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6 **Photoelectrocatalytic disinfection of water and wastewater: performance evaluation by qPCR**
7 **and culture techniques**

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18 **Abstract**

19 Photoelectrocatalytic oxidation (PEC) was evaluated as a disinfection technique using water and
20 secondary treated wastewater spiked with *Escherichia coli* and *Enterococcus faecalis*. PEC
21 experiments were carried out using a TiO₂/Ti-film anode and a zirconium cathode under simulated
22 solar radiation. Bacterial inactivation was monitored by culture and quantitative PCR (qPCR).
23 Inactivation was enhanced increasing the duration of the treatment and decreasing bacterial
24 population and the complexity of the aqueous matrix. *E. coli* cells were reduced approximately 6
25 orders of magnitude after 15min of PEC treatment in water at 2V of applied potential and an initial
26 concentration of 10⁷ CFU/mL; pure photocatalysis led to about 5 log reduction, while
27 electrochemical oxidation alone resulted in negligible inactivation. The superiority of PEC relative
28 to PC can be attributed to a more efficient separation of the photogenerated charge carriers.
29 Regarding disinfection in mixed bacterial suspensions, *E.coli* was more susceptible than *E.faecalis*
30 at a potential of 2V. The complex composition of wastewater affected disinfection efficiency,
31 yielding lower inactivation rates compared to water treatment. qPCR yielded lower inactivation
32 rates at longer treatment times than culture techniques presumably due to the fact that the latter do
33 not take into account the viable but not culturable state of microorganisms.

34
35 **Keywords:** *E. coli* · *E. faecalis* · Disinfection · qPCR · TiO₂ · Voltage

36 INTRODUCTION

37 The growing demand for clean water and wastewater, free from disease-causing pathogenic
38 microorganisms, poses new challenges in the development of effective disinfection techniques.
39 Over the last decades, advanced oxidation processes (AOPs) have been recognised as an emerging
40 group of techniques, highly effective for the decomposition and mineralization of organic
41 compounds in aqueous samples (Malato *et al.* 2009; Chong *et al.* 2010). TiO₂ photocatalysis is an
42 important member of AOPs and its benefits regarding water and wastewater disinfection have been
43 demonstrated with respect to *Escherichia coli*, *Staphylococcus aureus* and *Enterococcus faecalis*
44 (Chen *et al.* 2010; Chatzisymeon *et al.* 2011; Rémy *et al.* 2011; Venieri *et al.* 2011). In most
45 photocatalytic applications the catalyst is employed as slurry of fine particles in a photochemical
46 reactor, resulting in certain difficulties, such as post-reaction catalyst recovery and low quantum
47 efficiencies (Koivunen & Heinonen-Tanski 2005; Egerton *et al.* 2006; Marugán *et al.* 2006). These
48 problems may be addressed by immobilization of TiO₂ on a conducting support and application of a
49 potential bias, so as to reduce the recombination of charge carriers, which is the main limitation of
50 the process photonic efficiency (Marugán *et al.* 2009; Egerton 2011).

51 These modifications have led to the development of photoelectrocatalysis (PEC), which
52 consists of the application of a constant bias anodic potential usually to a TiO₂-based thin film
53 acting as a photoanode. The photoinduced electrons are then continuously extracted from the anode
54 by an external electrical circuit to be injected into the cathode (Sirés & Brillas 2011). The constant
55 current density or bias potential applied to the semiconductor electrode promotes the efficient
56 separation of electron-hole pairs and accelerates the production of photogenerated oxidizing species
57 onto the catalyst surface (Martínez-Huitle & Brillas 2009). In this view, it is not surprising that PEC
58 has recently found a wide range of environmental applications, including degradation of organic
59 pollutants and pharmaceuticals, detoxification of aqueous samples, wastewater remediation and
60 microbial inactivation (Fraga *et al.* 2009; Martínez-Huitle & Brillas 2009; Nissen *et al.* 2009;
61 Egerton 2011; Frontistis *et al.* 2011; Li *et al.* 2011; Sirés & Brillas 2011).

62 PEC seems to be a promising and efficient tool, regarding the effective inactivation of
63 pathogens which are contained in water and wastewater, contributing in the control of waterborne
64 diseases. The bactericidal function of this technique can be attributed to the oxidation properties of
65 photocatalytically generated active oxygen species (AOS), which cause damage to cellular
66 membrane and further destruction of bacterial structure (Li *et al.* 2011). The studies referred to PEC
67 as means of disinfection highlight the importance of certain parameters like applied voltage,
68 bacterial concentration, treatment time and the aqueous matrix, which are considered determining
69 factors of microbial inactivation (Martínez-Huitle & Brillas 2009). Generally, the overall evaluation
70 of PEC, as a method for disinfection, is mainly performed with the use of a faecal bacterial
71 indicator, applying conventional microbiological protocols, based colony-forming units (CFU)
72 counts on selective culture media (Egerton *et al.* 2006; Philippidis *et al.* 2010; Egerton 2011; Li *et*
73 *al.* 2011). Yet, the environment of PEC treatment may be considered rather stressful for microbial
74 populations, inducing their entrance to the viable but not culturable (VBNC) state. Within this state,
75 bacteria demonstrate metabolic activity and maintain their pathogenic features but they are not
76 recoverable in culture media, leading to false negative results (Muela *et al.* 2008). The application
77 of molecular methods surpasses this limitation, based on nucleic acid identification and
78 quantification. Among them quantitative PCR (qPCR) has proven to be sensitive yielding accurate
79 quantitative results and allowing for new approaches in waterborne pathogen research (Lee *et al.*
80 2006; Shannon *et al.* 2007).

