. fluorescens*. (I) OG damages the membrane of bacterial cells and easily permeates into the intact cells and then (II) disrupts the activity of ETC on the cytoplasmic membrane to generate a high level of toxic ROS. (III) The polyphenolic hydroxyl group of OG would be oxidized by the photoradiation of BL to generate •OH. As a consequence, these ROS-triggered damages lead to microbial cell inactivation and death due to damage to intracellular molecules, such as the essential proteins, DNA, and lipids.





Fig. 1. (A) Evaluate the effect of OG or GA concentration and BL irradiation time on the 783 antimicrobial activity against P. fluorescens in the absence or presence of BL irradiation at 25 °C. 784 The number of bacteria in the suspension was determined by standard plate counting (Wavelength: 785 420 nm, irradiance: 212 mW/cm²). (a) OG concentration-dependent bactericidal activity. (b) The 786 effect of irradiation time on the bactericidal activity of OG-mediated PDI. (B) Time-dependent 787 788 bactericidal activity against (a) P. fluorescens and (b) S. aureus (wavelength: 420 nm, irradiance: 212 mW/cm²). Values and error bars indicate the mean and standard deviation, respectively. 789 Significant differences are shown, P<0.05 (*) and P<0.01 (**). ND: not detected. OG: octyl gallate, 790 GA: gallic acid, OG, NS: normal saline. 791





(B)

(A)













(D)







(F)



(G)



Fig. 2. (A) Measurement of uptake of OG or GA in P. fluoresens using fluorescent-based indicators 799 diphenylboric acid 2-aminoethyl ester (DPBA). P. fluorescens treated by OG or GA solution (0.4 800 mM) in the presence and absence of BL irradiation for 30 min. Absolute fluorescence values were 801 corrected by subtracting the fluorescence values for samples incubated in water and in dark. (B) 802 Total reactive oxygen species (ROS) were detected using DCFH-DA in P. fluorescens treated by 803 OG (0.4 mM) solution in the presence and absence of BL. (C) CLSM images of P. fluorescens 804 incubated with DCFH-DA. Photodynamic inactivation of P. fluorescens was evaluated at 30 °C in 805 presence of OG (0.4 mM) and simultaneously illuminated by BL for 20 min (DIC-differential 806 interference contrast). The initial bacterial density of P. fluorescens is ~8 Log CFU/mL. Pictures 807 were taken using $\times 63$ oil immersion lens with a zoom factor of 2. (**D**) (**a**) Detection of intracellular 808 reactive oxygen species (ROS) using DCFH-DA within P. fluorescens with or without scavengers 809 including CAT (1200 U/mL), DMSO (2.8 M), or TEMPOL (16 mM). (b) Antibacterial assessments 810 of OG alone or OG-mediated PDI with or without CAT (1200 U/mL), DMSO (2.8 M), or TEMPOL 811 (16 mM). The mixture of bacterial suspension and OG solution (0.4 mM) was irradiated with BL 812 for 20 min. (E) Generation of hydroxyl radicals in P. fluorescens treated by OG (0.4 mM) solution 813 in the presence and absence of BL irradiation. Values and error bars indicate the mean and standard 814 deviation, respectively. Significant differences (P < 0.05) between the groups are denoted by 815 different superscript letters. (F) Membrane damage as indicated by the fluorescence level of PI in P. 816 fluorescens treated by OG or GA solution with the final concentration of 0.4 mM in the presence 817 and absence of BL irradiation (wavelength: 420 nm, irradiance: 212 mW/cm², treatment time: 20 818

- 819 min). The initial bacterial density of P. fluorescens is ~8 Log CFU/mL. (G) Scanning electron
- microscopy (SEM) images of *P. fluorescens*. (a), (b) and (c) were SEM images of control, treatment
- by 0.4 mM OG and 0.4 OG+BL for 20 min, respectively. (H) Effects of the OG+BL treatment on
- the total protein of *P. fluorescens* treated by OG (0.4 mM) solution in the presence and absence of
- BL irradiation. (I) Effects of the OG-mediated PDI on the genomic DNA of *P. fluorescens. P.*
- *fluorescens* was treated by OG (0.4 mM) solution in the presence and absence of BL irradiation.
- 825 Values and error bars indicate the mean and standard deviation, respectively. Significant differences
- 826 (P < 0.05) between the groups are denoted by different superscript letters.





