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Environmental sustainability of light-driven processes for wastewater treatment applications

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1 **Environmental sustainability of light-driven processes for wastewater treatment**
2 **applications**

3

4 Spyros Foteinis^a, Alistair G. L. Borthwick^b, Zacharias Frontistis^c, Dionissios
5 Mantzavinos^c, Efthalia Chatzisyneon^{b*}

6

7 ^a Senior Engineer, Greek Public Power Corporation (PPC) Renewables S.A.,
8 Kapodistriou 3, Ag. Paraskeui, GR-15343, Attica, Greece

9 ^b School of Engineering, Institute for Infrastructure and Environment, University of
10 Edinburgh, Edinburgh EH9 3JL, United Kingdom

11 ^c Department of Chemical Engineering, University of Patras, Caratheodory 1,
12 University Campus, GR-26504 Patras, Greece

13 * Corresponding author: e.chatzisyneon@ed.ac.uk, tel.: +44(0)1316505711

14

15 **Abstract**

16 A comparative analysis is presented of light-driven advanced oxidation processes in
17 terms of environmental sustainability. Photochemical oxidation has proven a viable
18 option for treating emerging and priority pollutants at laboratory scale. Nevertheless,
19 as a nascent technology, photocatalysis is yet to be widely applied at large-scale water
20 treatment plants. This paper presents a powerful tool that should enable stakeholders
21 to develop sustainable, large-scale, photocatalytic treatment plants by providing
22 knowledge of environmental sustainability and hotspots (where technological flaws

23 have high environmental impact) and understanding as to how process sustainability
24 can be improved through scenario analyses. The following processes were examined:
25 natural and simulated solar photolysis, solar photo-Fenton without hydrogen peroxide
26 addition (solar/Fe), solar photo-Fenton (solar/Fe/H₂O₂), photolysis under UV-A
27 irradiation (UV-A), titania-mediated photocatalysis (UV-A/TiO₂), photolysis under
28 UV-C irradiation (UV-C), and UV-C treatment with hydrogen peroxide addition (UV-
29 C/H₂O₂). Actual life cycle inventory data were collected at bench scale, and the
30 environmental performances estimated by means of life cycle assessment. Effective
31 removal of 1 µg of 17α-ethynylestradiol per liter of wastewater, a commonly occurring
32 micropollutant and endocrine disrupting chemical, was used as the functional unit.
33 Solar photolysis exhibited an environmental footprint about 23 times higher than
34 solar/Fe. Solar/Fe/H₂O₂ minimized the environmental footprint. Being energy
35 intensive, simulated solar irradiation had a much higher (~ 5-fold) environmental
36 footprint than natural solar light. UV photolysis exhibited low environmental impact,
37 with UV-C found to be about 3 times more environmentally friendly than UV-A
38 photolysis. Addition of TiO₂ to UV-A and H₂O₂ to UV-C caused their total
39 environmental impacts to decrease by about 97% and 88%, implying that UV-A/TiO₂
40 was better than UV-C/H₂O₂. In terms of total environmental footprint, the AOPs
41 descend in the following order: solar photolysis > UV-A > UV-C > solar/Fe > UV-
42 A/TiO₂ > UV-C/H₂O₂ > solar/Fe/H₂O₂. The environmental sustainability of all
43 processes was directly proportional to treatment efficiency but inversely proportional
44 to treatment time (due to the large energy input per unit time). Although reagent use
45 (i.e. titania, iron, and hydrogen peroxide) was not associated with high environmental
46 impact, its addition greatly improved process efficiency as well as environmental
47 sustainability. For all examined light-driven processes, the main environmental

48 hotspot was electricity consumption. Introduction of renewable energy sources could
49 reduce the environmental footprint of oxidation processes by up to 87.5%.

50

51 **Keywords:** water purification; estrogens; photocatalysis; LCA; EDCs; EE2

52

53

54

55 **1. Introduction**

56 Trace- or micro-pollutants are synthetic chemicals of emerging environmental and
57 health concern that have recently been detected **in the aquatic environment** (Tiedeken,
58 2017). **Several hundred EDCs have been measured in humans and wildlife, even in**
59 **such remote places as the Arctic** (Birnbaum, 2013). There is growing evidence that
60 these pollutants have adverse effects on human health and living organisms. Trace-
61 pollutants can act, or have the potential to act, as endocrine-disrupting chemicals
62 (EDCs) that cumulatively interfere with the endocrine system of living organisms and
63 cause genetic abnormalities, infertility, feminization, increased cancer rates, trigger
64 Alzheimer disease, etc. (Rochester, 2013). EDCs derive from the chemical processing
65 industry in the form of drugs, surfactants, cosmetics, and other personal care products,
66 which usually end up in the sewage system. Synthetic estrogens are EDCs that are
67 found in increasing concentrations in natural waters (Zhang et al., 2014) and
68 wastewater (Mohagheghian et al., 2014). A representative synthetic estrogen is 17 α -
69 ethynylestradiol (EE2), which is the basic component of the contraceptive pill. EE2 is
70 more stable in an aqueous environment and has greater estrogenic potency (~11–27

71 times) than natural estrone (E1) and estradiol (E2). Continuous exposure to EE2, even
72 to concentrations of $\mu\text{g/L}$, has been found to cause bodyweight loss, accelerate
73 vaginal opening, alter estrous cycles in young animals, and damage fish populations
74 (Frontistis et al., 2015).