81 The aim of this work was to investigate the inactivation of faecal indicators in water and
82 wastewater by means of photoelectrocatalytic oxidation, applying conventional cultural methods
83 and qPCR. This was implemented using representative strains of faecal indicator bacteria, namely
84 *Escherichia coli* and *Enterococcus faecalis*. The effect of operational conditions such as bacterial
85 concentration, potential, treatment time, and aqueous matrix were also examined.

86

87 MATERIALS AND METHODS

88

89 **Bacterial strains and wastewater**

90 The bacterial strains used as reference faecal indicators in the present study were Gram-
91 negative *E. coli* ATCC 23716 and Gram-positive *E. faecalis* ATCC 14506 (American Type Culture
92 Collection, Rockville, Md. USA). Suspensions of bacterial indicators were prepared in sterile
93 distilled water, which was used as sample for water disinfection experiments. The concentration of
94 bacterial cells in the suspension was estimated measuring its optical density at 600 nm (Shimadzu
95 UV1240 spectrophotometer). The bacterial concentrations used in this study were within the range
96 of 10^4 - 10^8 CFU/mL. Plate counts were also performed for accurate bacterial count.

97 Wastewater disinfection experiments were carried out in real wastewater collected from the
98 outlet of the activated sludge unit (prior to chlorination) of the municipal wastewater treatment plant
99 of Chania, W. Crete, Greece. Its main characteristics were determined according to Standard
100 Methods (1999) as follows: the chemical oxygen demand and dissolved organic carbon were 26 and
101 7.8 mg/L, respectively, the concentration of chlorides, sulfates, nitrates, nitrites, bicarbonates and
102 total solids were 222.1, 60.3, 25.9, 57.1, 182.1 and 7 mg/L, respectively, while the pH was 7.8.
103 Sterilization of the sample was performed prior to any photoelectrocatalytic treatment in order to
104 adjust the initial bacterial concentration at the desired level.

105

106 **Disinfection experiments**

107 Photocatalytic experiments were performed using a solar radiation simulator (Newport,
108 model 96000) equipped with a 150 W xenon ozone-free lamp and an Air Mass 1.5 Global Filter
109 (Newport, model 81094), simulating solar radiation reaching the surface of the earth at a zenith
110 angle of 48.2°. According to the spectral irradiance data given by the manufacturer, simulated solar
111 radiation contains about 5% UV-A radiation, and 0.1% UV-B radiation, while the filter cuts
112 radiations with wavelengths lower than 280 nm. The incident radiation intensity on the

113 photochemical reactor in the UV region of the electromagnetic spectrum was measured
114 actinometrically using 2-nitrobenzaldehyde (Sigma-Aldrich) as the chemical actinometer (Willett &
115 Hites 2000; Galbavy *et al.* 2010) and it was found to be 5.8×10^{-7} einstein/(L×s), which corresponds
116 to an irradiance of 1.31×10^{-2} W/m². Reactions took place in an open, double-walled, cylindrical
117 glass vessel under continuous stirring.

118 Constant potentials (2 and 5V) were applied between the TiO₂/Ti anode and the zirconium
119 cathode using a galvanostat-potentiostat (Amel Instruments, model 2053). Details concerning the
120 experimental setup, as well as the preparation and characterization of the TiO₂/Ti-film are given
121 elsewhere (Frontistis *et al.* 2011). In brief, the anode surface was 5.8 cm² and had 75:25
122 anatase:rutile phase composition. The reactant mixture had a liquid holdup of 60 mL. The reaction
123 temperature was maintained constant at 25±1°C with a temperature control unit. All experiments
124 were conducted in the presence of 1% (w/v) Na₂SO₄ aqueous solution serving as the electrolyte.

125 Water samples were processed applying (a) photoelectrocatalytic treatment, (b)
126 photocatalytic treatment (PC: solar radiation alone) and (c) electrochemical oxidation (EO:
127 application of voltage only in dark conditions). Photoelectrocatalytic treatment was also applied for
128 disinfection of wastewater samples. All disinfection experiments were performed in triplicate. In a
129 typical experiment, the inoculated aqueous solution was loaded to the cell, left in the dark under
130 stirring for 20 min in order to equilibrate and then exposed to solar radiation and/or potential. At
131 specific time intervals about 3 mL of the reaction solution were withdrawn. Half of the quantity was
132 immediately analyzed with respect to viable bacterial cells applying conventional culture method
133 and the rest was used for DNA extraction and PCR amplification.

134

135 **Culture method**

136 The detection and enumeration of *E. coli* and *E. faecalis* in the solution were performed
137 using the serial dilution streak plate procedure. The media used in the study were HiCrome
138 Coliform Agar (HiMedia Laboratories) and Slanetz & Bartley medium (OXOID) for *E. coli* and *E.*

139 *faecalis*, respectively. Incubation took place at 37°C. Bacterial counts were performed after 20-24 h
140 and 48 h for *E. coli* and *E. faecalis* determination, respectively.

141

142 **DNA extraction**

143 Genomic DNA was extracted performing chemical lysis and phenol/chloroform/isoamyl
144 alcohol (25:24:1) extraction. Namely, the cells were spun for 2 min and were lysed for 1 h at 37°C
145 with 300 µL of lysozyme lysis buffer (100 mM NaCl, 500 mM Tris [pH 8], lysozyme 10 mg/mL)
146 and 3 µL of 20 mg/ml proteinase K. Then, 200 µL of SDS lysis buffer (100 mM NaCl, 500 mM Tris
147 [pH 8], 10% [w/v] SDS) were added, followed by incubation at 65°C for 10 min. The solution was
148 extracted with 750 µL of chloroform/isoamyl alcohol (24:1), spun, and the aqueous phase was re-
149 extracted with phenol/chloroform/isoamyl alcohol (25:24:1). Ethanol purification step was
150 performed and the quantity and purity of all DNA samples were determined measuring their
151 absorbance at 260 nm and estimating the ratio of absorbance values at 260 nm and 280 nm,
152 respectively.

