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Evaluation of Fenton and modified Fenton oxidation coupled with membrane distillation for produced water treatment: Benefits, challenges, and effluent toxicity

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1	Evaluation of Fenton and modified Fenton oxidation
2	coupled with membrane distillation for produced water
3	treatment: Benefits, challenges, and effluent toxicity
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19 ABSTRACT

20 Membrane distillation is a promising technology to desalinate hypersaline produced waters. 21 However, the organic content can foul and wet the membrane, while some fractions may pass 22 into the distillate and impair its quality. In this study, the applicability of the traditional 23 Fenton process was investigated and preliminarily optimized as a pre-treatment of a synthetic 24 hypersaline produced water for the following step of membrane distillation. The Fenton process was also compared to a modified Fenton system, whereby safe iron ligands, i.e., 25 26 ethylenediamine-N,N'-disuccinate and citrate, were used to overcome practical limitations of the traditional reaction. The oxidation pre-treatments achieved up to 55% removal of the 27 dissolved organic carbon and almost complete degradation of the low molecular weight toxic 28 organic contaminants. The pre-treatment steps did not improve the productivity of the 29 30 membrane distillation process, but they allowed for obtaining a final effluent with 31 significantly higher quality in terms of organic content and reduced Vibrio fischeri inhibition, 32 with half maximal effective concentration (EC_{50}) values up to 25 times those measured for the raw produced water. The addition of iron ligands during the oxidation step simplified the 33 34 process, but resulted in an effluent of slightly lower quality in terms of toxicity compared to the use of traditional Fenton. 35

36

37 **Keywords**: membrane distillation; produced water; advanced oxidation; iron ligands;

38 ecotoxicity

39 HIGHLIGHTS

- Thermal and modified Fenton degraded target contaminants in produced water.
- The oxidative pre-treatment decreased organics in the membrane distillation effluent.
- 42 Coupled oxidation and membrane distillation decreased the toxicity of the final effluent.
- Traditional Fenton pre-treatment provided the best effluent in terms of toxicity.
- Modified Fenton nearly degraded all target contaminants in hypersaline solutions.

GRAPHICAL ABSTRACT



49 **1. Introduction**

50 Despite the current transition to more sustainable sources of energy, oil and gas extraction still plays a significant role in the energy sector and wastewater treatment is quickly emerging 51 52 as one of the most significant challenges of this industry. Indeed, the so-called produced water (PW) is the largest waste stream generated in oil and gas extraction activities 53 (Ahmadun et al., 2009). Considering both onshore and offshore sites, the global PW 54 production has increased from 150 million to around 300 million barrels per day from 1990 55 to 2015 (Ahmadun et al., 2009; Liu et al., 2021). The average water cut, namely, the amount 56 of water volume produced per oil volume, is roughly 3:1 (Ahmadun et al., 2009; Jimenez et 57 al., 2018; Liu et al., 2021; McCormack et al., 2001). As oilfields age during the current 58 59 energy transition, the water cut will also increase, together with wastewater treatment difficulties (Igunnu and Chen, 2014). PW is a highly complex matrix, rich in organic and 60 inorganic compounds, with widely diverse composition as a function of geological formation, 61 age of the oilfield, and type of hydrocarbon product being extracted (Ahmadun et al., 2009; 62 Estrada and Bhamidimarri, 2016). However, the major compounds are typically dispersed 63 oils, dissolved organics (e.g., phenols, benzene, toluene, xylenes), dissolved minerals (e.g., 64 sodium chloride, calcium and magnesium salts), and natural organic matter (NOM) 65 66 (Ahmadun et al., 2009; Al-Ghouti et al., 2019; Coha et al., 2021; Estrada and Bhamidimarri, 2016; Jimenez et al., 2018; Kabyl et al., 2020; Liu et al., 2021; Neff et al., 1992). 67