75 Due to their xenobiotic and non-biodegradable nature, conventional biological
76 wastewater treatment plants (WWTPs) cannot effectively remove EDCs, which in
77 turn are discharged into receiving waters. To overcome this, it is necessary to add
78 robust tertiary treatment technologies to existing WWTPs. Of the technologies
79 available for the removal of EDCs, light-driven advanced oxidation processes (AOPs)
80 offer considerable promise. The effectiveness of AOPs is mainly due to the formation
81 of reactive oxygen species (ROS), such as hydroxyl radicals (HO^\bullet), which
82 subsequently oxidize the organic content of water samples. AOPs include solar, UV-
83 A and UV-C photolysis and photocatalysis, usually accelerated by adding titania
84 (TiO_2) (i.e. heterogeneous catalysis) (Lee et al., 2017), hydrogen peroxide (H_2O_2)
85 and/or iron (Fe^{2+}) to form the photo-Fenton reagent (i.e. homogeneous catalysis)
86 (Clarizia et al., 2017). To date, several studies have investigated the treatment of EE2-
87 contaminated water by means of UV photocatalysis (Madsen and Sogaard, 2012), UV
88 photolysis (Marinho et al., 2013; Sichel et al., 2011), and solar photocatalysis (Kim et
89 al., 2017; Koutantou et al., 2013). Even so, apart from UV-C photolysis, other light-
90 driven AOPs are still nascent technologies, not yet applied at industrial-scale.

91 AOPs are energy intensive, with high operating cost and elevated
92 environmental footprint (Chatzisyneon et al., 2013). Solar photo-Fenton AOPs have
93 high chemical demand, and generate residual fluxes with negative environmental
94 impacts, such as sludge contaminated by metal ions, exhausted solid catalysts, etc.
95 (Rodríguez et al., 2016). Previous research has focused on the degradation efficiency

96 and techno-economic feasibility of AOPs, without detailed consideration of
97 environmental sustainability (Rodríguez et al., 2016). A brief review of existing
98 studies on AOPs environmental sustainability is given by (Ioannou-Ttofa et al., 2016).

99 In order for AOP technology to reach prototype-scale applications, it must be
100 acceptable from an environmental perspective. To achieve this, the environmental
101 sustainability of each AOP should first be assessed at bench- or pilot-scale, in order to
102 identify merits and drawbacks, establish the main environmental impact hotspots, and
103 assess ways of reducing the total environmental footprint through scenario and
104 sensitivity analyses. By determining the optimal environmental performance of AOPs,
105 the technology could be effectively scaled up to sustainable, large-scale applications
106 in water treatment works.

107 This paper describes a comparative life cycle assessment (LCA) of seven well-
108 established light-driven AOPs, namely: solar, solar/Fe, solar/Fe/H₂O₂, UVA,
109 UVA/TiO₂, UVC, and UVC/H₂O₂. The aim is to identify the strengths and
110 weaknesses of AOPs from an environmental sustainability perspective, thus enabling
111 process scale up. LCA methodology is employed, in accordance with ISO 14040 and
112 ISO 14044 (ISO, 2006a, b), using SimaPro 8. The assessment was made using life
113 cycle inventory (LCI) data collected from bench-scale experiments, rather than
114 extracted from a database. The results should provide researchers, decision- and
115 policy-makers, and the water treatment industry with a better understanding of the
116 environmental sustainability of light-driven AOPs, which in turn should help advance
117 the technology so that it becomes ready for industrial-scale application. To the best of
118 the authors' knowledge this is the first study to date dealing with LCA of several
119 light-driven oxidation processes. Many publications focus on comparing several
120 irradiation sources in terms of ability to decontaminate/disinfect water and

121 wastewater. Assessment of environmental sustainability of such processes, including
122 both solar and UV-irradiated techniques, is presently missing from the literature.

123 Of the various methodologies used to assess the environmental sustainability
124 of a product or process, the most commonly utilized are multi-criteria analysis
125 (MCA), environmental performance indicators (EPIs), and life cycle assessment
126 (LCA) (Hermann et al., 2007). MCA compares and ranks alternative options, and
127 evaluates environmental consequences according to established criteria. However, its
128 weakness lies in the subjectivity of the weighting step, necessary to evaluate different
129 criteria. EPIs estimate the current or past environmental performance of an
130 organisation and compare it against a set of targets; however, the usefulness of EPIs is
131 limited by insufficient data availability (Hermann et al., 2007). LCA offers an
132 effective means of including environmental considerations in the design, production,
133 use, and disposal of a product (Foteinis et al., 2011). LCA is a tool for the systematic
134 evaluation of environmental impacts, which provides insight into the overall
135 performance and relative contributions of different stages within the product lifespan
136 (Hermann et al., 2007).

137

138 **2. Materials and methods**

139 Data used in the comparative LCA analysis were obtained from laboratory
140 experiments, described by Frontistis et al. (2011, 2012, 2015). All experiments were
141 carried out under the same ambient temperature and water conditions. Table 1 lists the
142 optimum operating conditions assayed for each light-driven process. In all cases, the
143 wastewater sample was stirred by a 50 W magnetic stirrer and the ambient
144 temperature kept constant at 25 ± 2 °C. Energy required to keep the temperature

145 constant was external to system boundaries, while the stirrer was assumed to operate
146 at 30 W (i.e. not at full power). At industry scale, wastewater pumping would replace
147 the magnetic stirrer. Simulated solar irradiation was emitted by a Newport, model
148 96000, 150 W solar simulator system. The UV-A and UV-C experiments were
149 conducted in an immersion well, batch type, laboratory-scale photoreactor (Ace
150 Glass, Vineland, NJ, USA). UV-A irradiation was provided by a 9 W lamp (Radium
151 Ralutec, 9W/78, 350–400 nm). UV-C irradiation was provided by an 11 W low-
152 pressure mercury lamp (Phillips, TUV PL-S). The Fe²⁺ ionic solution used in the
153 experiments was in the form of FeSO₄·7H₂O (≥ 99%, Sigma-Aldrich). H₂SO₄ was
154 added in order to regulate the initial water pH. TiO₂ P25 was donated by Evonik
155 Industries, and H₂O₂ (35% w/w) was purchased from Merck.