153

154 **Monitoring of *E. coli* and *E. faecalis* through qPCR**

155 The SYBR green method was chosen for the quantification of both bacterial indicators using
156 the StepOne Plus System (Applied Biosystems Inc., Foster City, CA, USA). The *gadAB* gene was
157 used as a target for *E. coli* quantification through qPCR, and the primers were as follows: forward
158 primer 5'-GCG TTG CGT AAA TAT GGT TGC CGA-3' (*gadrt-1*) and reverse primer 5'-CGT
159 CAC AGG CTT CAA TCA TGC GTT-3' (*gadrt-2*) (Chen *et al.* 2006). The product size for this
160 primer set is 305 bp. The set of primers for *E. faecalis* detection and quantification was designed
161 according to the sequences of 16S rRNA, which are available in GenBank (Bartosch *et al.* 2004).
162 The primer pair contained a forward primer 5'-AACCTACCCATCAGAGGG -3' (Efs130F) and a
163 reverse primer 5'-GACGTTTCAGTTACTAACG -3' (Efs490R), which yield a 360 bp PCR product.
164 Triplicate PCR reactions were carried out with Quantimix Easy SYG Kit (Biotools) to a final

165 volume of 20 μ L. The mixed qPCR solution contained 2XPCR master mix, 0.5 mM of each primer,
166 30 nM of reference dye (Rox) all diluted to the final volume of the reaction mixture with
167 DNase/RNase free water. PCR reactions were carried out at a temperature profile of 10 min initial
168 denaturation at 95°C, followed by 40 cycles each of denaturation at 95°C for 0.5 min, annealing at
169 57°C for 1 min, and extension at 72°C for 0.5 min. Deionized water and DNase-treated *E. faecalis*
170 and *E. coli* served as negative controls. Also, melt curve analyses were conducted from 55 to 95°C.

171 To determine the detection sensitivity of the qPCR assay, a series of 10-fold diluted pure
172 culture genomic DNA of each bacterium was tested for qPCR amplification and cycle threshold
173 (C_T). Standard curves were generated with *E. coli* and *E. faecalis* DNA.

174

175 **RESULTS AND DISCUSSION**

176

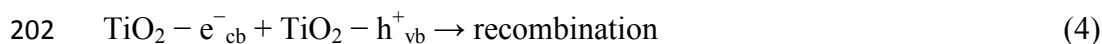
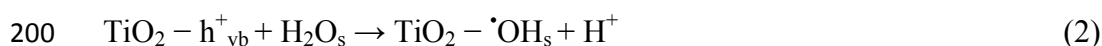
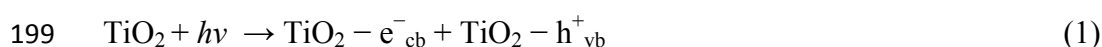
177 **Efficiency of photoelectrocatalysis in relation to photocatalysis and electrochemical oxidation**

178 The decrease of bacterial population in water samples was investigated under three distinct
179 conditions: (a) photoelectrocatalytic treatment (PEC: simultaneous application of solar radiation and
180 2V bias potential); (b) photocatalytic treatment (PC: solar radiation alone); (c) electrochemical
181 oxidation (EO: application of voltage only in dark conditions).

182 The results, summarized in Fig. 1, show that the application of a 2V potential during PEC increased
183 the extent of disinfection compared to PC. For instance, PEC led to a 5.9 log reduction after 15 min,
184 with the respective values of PC and EO being 4.9 and 0.2. However, both PEC and PC resulted in
185 an approximately 7 log reduction after 120 min of treatment, while in the absence of radiation
186 bacterial inactivation was inadequate (i.e. 1.5 log reduction); this clearly shows that EO is not a
187 suitable disinfection process at the conditions under consideration as it is not able to achieve total *E.*
188 *coli* inactivation in aqueous suspensions. Control runs were also performed in the absence of
189 radiation and potential showing no change in bacterial population after 120 min of contact time
190 (data not shown).

191 Illumination of a semiconductor-electrolyte interface with photons having energy greater
 192 than its band gap energy generates electron-hole pairs at the anode electrode surface. The
 193 simultaneous application of a bias positive to the flat-band potential produces a bending of the
 194 conduction and valence bands which, in turn, causes a more effective separation of the
 195 photogenerated carriers within the space charge layer (Morrison 1980). The potential gradient
 196 forces the electrons towards the cathode, thus leaving the photogenerated holes to react at the anode
 197 with H₂O and/or OH⁻ to yield hydroxyl radicals, i.e.:

198 *Anode (working electrode):*



203 *Cathode (counter electrode):*



205 where the subscripts cb and vb denote the conduction and valence bands, respectively, h⁺ and e⁻
 206 denote the photogenerated holes and electrons, respectively, while the subscript s refers to species
 207 adsorbed onto the photoanode surface.