(**A**)







Fig. 3. Effect of OG coupled with BL irradiation on biofilm formation. (A) Surviving planktonic 832 cells in the culture supernatant treated with OG in the absence or presence of BL irradiation for 15 833 min (wavelength 420 nm, irradiance: 94.8 mW/cm²). (B) Relative biofilm formation for 60 h and (C) 834 3D CLSM images of *P. fluorescens* biofilms after different conditions. (a) The untreated control 835 was further incubated for 60 h without OG and BL irradiation. (b) P. fluoresens were treated with 836 BL irradiation alone for 15 min and further incubated for 60 h. (c) P. fluoresens were treated with 837 0.05 mM or (d) 0.1 mM OG alone for 15 min and further incubated for 60 h. (e) P. fluoresens were 838 treated with 0.05 mM OG or 0.1 mM OG (f) combined with BL irradiation for 15 min and further 839 incubated for 60 h. The first and second columns display bacteria labeled with LIVE/DEAD 840 BacLight SYTO 9 (green) and PI (red) and the third column displays the merged pictures. Values 841 and error bars indicate the mean and standard deviation, respectively. Significant differences 842 (P < 0.05) between the groups are denoted by different superscript letters. 843

844




	SYTO 9	PI	Merged	
(a) OG(-)BL(-) Control	20 02 02 02 0 0 0 0 0 0 0 0 0 0 0 0 0	0 0 0 2 0 4 0 0 0 0 0 0 0 0 0 0 0 0 0 0	a^{0} a^{0} a^{0} a^{0} a^{0} a^{0} b^{0} a^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} a^{0} b^{0} b^{0} a^{0} b^{0	
(b) OG(-)BL(+)	$\begin{array}{c} 0.0\\ 0.2\\ 0.4\\ y (mm) \\ 0.6\\ 0.9\\ 0.9\\ 1\\ 1\\ 0.0\\ 0\\ 1\\ 1\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\ 0\\$	$y_{(mm)} = 0.000 + 0.00000 + 0.000000 + 0.000000 + 0.00000 + 0.0000 + 0.0000 + 0.0$	$\begin{array}{c} 0.0\\ 0.0\\ 0.2\\ 0.4\\ y(mm) \\ 0.6\\ 0.0\\ 0.9\\ 1.0\\ 0.0\\ 1.0\\ 0.0\\ 0 \\ 1.0\\ 0.0\\ 0 \\ 1.0\\ 0.0\\ 0$	
(c) OG(+)BL(-)	$ \begin{array}{c} & & & & & & & & & & & & & & & & & & &$	00 00 01 02 04 05 04 05 05 05 05 05 05 05 05 05 05	00 00 00 00 00 00 00 00 00 00 00 00 00	
(d) OG(+)BL(+) 30 min	y (mm) 0.3 1 1 0 0 0 0 0 0 0 0 0 0 0 0 0	00 00 00 00 00 00 00 00 00 00 00 00 00	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	



851 Fig. 4. Effect of OG+BL on preformed biofilms (wavelength 420 nm, irradiance: 212 mW/cm²). (A) Relative biofilm. (B) 3D CLSM images of *P. fluoresens* biofilms after different treatments. (a) The 852 853 untreated control constituted a 60 h-old biofilm further cultivated for 60 min without OG and BL irradiation. (b) *P. fluoresens* 60 h-old biofilms were treated with BL irradiation alone for 60 min. (c) 854 P. fluoresens 60 h-old biofilms were treated with 0.4 mM OG for 60 min. (d) P. fluoresens 60 h-old 855 biofilms were treated with 0.4 mM OG coupled with BL irradiation for different exposure times (d) 856 30 min, (e) 60 min, and (f) 90 min. The first and second columns display bacteria labeled with 857 LIVE/DEAD BacLight SYTO 9 (green) and PI (red) and the third column displays the merged 858 pictures. (C) Quantitative analysis of surviving cells in biofilms or (D) in the culture supernatant. 859 Values and error bars indicate the mean and standard deviation, respectively. Significant differences 860 (P < 0.05) between the groups are denoted by different superscript letters. 861 862



Fig. 5. The changes in preservation quality indices of Chinese giant salamander during storage at 4 865 °C. (A) Results of bactericidal assessments of OG/PLA NFs combined with BL irradiation against 3 866 Log CFU/mL bacteria. ND: not detected. (B) The change of colonies of Chinese giant salamander 867 during the storage for 3, 6, 9, 12 and 15 days at 4 °C. The fresh salamander meat was packed using 868 OG/PLA NFs and then immediately treated with BL irradiation for 30 min. (C) The PCA of Chinese 869 giant salamander with different treatments: (a) fresh salamander meat; (b) control; (c) control+BL; 870 (d) OG/PLA NFs; (e) OG/PLA+BL; (b-e, 4 °C, 15 days). The salamander meats were singly sealed 871 with OG/PLA NFs and irradiated immediately with BL for 30 min at a distance of 1 cm. Significant 872 differences (P < 0.05) between the groups are denoted by different letters. 873 874