156

157 **3. Environmental sustainability analysis**

158 To assess the environmental sustainability of light-driven AOPs, LCA methodology
159 was employed, as detailed in ISO 14040 and 14044 (ISO, 2006a, b). Bench-scale
160 experimental results were utilized by the environmental model. The timespan covered
161 2010 to the present date, the geographical boundaries encompassed Greece and
162 similar countries, and average technology was assumed. For the foreground system,
163 primary inventory data were collected for laboratory-scale experiments, while, for the
164 background system, data were used regarding the most recent average technology
165 (e.g. for electricity the average technology mix in Greece was imported from the
166 ecoinvent database).

167

168 **3.1 Functional unit**

169 The functional unit selected to quantify the performance of a light-driven AOP was
170 the effective removal of 1 μg EE2 per liter of treated wastewater. The life cycle
171 inventory (LCI) for each AOP under study was then normalized per functional unit
172 (ISO, 2006a, b) in order to study the environmental performance of the different
173 technologies. Attributional life cycle assessment (ALCA) was used because it
174 estimates the environmental impacts of a product or system according to the delivery
175 of a specified quantity of the functional unit (Chatzisyneon et al., 2016).

176 **3.2 System boundaries and life cycle inventory (LCI)**

177 The system boundaries define which unit processes (the smallest elements for which
178 input and output data are quantified in the LCI) are included within the LCA (ISO,
179 2006a). Energy and raw material requirements, waterborne emissions, and the
180 materials' disposal or recycling are included within system boundaries.

181 For the solar and UV photoreactor AOPs, LCI data could not be identified and
182 so their primary materials, i.e. glass, lamps, and the stirrer, were taken into account. It
183 was assumed that both photoreactors have similar dimensions and materials, and that
184 all experiments were carried out at the same ambient temperature. Two different
185 scenarios were examined for the solar AOPs. The first scenario comprised the
186 photoreactor and lamp (i.e. simulated solar irradiation), whereas the second scenario
187 did not include the lamp (i.e. natural solar irradiation). The latter scenario is closer to
188 actual operating conditions of solar AOPs. Following Ioannou-Ttofa et al. (2017), the
189 photoreactor glass was assigned a useful lifespan of five years (10 h/d operation, all
190 year round). Recycling was also incorporated. Photoreactor lamps are not included in
191 SimaPro's proprietary life cycle inventory (LCI) databases, and so the LCI data were
192 obtained from relevant literature (Garrett and Collins, 2009; OSRAM, 2016). The

193 data were re-scaled according to the power requirements of each process and input to
194 SimaPro in order to simulate the environmental impact of each lamp under study.
195 Data on the stirrer used to mix effluent were not available in SimaPro's proprietary
196 LCI databases, and so were substituted by relevant data concerning the LCI of a low-
197 power motor (AAB, 2002), re-scaled to fit the rated output of the stirrer under study,
198 and used as input to SimaPro.

199 Information on the Fe^{2+} ion as iron sulphate was supplied from the SimaPro
200 LCI databases. Residual Fe^{2+} in the treated wastewater was also taken into account as
201 waterborne emission. Data on H_2O_2 and H_2SO_4 reagents were obtained from
202 proprietary LCI databases. Energy used to drive each process was supplied as
203 electricity from the Greek energy grid, which is fossil fuel-dependent and comprises
204 54% lignite, 11% crude oil, 17% natural gas, and 18% renewable energy (Ioannou-
205 Ttifa et al., 2016). To carry out the comparative analysis, from an environmental
206 perspective, of light-driven AOPs, the final use and disposal route of treated effluent
207 was taken to be external to system boundaries. In other words, **cradle-to-gate** (treated
208 effluent) was used.

209

210 Table 1.

211

212 **3.3 Life cycle impact assessment (LCIA)**

213 Life cycle impact assessment (LCIA) relates the data inventory to specific
214 environmental impacts and damages (ISO, 2006a, b). ReCiPe was chosen for the
215 LCIA as a robust method that comprises both midpoint and endpoint impact/damage
216 approaches which examine different stages in the cause-effect chain to calculate

217 impact (Chatzisyneon et al., 2016). The endpoint, or damage-oriented, approach
218 translates environmental impacts into issues of concern, such as human health, natural
219 environment, and natural resources. Endpoint results are associated with higher levels
220 of statistical uncertainty, compared to midpoint, due to data gaps and assumptions
221 stacking up along the cause-effect chain, but are easier for decision- and policy-
222 makers to comprehend (Chatzisyneon et al., 2016). Given that this is a comparative
223 LCA, results are compared using the following three endpoint damage categories:
224 “Human Health”, “Resources”, and “Ecosystems”. These can be also aggregated into
225 a single score, which makes interpretation simpler.

226 A hierarchist perspective (H), based on the most common policy principles,
227 was invoked within ReCiPe along with European normalization and average
228 weighting. Decisions whether or not to include information in the H model are based
229 on mean scientific consensus, and it assumes that, with proper management,
230 environmental impacts can be avoided (Chatzisyneon et al., 2016), thus fitting better
231 the goal and scope of the comparative analysis.

232 Moreover, in order to ensure accuracy and transparency of the LCA, the
233 primary LCI data along with data used for the background system were verified
234 against information from the open literature (Chatzisyneon et al., 2013; Gimenez et al
235 2015). Light-driven AOPs comprise a nascent technology for wastewater treatment,
236 and so comparative environmental studies based on similar operating conditions and
237 similar initial organic loads are needed; however, information on these important
238 parameters is scarce.