208 Solar radiation has the potential to inactivate bacterial cells. Moreover, in the presence of a
 209 catalyst, disinfection is mainly achieved by the action of oxidative radicals released from irradiated
 210 TiO₂ (Chen *et al.* 2010). PEC under sunlight radiation has better performance than PC since the
 211 application of potential is believed to suppress the rate of electron-hole recombination, thus
 212 enhancing photocatalytic rates (Baram *et al.* 2009). The holes which have been left by electrons are
 213 accelerated into the semiconductor surface, where they are available to react with organisms in the
 214 water. The obtained results are in accordance to other studies which highlight PEC as an efficient
 215 disinfection method, in terms of inactivating mainly *E. coli* and Gram-negative bacteria (Philippidis
 216 *et al.* 2010; Rahmawati *et al.* 2011). Generally, the benefits of electric field enhancement have been

217 demonstrated not only for coliforms but also for *Clostridium perfringens* spores and the recalcitrant
218 *Cryptosporidium parvum* (Egerton *et al.* 2006).

219

220 **Effect of initial bacterial concentration**

221 The influence of the initial *E. coli* concentration on PEC disinfection ability is presented in
222 Fig. 2. Conventional culture technique (Fig. 2a) showed that the rate of PEC disinfection was
223 inversely proportional to *E. coli* concentration in water samples. Total inactivation was achieved in
224 relatively short treatment time (i.e. within approximately 15 min) only when bacterial inoculum
225 contained 10^5 CFU/mL, while at higher concentrations residual *E. coli* cells were determined even
226 after 60 min. For instance and at an initial concentration of 10^8 CFU/mL, a 5.5 log reduction was
227 achieved within 30-60 min of treatment, beyond which no further inactivation was recorded.

228 SYBR green qPCR was performed to detect the potential presence of live non-culturable
229 bacterial cells. Our effort was to evaluate the consistency of *E. coli* cells and taking into account
230 that cultivability is not synonymous of viability (Rémy *et al.* 2011). qPCR showed different periods
231 of microbial inactivation (Fig. 2b) compared to plate counts. Inactivation rates were quite similar
232 when initial genome copies were of 10^7 and 10^5 /mL, as there was no enhancement of the process
233 after 40 min of treatment. During PEC, cell injury and loss of viability are achieved as cellular
234 membrane is destructed through direct contact with the photoanode (Li *et al.* 2011). Furthermore,
235 radiation induces DNA lesions, damaging nucleic acids and making them functionless. The
236 hydroxyl radical (and other AOS), directly generated by this process, is the main cause of DNA
237 destruction, which in turn leads to cell death (Sinha & Häder 2002; Gogniat & Dukan 2007).
238 According to our results, even after 120 min of treatment, genome copies were detected and
239 quantified in the water sample, reaching a plateau without any further decrease. This profile of
240 genome copies in the reaction mixture could be attributed to the detection sensitivity of the method.
241 qPCR is acknowledged as a reliable and sensitive molecular method, whose detection limit may be
242 in the region of 100 fg of *E. coli* genomic DNA (Lleo *et al.* 2005; Shannon *et al.* 2007). However,

243 in the present study environmental samples were processed which may inhibit qPCR or restrict its
244 detection limits. On the other hand, qPCR seems to be valuable in detecting the non-culturable
245 bacterial strains, which are induced under the stressed conditions of PEC. The so-called viable but
246 non culturable (VBNC) state of microorganisms is very common when dealing with environmental
247 samples or disinfection techniques, making plating counts inadequate for accurate bacterial
248 enumeration. Bacteria in this state retain their metabolic activity and pathogenic features, posing
249 danger for public health, while they are not recoverable in standard culture media. On the contrary,
250 molecular biology methods are capable of detecting these strains contributing in a more reliable
251 microbial evaluation of environmental samples (Wéry et al. 2008). Nevertheless, certain attention
252 should be paid on designing PCR experiments, including the amplicon length of the reactions, since
253 it is strongly correlated to the recorded disinfection efficiency. In the present study, SYBR green
254 PCR reactions yielded a 305 bp product for *E. coli* identification and quantification, resulting in
255 reliable and acceptable evaluation of PEC disinfection efficiency (Süb *et al.* 2009). Also, other
256 points under consideration concerning qPCR is that reactions can be affected by nucleic acid
257 contamination, leading to false positive results and the formation of primer - dimers. Some of the
258 possible sources of contamination are cross-contamination between samples, contamination from
259 laboratory equipment and carryover contamination of amplification products and primers from
260 previous PCRs. Therefore, appropriate measures should be taken to increase the reactions
261 specificity and all experiments should be performed with extra care to minimize contamination
262 risks.

263

264 **Effect of mixed bacterial populations**

265 In a set of experiments, an attempt was made to evaluate the disinfection efficiency of PEC
266 at 2V in water samples containing two bacterial populations. In this sense, sterile water samples
267 were inoculated with either 10^7 CFU/mL *E. coli* or 10^7 CFU/mL *E. faecalis* or both. The results
268 from culture technique and qPCR are summarized in Fig. 3.

269 Screening the inactivation profiles of the tested bacterial strains individually, *E. coli* seems
270 to be less resistant than *E. faecalis* at the experimental conditions in question. This outcome, which
271 is more obvious in qPCR results, was quite expected and in agreement with other studies dealing
272 with the inactivation of various bacterial populations (Cho *et al.* 2011; Li *et al.* 2011). Enterococci
273 are Gram positive bacteria, possessing a thick peptidoglycane cell wall and an additional outer
274 membrane containing two lipid bilayers, which provide them high complexity and potential of
275 preserving their viability during PEC treatment. On the other hand, *E. coli* cells require longer
276 treatment periods so as to achieve equal inactivation rates. In this sense, attention should be paid to
277 the differences among main aquatic microbial indicators, which may lead to different PEC
278 inactivation efficiency (Cho *et al.* 2011).