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SURPPORTING INFORMATION

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1. Method

31 *1.1. Quantification of phenolics adsorption through CLSM*

Quantification of phenolics (OG or GA) adsorption was conducted according to the method 32 reported by Wang et al. (2017) with some modifications. The P. fluorescens cells were cultured in 33 LB broth at 30 °C for 14-16 h. The bacterial cells were harvested and washed three times with PBS 34 (0.1 M, pH=7.2). Then the bacterial cells were resuspended in PBS to 8 Log CFU/mL and phenolics 35 were added to the resuspension solution to a final concentration of 0.1 mM. Cells were incubated at 36 30 °C for 30 min and then harvested. For the quantification of phenolics adsorption, 450 µL of 37 DPBA solution (DMSO, 0.2% w/v) was added to the above bacterial cells and then incubated for 5 38 min. The images of the cells were captured using confocal laser scanning microscopy (CLSM, 39 Leica TCS SP8, Germany). 40

41

42 1.2. Hydroxyl radical experiments using flow cytometry

Flow cytometry was used to detect hydroxyl radical formation. Hydroxyphenyl-fluorescein (HPF) is 43 an anthracene derivative of fluorescein which becomes fluorescent when it was activated by 44 hydroxyl radical (Shi et al., 2021). The bacterial culture with 8 Log CFU/mL P. fluorescens was 45 treated with or without OG under BL exposure for 20 min. Samples without irradiation were chosen 46 to be controlled. After various treatments, fluorescent reporter dye 3'-(p-hydroxyphenyl) fluorescein 47 (HPF, Invitrogen) was added into the solution to reach a final concentration of 10 µM, and the 48 mixture was incubated at 37 °C for 30 min. Then, samples were centrifuged at 10000×g for 5 min 49 and washed twice in PBS to remove excess dye. The cell pellet was resuspended in PBS for the 50 following measurement. For visualization of intracellular •OH in P. fluorescence, the fluorescent 51 signal ($\lambda ex/\lambda em = 490/515$ nm) was recorded by CytoFLEX flow cytometry (Beckman Coulter, 52 USA). Calibrite beads (Becton Dickinson) were used for instrument calibration. Flow data were 53 processed and analyzed with MATLAB (The MathWorks). 54

55 *1.3. Investigation of the cell membrane damage*

56 To evaluate the cell membrane damage induced by OG+BL treatment, the fluorescence probe

propidium iodide (PI) and scanning electron microscope (SEM) was performed (Shi et al., 2020; 57 2021). In brief, a sample containing 8 Log CFU/mL P. fluorescens and either 0 or 0.4 mM OG was 58 exposed to BL for 20 min. Cells were then harvested from 1 mL volume of the incubated sample by 59 centrifuging at 10000×g at room temperature for 5 min twice with an intermediate washing step 60 using sterile DI water. The pellet was then re-suspended in 10 µM PI solution (1 mg/mL) following 61 dark incubation at room temperature for 15 min. Subsequently, the incubated solution was washed 62 with PBS and centrifuged for 2 min at 10,000×g. The pellet was resuspended in 500 mL PBS, and a 63 100 µL of each solution was transferred into a 96 well plate. The fluorescence spectrum was 64 measured at excitation wavelengths of 495 with emission range from 500 to 800 nm with a 65 microplate reader (Synergy H1 Multi-Mode Reader, BioTek, Winooski, VT, USA). Bacterial 66 suspension with water kept in the dark was used as a control, and its fluorescence intensity reading 67 was used as the reference to calculate the relative fluorescence unit (RFU) for other treatments 68 using the following equation: 69

70

$I_{corrected} = I_s - I_0$

where I_s stands for the fluorescence emission intensity reading of the sample received treatments and I_o stands for the signal intensity of the control sample.