239

240 **3.4 Energy consumption**

241 The energy consumption of artificial lighting constitutes a major fraction of the
242 operating costs in UV treatment. Bolton et al. (2001) introduced the electric energy
243 per order, E_{EO} , defined as the energy required for 90% degradation of a pollutant per
244 m^3 of contaminated water. E_{EO} (kWh/ m^3 /order), for a batch-operated reactor, is
245 calculated from the following equation:

$$246 \quad E_{EO} = \frac{P \times t \times 1000}{V \times 60 \times \log(C_i/C_f)} \quad (1)$$

247 where P is the electrical power of the irradiation source (kW), t is the irradiation time
248 (min), V is the volume of the treated effluent (L), and C_i and C_f are the initial and the
249 final pollutant concentrations (mg/L), respectively.

250

251 **4 Results and discussion**

252 To render the analysis both comprehensive and straightforward to follow, the results
253 for the solar and UV irradiation light sources are considered separately. Then, a
254 comparative analysis of all processes follows **in order** to identify the most promising
255 **result** in terms of environmental sustainability. Finally, a sensitivity analysis is carried
256 out using scenarios to investigate the effect of the main environmental hotspots and to
257 propose “greener” alternatives by which to improve sustainability.

258 **4.1 Environmental sustainability of solar-driven AOPs and effects of Fe^{2+} and** 259 **H_2O_2**

260 Results provided by ReCiPe for natural and simulated solar-driven
261 photolysis/photocatalysis at endpoint level (Figure 1) show that simulated (artificial
262 light) and natural solar photolysis yielded by far the highest environmental footprints
263 of ~ 11 mPt and ~ 2 mPt per functional unit, respectively. The environmental footprint

264 due to photolysis was ~ 23 times larger than that of simulated/natural solar/Fe, using
265 low reagent concentration (5 mg/L Fe²⁺), with scores of 0.477 mPt (artificial light)
266 and 0.089 mPt (natural light). For photolysis, as well as all other AOPs, the main
267 environmental hotspot was electricity use derived from Greece's fossil fuel-dependent
268 electricity mix. At the time of writing, electricity systems worldwide use fossil fuels
269 for bulk power generation (Berill et al., 2016) and so the foregoing results are
270 presently valid for Greece, Europe and beyond. Indirect impacts of the use of
271 electricity from fossil fuels can be traced mainly to the "Human Health" damage
272 category, followed by "Resources", and less so the "Ecosystem" (Chatzisyneon et al.,
273 2016). "Human Health" damage is affected by fossil-fuel mining and combustion,
274 which release toxic materials including metals, sulphur, and polycyclic aromatic
275 hydrocarbons (PAHs) to the environment (Chatzisyneon et al., 2016). Fossil-fuel
276 extraction and burning contribute to climate change. Natural gas extraction also
277 releases SO₂. Impacts from coal arise from tailpipe emissions after combustion and
278 emissions during blasting at coal mines (Berill et al., 2016). "Resources" damage is
279 primarily caused by depletion of fossil fuels for electricity generation and of mineral
280 resources used to construct equipment required for resource extraction, processing
281 and consumption, and to a lesser degree by equipment related to AOPs (i.e. the stirrer
282 and photoreactor). Turning to "Ecosystem" damage, phosphate leachate from coal
283 mining spoil landfill sites and the emission of nitrogen oxides from combustion of
284 fossil-fuel directly impact on acidification and eutrophication. Waterborne metal
285 emissions from coal power plants, natural gas extraction (particularly of bromine) and
286 from disposed coal mine spoil (nickel and magnesium) affect **ecotoxicity** (Berill et al.,
287 2016; Ioannou-Ttofa et al., 2016).

288 Use of simulated irradiation raised the environmental impact because the total
289 environmental footprint of simulated solar photolysis and photocatalysis is about a
290 factor of 5 higher than natural solar light. This is attributed to electricity consumption
291 by the lamp (~ 81.3% of total environmental footprint), and to a much lower degree to
292 the lamp material (~ 0.05% of total environmental footprint). In terms of material, the
293 stirrer (i.e. motor) contributed 12.4% and 2.3% to the total environmental footprints
294 for natural and simulated solar photolysis. Finally, the photoreactor material (glass)
295 made a very low contribution to the total environmental footprint, 0.257% and
296 0.0494% for natural and simulated solar photolysis, respectively, mainly because of
297 the long lifespan of glass whose recycling was included in the system boundaries.
298 The relatively high environmental footprint of solar photolysis is due to its low
299 treatment efficiency as it consumes energy during the stirring process while EE2 is
300 removed from wastewater.

301

302 Figure 1.

303

304 To study the environmental impacts of the more environmentally friendly
305 natural solar-driven AOPs, a separate comparison was undertaken, neglecting
306 photolysis and simulated solar irradiation. Figure 2 shows that the amount of oxidation
307 reagents used strongly affected the environmental sustainability of solar-driven AOPs,
308 with high reagent concentration improving the overall environmental sustainability of
309 solar AOPs. At low concentration of iron ions (5 mg/L Fe^{2+}) the total environmental
310 footprint of natural solar/Fe was estimated to be 0.089 mPt, whereas when the
311 concentration was increased to 15 mg/L the total environmental footprint reduced by

312 about half to 0.047 mPt per functional unit (Figure 2). When H₂O₂ was also added as
313 a reagent, the environmental sustainability of the process was further enhanced. More
314 specifically, when keeping the iron ion concentration constant at 5 mg/L and adding
315 10 mg/L H₂O₂ the total environmental footprint of the process was ~0.01 mPt per
316 functional unit, and by increasing the H₂O₂ concentration to 17.2 mg/L the
317 environmental footprint of the process achieved a minimum of $\sim 0.356 \times 10^{-3}$ mPt per
318 functional unit.