279 When *E. coli* and *E. faecalis* were inoculated together in water samples, the extent of
280 inactivation for either strain was lower than that of their individual treatment. This may be
281 attributed partially to the elevated initial bacterial concentration, whose influence on PEC has been
282 discussed previously (Fig. 2). Other possible explanations would include the presence of
283 competitive microorganisms and the interaction amongst them and/or the competition for AOS
284 between the bacteria and the organic by-products, released to the solution by the inactivated
285 bacteria (Baram *et al.* 2009). In the present study, although PEC seems to be capable of inactivating
286 the used faecal indicators to a certain extent, the residual cells raise concerns about the suitability of
287 the method for disinfection of complex samples, containing various bacterial populations.

288 These findings are more obvious in qPCR results (Fig. 3b), which reveal longer periods
289 required for bacterial decrease. In the present study, considerable genome copies/mL were recorded
290 after 90 min of treatment, which is in contrast to plate counts. These extended periods could reflect
291 the time necessary to mutate DNA to a point that can no longer be assayed. Cells may have been
292 completely inactivated long before that point. On the other hand, concerns are raised about the
293 suitability of the PCR method to establish a correlation between amplified bacterial DNA and viable
294 *E. coli* and *E. faecalis*. However, because the half-life of the DNA released in the environment is

295 considered to be very short owing to the presence of numerous nucleases it might be deduced that
296 the DNA detected is that contained in non-culturable cells (Lleo *et al.* 2005).

297

298 **Disinfection in real wastewater**

299 PEC was also employed to disinfect biologically treated effluents taken just before the
300 chlorination step. Sterilized samples were inoculated with approximately 10^7 CFU/mL or 10^4
301 CFU/mL *E. coli* (Fig. 4). In this set of experiments, a higher value of applied potential (5V) was
302 chosen to highlight its effect on bacterial inactivation. Given that (i) the aqueous matrix was real
303 wastewater, which is generally considered as a “complex” sample containing various organic and
304 inorganic components and (ii) raising the anodic potential enhances photocatalytic rates (Baram *et*
305 *al.* 2009), runs were performed at 5V.

306 As seen in Fig. 4a, total *E. coli* inactivation occurred in almost 15 min when the initial cell
307 density was 10^4 CFU/mL and this increased to 90 min at 10^7 CFU/mL. The degree of disinfection
308 was also determined by means of qPCR, estimating bacterial genome copies remained in the
309 aqueous solution after treatment (Fig. 4b). Comparing findings from both procedures there was a
310 significant contrast, as already seen in previous experiments conducted in the present study. qPCR
311 showed that even after 90 min of PEC treatment at 5V potential, genome copies of the bacterial
312 indicator decreased by only 5 orders of magnitude, while colonies of *E. coli* were not recoverable at
313 the same time. Apart from factors such as the detection limit of the method or VBNC cells, which
314 have been mentioned previously, an important parameter under consideration is the particulate
315 matter present in wastewater. This aids in the resistance of microorganisms to disinfection, as it
316 may interfere by physically shielding bacterial cells and protecting the integrity of the contained
317 DNA. Therefore, the extent of DNA damage is limited, which inevitably results in high numbers of
318 genome copies detected by qPCR, even after long periods of treatment. In the present study,
319 considerable genome copies/mL were recorded, highlighting the importance of wastewater
320 composition and its bacterial content with respect to PEC efficiency. In this sense, plate counts

321 should be matched to qPCR results in order to perform a reliable and complete evaluation of
322 disinfection techniques, such as PEC and other photocatalytic treatments.

323 Furthermore, part of the photogenerated AOS may be wasted to attack the organic carbon of
324 the wastewater (about 8 mg/L which typically consists of highly resistant humic-type compounds
325 and biomass-associated products) and/or scavenged by bicarbonates, sulfates and chlorides (whose
326 collective concentration is about 460 mg/L) rather than inactivate pathogens. This could be
327 overcome increasing AOS concentration through raising the applied potential (Baram *et al.* 2009)
328 and would possibly justify the nearly identical *E. coli* profiles shown in Figs. 2a and 4a at an initial
329 concentration of 10^7 CFU/mL.

330

331 **Conclusions**

332 • Photocatalytic inactivation of *E. coli* in water samples is enhanced applying a positive potential
333 on TiO₂/Ti-films under simulated solar radiation. At the conditions in question, initial bacterial
334 density plays an important role, as it affects adversely the required treatment time for complete
335 microbial inactivation.

336 • *E. faecalis* are more resistant than *E. coli* bacteria during PEC treatment in water samples.
337 Furthermore, the disinfection efficiency is directly affected by the sample composition and the
338 contained mixed bacterial populations. In complex samples, bacteria are more physically
339 protected, resulting in lower inactivation rates.

340 • There was a considerable contrast between colony counts and qPCR results, concerning the
341 required time for total bacterial inactivation. qPCR data revealed longer periods required for
342 complete bacterial inactivation, compared to the time estimated by culture method. Stressed
343 conditions generated during PEC treatment may induce VBNC state of bacteria, which could
344 explain the discrepancy between the applied quantitative methods. Additionally, in order to
345 establish the accuracy and reliability of qPCR certain factors should be under consideration.

346 These include detection limits of the method, amplicon size, primer sequence, cycling conditions
347 and inhibitors present in the processed sample.

- 348 • For large scale applications, the use of sunlight as renewable energy source would undoubtedly
349 be beneficial to induce photocatalytic reactions, as well as provide the necessary electric field,
350 thus promoting the principle of sustainable development.