73

74 Scanning electron microscopy (SEM) was conducted to analyze the damage to the cell membrane caused by the OG+BL treatment. Briefly, the cultured samples containing 8 Log CFU/mL of P. 75 76 fluorescens were treated with BL for 20 min. Controls consisted of bacterial suspension without OG in the dark and only exposure to OG. Following treatment, 1 mL bacterial suspension was 77 78 centrifuged for 5 min at 6000 rpm and washed with PBS, then, the samples were resuspended with 2.5% glutaraldehyde for overnight at 4 °C. Finally, the samples were dehydrated sequentially in 25, 79 80 50, 75, 95% and 100% (v/v) aqueous solutions of ethanol following washed with PBS. Cells were placed on a SEM stub and coated with gold to observe the morphology with a Hitachi T-1000 81 scanning electron microscope (Hitachi High-Technologies Corporation, Tokyo, Japan) an 82 acceleration voltage of 15 kV. 83

84

85 *1.4. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE)*

SDS-PAGE analysis was performed to determine the alternations of bacterial proteins before and after the treatment of OG and BL irradiation. OG-mediated PDI treated *P. fluorescens* (8 Log CFU/mL) was collected by centrifugation (10000×g, 4 °C) for 5 min, and then washed 3 times using PBS (0.1 M, pH 7.2). The proteins were obtained using kits (BestBio, Shanghai, China) and then mixed with loading buffer at 100 °C for 10 min. Subsequently, the Marker (Takaba, Dalian,
China) and proteins were run through the 12% separating gel and 3% stacking gel at 100V for 90
min. The gel was stained using Coomassie brilliant blue (R-250) stain solution for 4 h. Afterward,
the gel was destained with a common decoloring agent.

94

95 1.5. Genome integrity determination

The damage of DNA caused by OG+BL was investigated by the agarose gel electrophoresis. Briefly, the OG+BL treated *P. fluorescens* (~8 Log CFU/mL) was centrifugated at 6000×g for 5 min, and the bacterial DNA was extracted using a DNA extraction kit (Sangon, Shanghai). The samples were separated by 1% agarose. The parameters of the DNA electrophoresis instrument were set to a voltage of 100 V for 30 mins. The gel images of samples were observed and recorded on a ChemiDoc MP Imaging System (Bio-Rad, Hercules, CA, USA).

102

103 *1.6. Confocal laser scanning microscopy (CLSM)*

To observe the formation of the biofilm after various treatments, samples were captured by CLSM 104 refer to Seo and Kang (2020) with some modifications. P. fluorescence was treated as described 105 above, Then, bacteria were cultured in a Confocal Quartz Petri Dish at 30 °C for 60 h, after 106 cultivation, planktonic cells were removed by gently washing the dishes with 0.9% sterile normal 107 saline. For visual observation of the performed biofilms after different treatments, P. fluorescence 108 was cultured in a Confocal Quartz Petri Dish using the method above, and then planktonic cells 109 were removed by gently washing the dishes with PBS, after that, preformed biofilms were treated 110 with different conditions. Treated biofilm was then stained using a LIVE/DEAD Baclight Bacterial 111 Viability Kits (Molecular Probes, Invitrogen, USA) according to the manufacturer's instructions. 112 The biofilm was washed with PBS and visualized using a CLSM. The 20×objective was used to 113 monitor STYO 9 fluorescence excited at 488 nm and emitted at 493-580 nm, propidium iodide 114 fluorescence excited at 552 nm and emitted at 566-719 nm. 115

116

117 *1.7. Preparation of nanofibers (NFs)*

The NFs were prepared using the method reported by us (Shi et al., 2020) with some modifications.
Briefly, the solutions used for electrospinning were obtained by dissolving poly(lactic acid) (PLA)
in hexafluoroisopropanol (HFIP) (9%, *w/v*, 0.9 g of PLA in 10 mL HFIP) at room temperature under

stirring for 3 h. The OG/ β -cyclodextrin inclusion complexes (OG/ β CD, 6% and 8%, w/v) were 121 added to the solution with magnetically stirring for 12 h at room temperature to obtain the 122 homogeneous solution for electrospinning. The prepared solution was put into a 10 mL 123 polypropylene syringe fitted with a 23G (outer/inner diameter; 0.64 mm/0.33 mm) metallic needle. 124 The solution flow rate was 0.9 mL h⁻¹. The loaded syringe was fixed horizontally with a syringe 125 pump (Baoding longer, LSP03-1A) and the electrode of the high voltage power supply (Tianjin 126 Dongwen, DWLP303-1ACDB) was connected to the metal needle tip. The working distance 127 between the needle tip and the ground electrode was 10 cm. The electrospinning voltage was 18 kV. 128 The electrospinning temperature and the relative humidities were 25 °C and 45%, respectively. The 129 collected fibers were vacuum dried for 24 h to remove solvent residue and their morphology was 130 observed via SEM (Hitachi T-1000, Hitachi High-Technologies Corporation, Tokyo, Japan) (see Fig. 131 S1) and the preservation of giant salamander (Fig. S4). 132

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134 1.8. In situ antibacterial activity of OG/PLA NFs