319 This large reduction is attributed to: (a) increased degradation efficiency at
320 higher H₂O₂ concentration (Table 1); (b) lower treatment time (15 min for 10 mg/L
321 H₂O₂, and 1 min for 17.2 mg/L H₂O₂) and hence reduced energy consumption; and (c)
322 use of low amounts of H₂O₂, a non-toxic chemical without elevated environmental
323 impact. As mentioned before, the environmental impacts of solar/Fe can be traced
324 back to Greece's fossil fuel-dependent electricity mix used to drive the stirrer. The
325 contributions of electricity consumption to the total environmental footprint of natural
326 solar/Fe (5 mg/L and 15 mg/L), natural solar/Fe/H₂O₂ (10 mg/L) and natural
327 solar/Fe/H₂O₂ (17.2 mg/L) were 87.4%, 87.3% and 86.5%. The photoreactor and the
328 stirrer-drive motor made material contributions of 0.256 ± 0.02 % and 12.35 ± 0.05
329 %. As a non-hazardous reagent when in small concentrations, Fe²⁺ had a negligible
330 effect in all cases (its biggest score was 0.058% in natural solar/Fe/H₂O₂ (17.2
331 mg/L)). Similarly, the addition of miniscule amounts of H₂SO₄ in concentrations of
332 about 50 μ L/L led to it also making a negligible contribution. For natural
333 solar/Fe/H₂O₂, addition of hydrogen peroxide at concentrations of 10 mg/L and 17.2
334 mg/L contributed ~0.037% and 0.943% to total environmental footprint. The latter,
335 higher percentage contribution is related to the overall low environmental footprint of
336 the process (0.356×10^{-3} mPt) and the higher quantity of hydrogen peroxide used (and

337 the knock-on increased energy and materials required for its synthesis). It should be
338 noted that no H₂O₂ emissions (e.g. airborne, waterborne) or harmful by-products **were**
339 assumed to be generated during treatment.

340

341 Figure 2.

342

343 **4.2 Environmental sustainability of UV-A and UV-C photocatalysis**

344 Figure 3 presents the environmental footprints of UV-A and UV-C
345 photolysis/photocatalysis in terms of “Human Health”, “Resources” and
346 “Ecosystems” endpoint damage categories. UV-A photolysis yields a higher
347 environmental footprint (0.309 mPt), whereas that of UV-C is about a factor of three
348 smaller (0.117 mPt). This is expected because UV-C treatment has a much higher
349 treatment efficiency due to the higher energy (Frontistis et al., 2015), compared to
350 UV-A treatment. In both cases the lamp materials hardly contributed to the total
351 environmental footprint, whereas the UV-C lamp required about 20% higher power
352 but also had significantly higher treatment efficiency (see Table 1). As a result, UV-C
353 removed 1 µg/L of EE2 at a much faster rate than UV-A treatment, requiring less
354 energy and contributing less environmental footprint per functional unit.

355 When reagents were added, the environmental footprint of both UV-A and
356 UV-C treatment **was** substantially reduced. Figure 3 shows that addition of titania (10
357 mg/L TiO₂) drastically reduced the total environmental footprint of UV-A treatment,
358 from ~309 µPt for UV-A photolysis to ~9.2 µPt for UV-A/TiO₂ heterogenous
359 photocatalysis. As far as UV-C treatment is concerned, the addition of H₂O₂ (10
360 mg/L) **also had** a profound effect, with the environmental footprint of UV-C

361 photolysis reducing from ~117 μPt for UV-C to ~13.8 μPt for UV-C/H₂O₂. These
362 large reductions (~97% for UV-A/TiO₂ and ~88% for UV-C/H₂O₂) are due to a
363 combination of improved treatment efficiency and reduced treatment time (Table 1).

364 As with solar-driven AOPs, the environmental sustainability of UV-driven
365 AOPs is enhanced by addition of small amounts of the non-hazardous reagents, TiO₂
366 and H₂O₂, leading to significant improvement in degradation efficiency and reduction
367 in treatment time, especially for UV-A treatment.

368 Electricity consumption makes the largest contribution to most damage
369 categories, reflected by its contribution to the total environmental footprint of UV-
370 driven AOPs of 88.3 ± 0.1 %. This score is dominated by electricity consumption by
371 the stirrer motor and, to a lesser degree, to the lamp(s). The stirrer motor as a material
372 was the next most important environmental hotspot with scores ranging from 9.15%
373 to 9.63% of the total environmental footprint. The lamp as a material contributed from
374 1.79% for UV-A to 2.35% for UV-C. The photoreactor as a material (glass)
375 contributed from 0.19% for UV-C to 0.29% for UV-A. The reagents TiO₂ and H₂O₂
376 contributed very low percentages, 0.236% for UV-A/TiO₂ and 0.0273% for UV-
377 C/H₂O₂, of the total environmental footprint. Even though TiO₂ had a higher impact
378 than H₂O₂, UV-C/H₂O₂ exhibited a slightly higher total environmental footprint than
379 UV-A/TiO₂, mainly due to the reduced treatment time of the latter (Table 1).