351

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454

455 **Figure captions**

456

457 **Figure 1**

458 *E. coli* inactivation in water samples during PEC, PC and EO. Conditions: 10^7 CFU/mL initial
459 bacterial concentration; 2V potential.

460 **Figure 2**

461 *E. coli* inactivation in water samples containing various initial bacterial concentrations during PEC
462 (2V potential), assessed by (a) the culture technique and (b) qPCR.

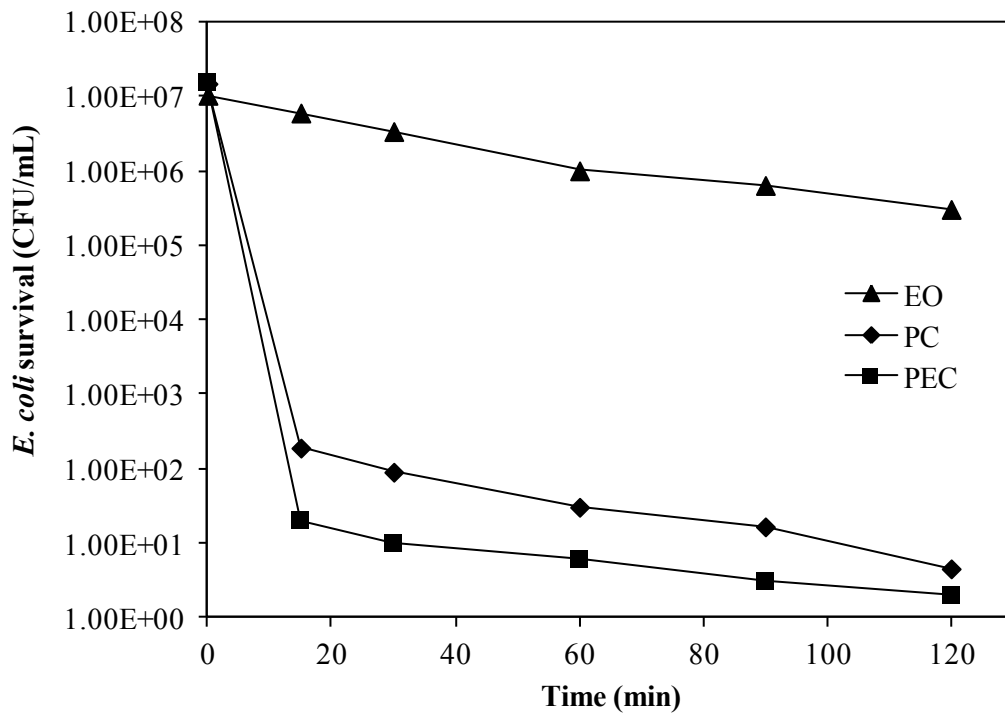
463 **Figure 3**

464 *E. coli* and *E. faecalis* inactivation in water samples during PEC (2V potential), assessed by (a) the
465 culture technique and (b) qPCR.

466 **Figure 4**

467 *E. coli* inactivation in real wastewater during PEC (5V potential), assessed by (a) the culture
468 technique and (b) qPCR.

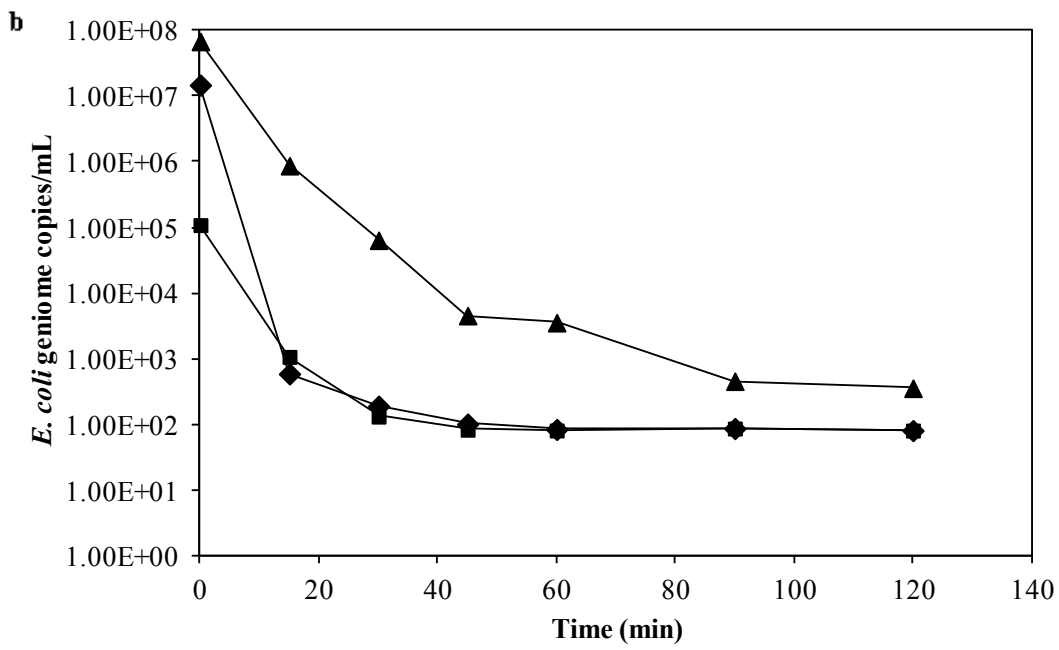
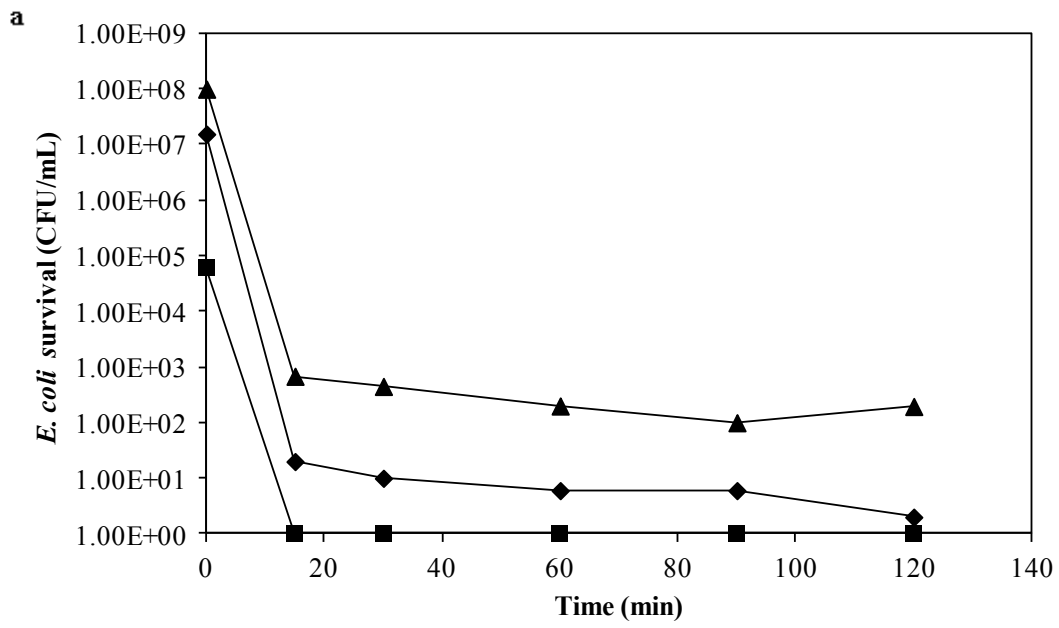
469 **Figure 1**