To assess the synergistic effect of OG/PLA NFs and PDI on the preservation of Chinese giant 135 salamander, the changes in the total viable count and flavor of samples during storage were studied. 136 The salamander meat cubes (4 cm×4 cm×1 cm, 10 g) were cut under aseptic conditions, then 137 inoculated by immersion in a bacterial suspension of P. fluorescens (3 Log CFU/mL) for 30 seconds 138 and then all samples were dried at room temperature for 10 min and the treated salamander meats 139 were singly sealed with OG/PLA NFs and followed immediately by BL exposure (Radiation 140 intensity: 213 mW/cm²) for 30 min at a distance of 1 cm. The bacteria were collected from the 141 surface of meat using PBS (0.1 M, pH 7.2). The viable cell count was determined by placing them 142 on LB agar plates after serial dilution. To further investigate the effect of OG/PLA NFs combined 143 with PDI on the preservation of Chinese giant salamander, the changes in the total viable count and 144 flavor of samples during the storage were studied. All samples were stored at 4 °C for 15 days. The 145 colonies were counted at intervals of 3 days. Meanwhile, to evaluate the quality of the giant 146 salamander with different treatments during the storage, the electronic nose was applied to analyze 147 the flavor on the salamander from batches control, control+BL, OG/PLA NFs and OG/PLA 148 NFs+BL at 4 °C for 15 days (for more details was given in SI-Section 1.2). Each group was 149 150 analyzed three times in parallel.

The flavor analysis using an electronic nose was also performed on salamander from batches 153 control, control+UV-A, OG/HPBCD NFs and OG/HPBCD NFs+UV-A at 4 °C for 15 days. Five 154 repetitions were performed for each group. The electronic nose (Figure S1) applied in the 155 experiment was also developed by us. It consists of a sampling system, a detector containing the 156 157 array of sensors, and pattern recognition for data recording. The sensor array is composed of fourteen different metal oxide sensors. Each sensor has a certain degree of affinity toward specific 158 chemicals or volatile compounds. The sensors' response to changes in conductivity induced by the 159 adsorption of gas-phase molecules and on subsequent surface reactions. Before the measurement, 160 the system of the electronic nose was cleaned with zero-air which was indoor air-filtered by active 161 carbon. The main purpose of zero-air was to clean the circuit and to return the sensors to the 162 baselines. During the measurement of icefish, the headspace gas of a sample was pumped into the 163 sensor chamber at a constant rate of 0.6 L/min through a tube connected to a needle. The response 164 of each sensor was expressed as a ratio of conductance (G/G0, G and G0 are conductances of the 165 sensors' response to the sample gas and the zero-air, respectively). The measurement procedure was 166 controlled by a special program. The measurement time was 160 s, which was sufficient for each 167 sensor to reach a stable value. The cleaning time was set to 100 s. The data was stored after the 168 measurement was completed. The same sample was paralleled 5 times. 169

170

172 **2. Results**

173

174

Table S1. MIC and MBC of GA or OG against *P. fluoresens* and *S. aureus*.

HO	R	P. fluoresens (G ⁻)		S. aureus (G ⁺)		$c \text{Log } P^a$	
но			MIC (mM)	MBC (mM)	MIC (mM)	MBC (mM)	6
Gallic acid	GA	Н	6.4	6.4	3.2	>6.4	0.4254
Octyl Gallate	OG	$C_{8}H_{17}$	0.1	3.2	0.05	0.1	4.6344

^a Theoretical estimate using ChemBioDraw Ultra 13.0 program. Hydrophobicity of GA and OG from their partition coefficient (Log*P*) analysis. Log*P* is defined as the decadic logarithm of the particular ratio of the concentration of a compound between the two solvents (octanol phase and water phase).

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- 180





Fig. S1. SEM images PLA-OG/ β CD nanofibers.

The morphological analysis of the composite electrospinning nanofibers containing OG was carried out using Hitachi T-1000 scanning electron microscope (Hitachi High-Technologies Corporation, Tokyo, Japan), with an acceleration voltage of 15 kV, as shown in Fig. S1. The composite nanofibers were set on a metallic stub and covered with gold under vacuum in an argon atmosphere.

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-
- 190







Fig. S2. Confocal laser scanning microscopy images of GA or OG in *P. fluorescens* stained with
 DPBA.







Fig. S4. Physical appearance of Chinese giant salamander treated with (a) control ; (b) BL; (c) OG/PLA NFs; (d) OG/PLA NFs + BL (a-d: 4 °C, 15days)

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Declaration of interest statement

We declare that we have no financial and personal relationships with other people or organizations that can inappropriately influence our work, there is no professional or other personal interest of any nature or kind in any product, service and/or company that could be construed as influencing the position presented in, or the review of, the manuscript entitled.