380

381 Figure 3.

382

383 **4.3 Environmental sustainability of solar versus UV-A and UV-C photocatalysis**

384 Given that photolysis invariably exhibited the highest overall environmental footprint,
385 the most promising photocatalytic processes were determined in terms of
386 environmental sustainability. Figure 4 presents a comparative analysis, using ReCiPe
387 impact assessment method, of natural solar/Fe, natural solar/Fe/H₂O₂, UV-A/TiO₂ and
388 UV-C/H₂O₂ photocatalysis. Natural solar/Fe/H₂O₂, at high reagent concentrations
389 (Fe²⁺ = 5 mg/L and H₂O₂ = 17.2 mg/L) yielded the lowest score (0.356 μPt per
390 functional unit) amongst all processes. For simulated solar irradiation, the total
391 environmental footprint of solar/Fe/H₂O₂ rose to 1.869 μPt, but nevertheless remains
392 substantially lower than all the other light-driven AOPs considered. Again, the
393 presence of iron and hydrogen peroxide oxidants, the reduced treatment time and
394 enhanced EE2 removal efficiency caused the energy demand per functional unit to be
395 minimized, lowering the environmental footprint. The next most environmentally
396 friendly AOPs were UV-A/TiO₂ (~9.2 μPt or ~96% higher than natural
397 solar/Fe/H₂O₂) and UV-C/H₂O₂ (~13.8 μPt). Both exhibited relatively high treatment
398 efficiency, with UV-A/TiO₂ requiring less treatment time to achieve EE2 removal
399 (Table 1), which meant less energy input and a lower environmental footprint than
400 UV-C/H₂O₂. Also, the lamp required higher energy to drive the UV-C/H₂O₂ process
401 (11W) than UV-A/TiO₂ (9 W). The lamps, photoreactor, and stirrer made low
402 contributions as materials to the total environmental footprints of the UV-C/H₂O₂ and
403 UV-A/TiO₂ processes. The contribution by the reagents, TiO₂ and H₂O₂, was
404 miniscule compared to electricity consumption. Finally, natural solar/Fe exhibited a
405 high overall environmental footprint, especially at low reagent concentration (5 mg/L
406 Fe²⁺) where the value was 0.089 mPt. For a high iron concentration (i.e. 15 mg/L), the
407 total environmental footprint was halved, to 0.047 mPt per functional unit (Figures 2
408 and 4).

409 In short, all the light-driven AOPs considered in this work were limited by the
410 same environmental hotspot, namely electricity consumption from Greece's fossil
411 fuel-dependent energy mix, which dominated the contributions to ReCiPe's damage
412 categories "Human Health" and "Resources". Similar findings were obtained by
413 (Chatzisyneon et al., 2013) who compared the environmental sustainability of UV-
414 A/TiO₂ with electrochemical and wet air oxidation processes for treatment of agro-
415 industrial wastewater.

416

417 Figure 4.

418

419 The present comparison is based on bench-scale experimental data. It is expected that
420 further benefits can be achieved for all AOPs examined, in terms of lowering the
421 environmental footprint per functional unit when the processes are scaled up. For
422 example, in prototype applications, the stirring processes, which required large energy
423 inputs at bench scale, will be replaced by pumping which is more energy efficient.
424 Given that it also consumes electricity, pumping is likely to be a prime environmental
425 hotspot (as also suggested by Foteinis et al. (2018) in a study of pilot-scale Fenton
426 processes for pharmaceutical wastewater treatment).

427 Energy consumption to degrade 90% of EE2 was also estimated in order to undertake
428 a more comprehensive comparative analysis of artificial light-driven oxidation
429 processes. The corresponding treatment time was estimated either using experimental
430 values from photocatalytic tests carried out by Frontistis et al. (2015), Frontistis et al.
431 (2012) and Frontistis et al. (2011) or by extrapolating the experimental values to
432 achieve 90% removal of EE2. The results are shown in Table 2, where it is observed

433 that UVA/TiO₂ process has the lowest energy demands followed by UVC/H₂O₂,
434 solar/Fe(5mg/L)/H₂O₂, UVC, UVA, solar/Fe(15mg/L), solar/Fe(5mg/L) and
435 simulated solar process. In principle, these results are consistent with those obtained
436 from LCA (Figure 4) confirming the high dependence of AOPs on electricity
437 consumption.

438

439 Table 2.

440

441 4.5 Sensitivity analysis

442 The main environmental barrier to light-driven AOPs under study is electricity
443 consumption from the Greek energy mix dominated by fossil fuels. Power systems
444 based largely on renewable energy sources (RES) perform much better regarding
445 climate change and other impact categories than systems based on fossil fuels (Berrill
446 et al., 2016). A sensitivity analysis was carried out involving three energy mix
447 scenarios solely based on RES, i.e. solar, wind, and hydropower, all naturally
448 abundant in Greece, Europe and beyond. Energy storage, curtailment, and grid
449 extension were neglected because the aim of scenario analysis is purely to illustrate
450 possible pathways and futures, rather than make forecasts or predictions (Kouloumpis
451 et al., 2015). Moreover, the extra impacts caused by energy storage and grid extension
452 are likely to be of such relatively small magnitude that the environmental benefits of
453 switching to renewables would not be undermined (Berrill et al., 2016). The use of
454 RES to meet the electricity needs of light-driven AOPs is expected to lead to
455 substantial improvement in their environmental sustainability. For example, use of an
456 electricity mix solely based on photovoltaic (PV) systems (i.e. 3 kWp single-Si panels

457 mounted on slanted roofs) decreases the total environmental footprint of solar AOPs
458 by about 85% and UV-driven AOPs by 87%. On the other hand, use of an electricity
459 mix solely based on wind energy (onshore wind turbines, capacity in the range from 1
460 to 3 MW) further improves the environmental sustainability of light-driven AOPs
461 because energy from wind turbines usually has a lower environmental impact than
462 solar PVs (Chatzisymeon et al., 2016). In this case, the total environmental footprint
463 of solar AOPs and UV-driven AOPs is decreased by about 81% compared to the
464 initial scenario. Finally, use of an electricity mix solely based on hydropower leads to
465 the largest decrease in total environmental footprint of light-driven AOPs by 86%
466 (solar) and 87% (UV) because hydropower is the most environmentally friendly RES
467 option (Ioannou-Ttofa et al., 2016).