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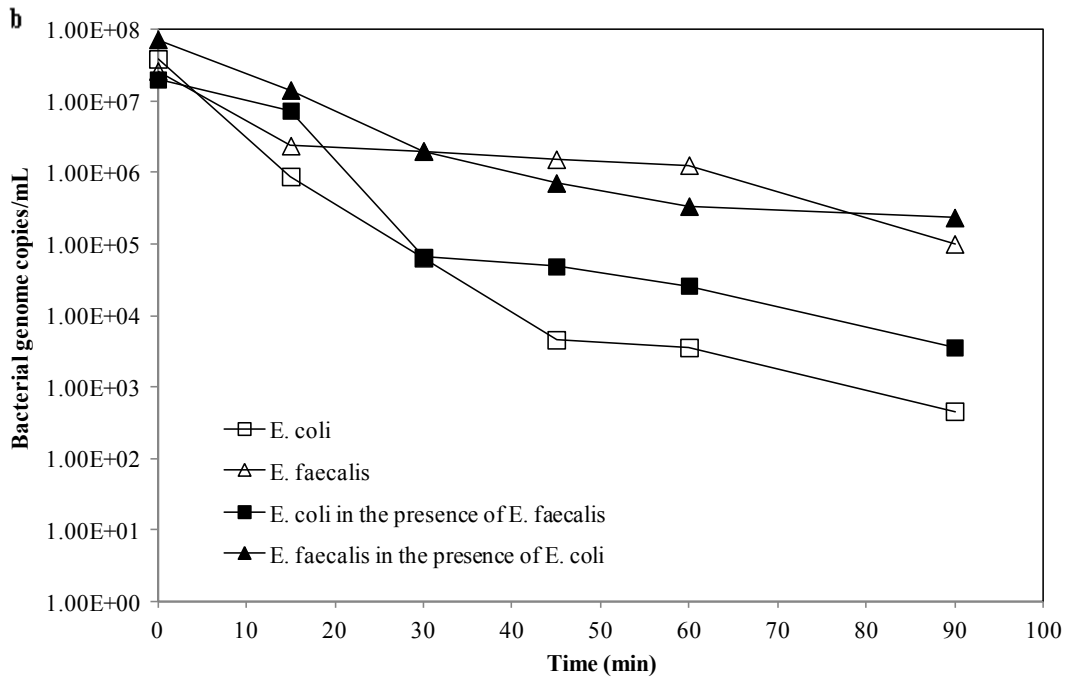
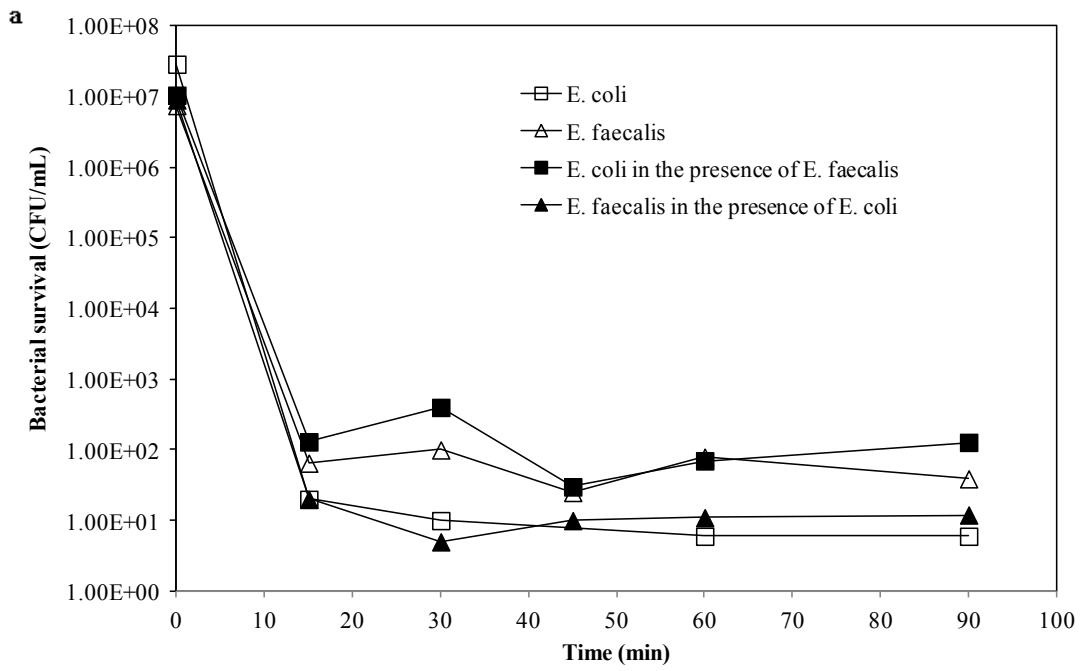
472 **Figure 2**