468 In all cases, the highest reduction in environmental footprint occurred for the
469 most energy intensive AOPs (i.e. simulated solar, UV-A, and UV-C, ordered from
470 higher to lower reduction), whereas the smallest reduction occurred for the most
471 energy efficient AOPs (i.e. solar/Fe/H₂O₂, solar/Fe, UV-A/TiO₂, and UV-C/H₂O₂,
472 ordered from lower to higher reduction). The order of light-driven processes in terms
473 of environmental sustainability remained the same for all scenarios; from higher to
474 lower score: natural or simulated solar > UV-A > UV-C > natural or simulated
475 solar/Fe > UV-A/TiO₂ > UV-C/H₂O₂ > natural or simulated solar/Fe/H₂O₂. Even so, it
476 should be noted that UV-A/TiO₂ and UV-C/H₂O₂ exhibited similar environmental
477 footprints when using RES.

478

479 **5. Conclusions**

480 This paper has investigated the environmental performance of light-driven AOPs at
481 removing an endocrine disruptor, EE2, from wastewater using actual life cycle
482 inventory (LCI) data. It was found that the environmental sustainability of light-
483 driven AOPs was directly proportional to treatment efficiency (which was expected
484 given that the chosen functional unit was the removal of 1 μg EE2 per liter of
485 wastewater), and was also inversely proportional to treatment time. Moreover,
486 electricity consumption from the fossil fuel-dependent Greek energy mix was the
487 main environmental hotspot for all examined AOPs. The Fe^{2+} , H_2O_2 , and H_2SO_4
488 reagents used in light-driven AOPs were associated with low environmental impacts
489 because the chemicals did not detrimentally affect health or the eco-system, no
490 harmful by-products were generated, and only low dosages were used. Use of RES to
491 meet the electricity needs of light-driven AOPs substantially improved their
492 environmental sustainability, by up to 87% for solar- and 88% for UV-driven AOPs.

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555 northeastern China. *Archives of environmental contamination and toxicology* 66, 361-369.

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559 **List of Tables**

560 Table 1: Experimental data, taken from Frontistis et al. (2011, 2012, 2015), used to
561 build the LCI of light-driven AOPs.

562 Table 2: Electrical energy (E_{EO}) consumed by several light-driven oxidation
563 processes.

564 Table 1.

Light-driven process	Irradiation power, W	[TiO ₂], mg/L	Power for water stirring, W	[Fe ²⁺], mg/L	H ₂ O ₂ , mg/L	Treatment time, min	EE2 removal, µg/L	Reference
Solar	150	-	30	-	-	60	2	(Frontistis et al., 2015)
Solar/Fe	150	-	30	5	-	60	46	(Frontistis et al., 2015)
	150	-	30	15	-	60	86	(Frontistis et al., 2015)
Solar/Fe/H ₂ O ₂	150	-	30	5	10	15	98	(Frontistis et al., 2015)
	150	-	30	5	17.2	1	196	(Frontistis et al., 2011)
UVA	9	-	30	-	-	60	17	(Frontistis et al., 2015)
UVA/TiO ₂	9	750	30	-	-	10	95	(Frontistis et al., 2012)
UVC	11	-	30	-	-	60	47	(Frontistis et al., 2015)
UVC/H ₂ O ₂	11	-	30	-	10	15	100	(Frontistis et al., 2015)

565

566

567 Table 2.

Light-driven process	Irradiation power, kW	Volume, L	Treatment time to remove 90% of EE2, min	<i>E</i> _{EO} , kWh/m ³ /order	Reference
Solar	0.150	0.3	2251	18758	(Frontistis et al., 2015)
Solar/Fe(5mg/L)	0.150	0.3	115	958	(Frontistis et al., 2015)
Solar/Fe(15mg/L)	0.150	0.3	70	583	(Frontistis et al., 2015)
Solar/Fe(5mg/L)/H ₂ O ₂	0.150	0.3	2	17	(Frontistis et al., 2015)
UVA	0.009	0.3	312	156	(Frontistis et al., 2015)
UVA/TiO ₂	0.009	0.3	7	4	(Frontistis et al., 2012)
UVC	0.011	0.3	113	69	(Frontistis et al., 2015)
UVC/H ₂ O ₂	0.011	0.3	10	6	(Frontistis et al., 2015)

568

569

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571 Figure 1. Environmental footprint of natural and simulated solar (a) photolysis and (b)
572 photocatalysis per functional unit, i.e. removal of 1 μg EE2 per liter of wastewater.

573 Figure 2. Environmental footprint of natural solar photocatalysis for removal of 1 μg
574 EE2 per liter of wastewater. Inset: environmental footprint of natural solar/Fe/H₂O₂
575 (17.2 mg/L).