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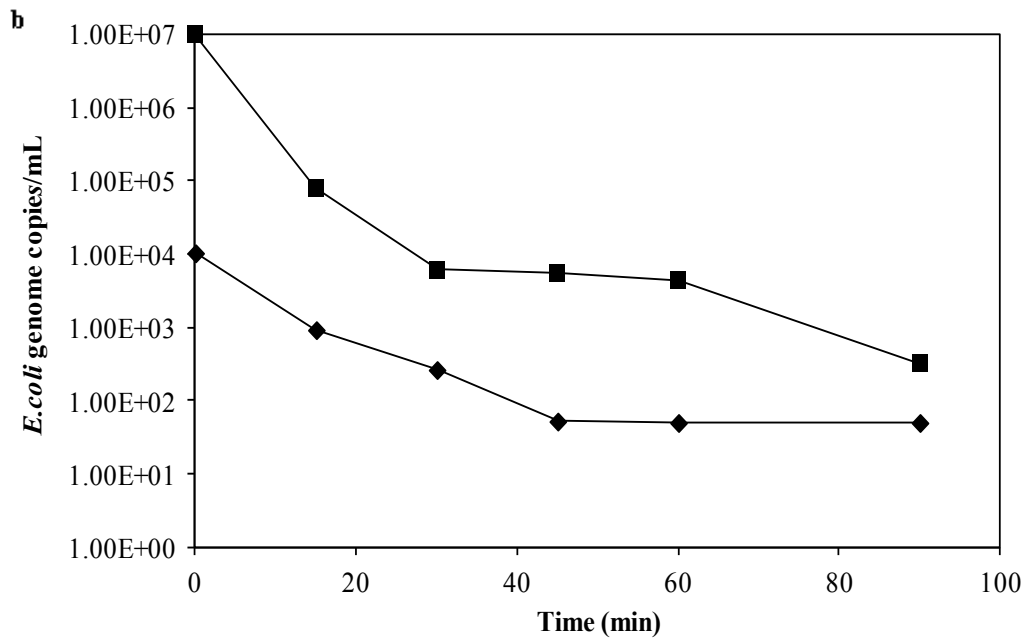
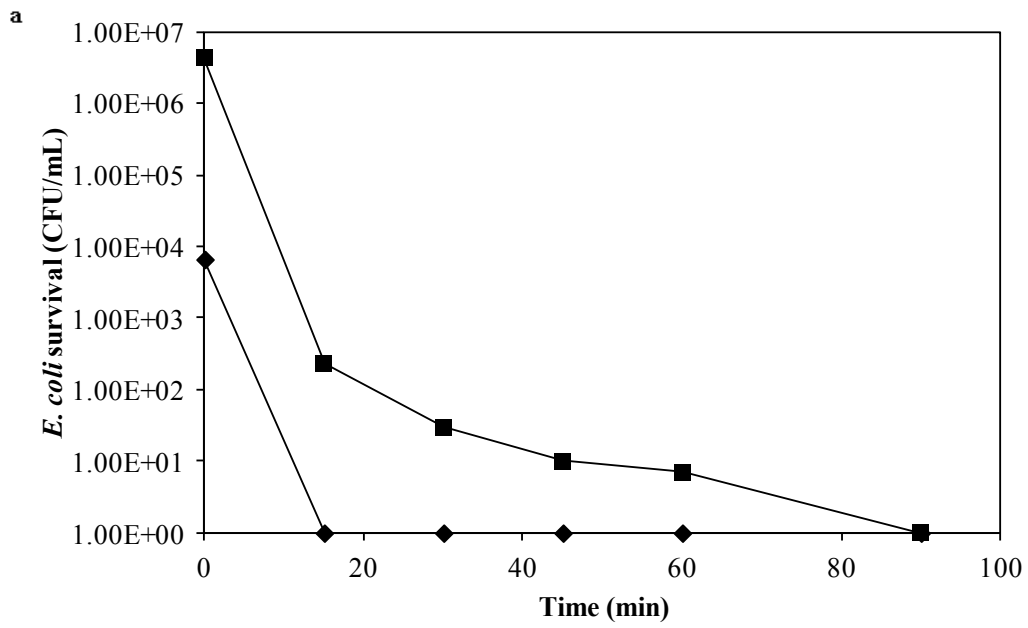
475 **Figure 3**



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478 **Figure 4**



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