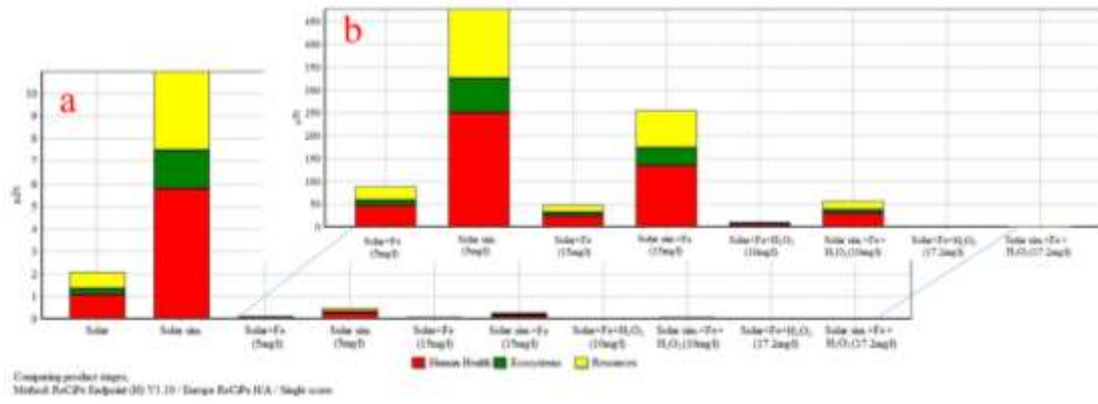
576 Figure 3. Environmental footprint of UV-A and UV-C AOPs per functional unit
577 (removal of 1 μg EE2 per liter of wastewater).

578 Figure 4. Environmental footprint of natural solar, UV-A and UV-C photocatalysis
579 for removal of 1 μg EE2 per liter of wastewater.

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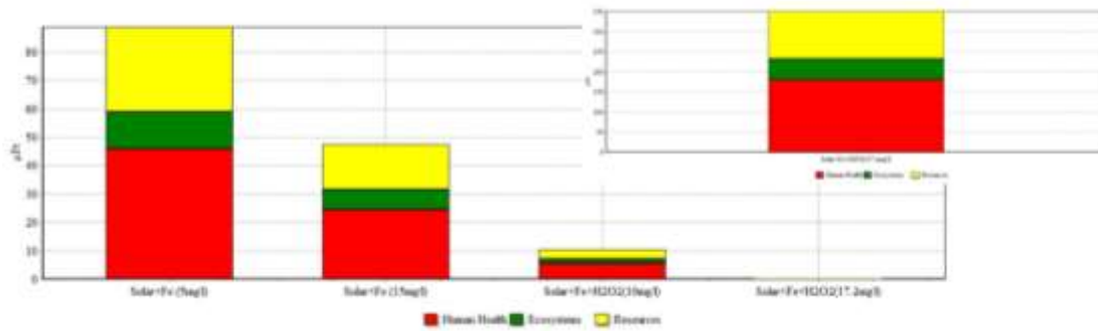
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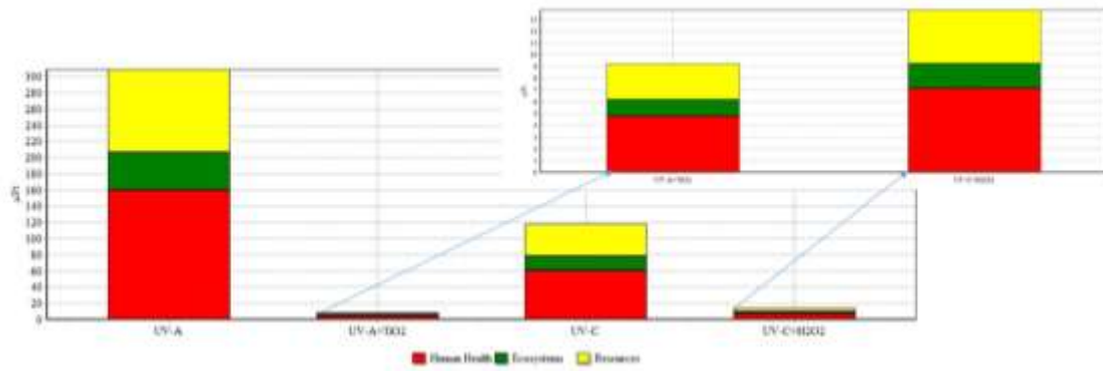
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589 Figure 2.

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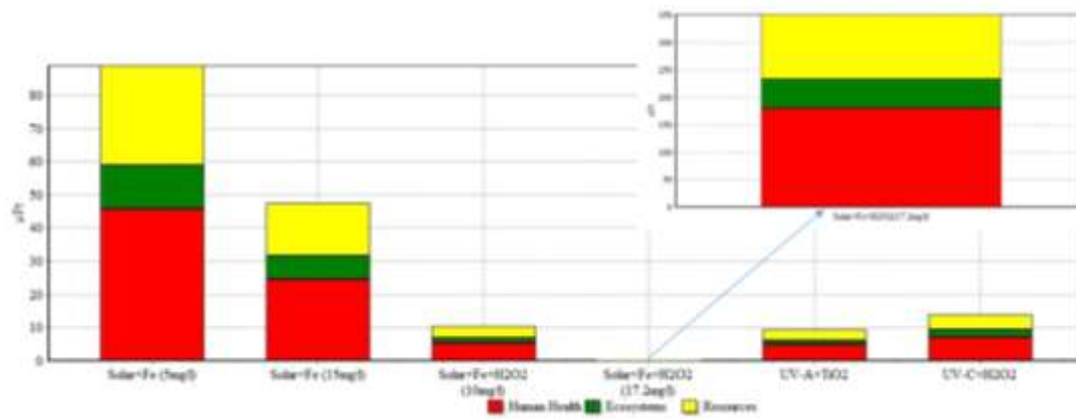
Method ReCIPs Endpoint (B) V1.10 - Europe ReCIPs HIA - Single score

594 Figure 3.

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Method ReCIPs Endpoint (B) V1.10 - Europe ReCIPs HIA - Single score

599 Figure 4.

600