The oil and gas industry faces increasing pressure to limit its environmental footprint (Mohammad-Pajooh et al., 2018). Expensive treatment trains for a multi-contaminated water, as well as water scarcity and increasing international attention to environmental issues, are the drivers pushing this industry to use water more sustainably, bringing along the concepts of water reuse and safe water discharge. As a result, innovative, environmentally focused, and reliable methods of meeting water treatment demands, capable of operating in this 74 specific application are being developed (Estrada and Bhamidimarri, 2016; Liu et al., 2021; Mohammad-Pajooh et al., 2018). These technologies should be versatile to meet the 75 76 requirements of low cost and compactness, the latter characteristic being especially important 77 in offshore activities (Kabyl et al., 2020; Liu et al., 2021). One of the target parameters that needs abatement to allow for water reuse or safe discharge is salinity, which presents average 78 values of 50-100,000 ppm in PW (Coha et al., 2021; Estrada and Bhamidimarri, 2016). Such 79 salt concentration can plug the reinjection well or may be toxic if PW is discharged in the 80 81 environment without desalination (Aquiliina, 2012; Ariono et al., 2016; Canedo-Arguelles et 82 al., 2019; Chen et al., 2016; Kleinitz et al., 2001; Liu et al., 2021). Membrane distillation (MD) is a promising emerging technology, capable to extract high-quality effluents from 83 84 hypersaline solutions using low-grade energy and with relatively low capital cost, due to the 85 absence of high pressure and high temperature components (Chen et al., 2017; Han et al., 86 2017; Howell, 2004; Lin et al., 2014; Shaffer et al., 2013). In recent studies, MD was successfully tested on hypersaline PW (Han et al., 2017; Li et al., 2014; Ricceri et al., 2019). 87 88 However, since the MD membranes are highly hydrophobic (PTFE and PVDF membranes 89 are generally used), this process may present important practical limitations in the presence of 90 a large and broad content of organic compounds, such as for typical PW (Estrada and 91 Bhamidimarri, 2016; Gonzalez et al., 2017). Organics may either induce wetting phenomena 92 or freely pass through the hydrophobic membrane and end up in the final effluent (Chen et 93 al., 2017; Franken et al., 1987; Kargbo et al., 2010; Wang et al., 2018). Wetting phenomena 94 in MD occur when the transmembrane pressure (ΔP) exceeds the liquid entry pressure (LEP), according to eq. (1), thus allowing the contaminated feed water to pass undisturbed through 95 96 the porous membrane (Franken et al., 1987; Horseman et al., 2021a; Ricceri et al., 2019).

97

$$\Delta P \ge LEP \tag{1}$$

98 LEP is defined according to eq. (2):

99
$$LEP = -\frac{2B\gamma cos\theta}{r}$$
(2)

100 where γ is the feed water surface tension, θ is the intrinsic contact angle between the feed 101 water and the solid membrane material, r is the equivalent pore radius, and B is a geometric 102 factor accounting for the noncylindrical nature of the membrane pore geometry (B = 1 for)perfectly cylindrical pores). A low surface tension γ reduces also $\cos\theta$, thus facilitating 103 membrane pore wetting. The degradation of toxic organic compounds in the feed solution 104 105 both increases its surface tension and thwarts their interaction with the hydrophobic 106 membrane, potentially allowing for a more efficient MD process and a more effective management of the PW. 107

108 The list of available technologies and processes counts a plethora of options to remove or partially degrade organic compounds (Adewumi et al., 1992; Ahmadun et al., 2009; Chang et 109 110 al., 2019a; Chang et al., 2019b; Estrada and Bhamidimarri, 2016; Liu et al., 2021; Shang et 111 al., 2019; Tang et al., 2021). Activated carbon adsorption and sand filtration are low-cost treatment processes; however, they produce harmful waste since they do not degrade the 112 113 toxic organic compounds. Biological treatment is not effective toward biorecalcitrant organic compounds, such as benzene, toluene, xylenes (BTX), and requires large plants with long 114 115 retention times, hence not available for offshore platforms (Ayed et al., 2017). Electrochemical, photocatalytic, and ozone-based oxidations are growing rapidly, but they 116 117 are still currently associated with high capital costs and with difficulties in practical 118 implementation (Ahmadun et al., 2009; Bessa et al., 2001; Coha et al., 2021; Dalmacija et al., 1996; Ma and Wang, 2006; Ricceri et al., 2019; Shokrollahzadeh et al., 2012). Among 119 advanced oxidation processes (AOPs), the Fenton reaction involves the use of iron sulfate 120 121 and hydrogen peroxide to generate highly reactive hydroxyl radicals able to oxidize almost

all the organic compounds (Coha et al., 2021; Haber et al., 1934; Miklos et al., 2018). The
main reaction is as follows:

$$Fe(II) + H_2O_2 \rightarrow Fe(III) + {}^{\bullet}OH + OH^-$$
(3)

The Fenton process is a promising candidate to treat PW both onshore and offshore since it is 125 versatile, characterized by high kinetics also at room temperature, and capable to remove 126 organics from a multi-contaminated matrix (da Silva et al., 2015; Jiménez et al., 2019; 127 Jiménez et al., 2017; Jin and Davarpanah, 2020; Ruiz-Aguirre et al., 2017; Zhai et al., 2018). 128 This method has some limitations, mainly the need for acidic pH to avoid iron hydroxide 129 precipitation, and the production of sludge once neutral pH is restored (Diya'uddeen et al., 130 131 2012). A modified Fenton process encompassing the addition of an iron ligand helps overcoming these precise challenges (Chahbane et al., 2007; Farinelli et al., 2020; Farinelli et 132 al., 2019; Messele et al., 2019). However, the literature lacks reports about the application of 133 134 Fenton processes carried out in presence of iron ligands to treat PW.

135 The first objective of this work is to evaluate a coupled system including Fenton (or 136 modified Fenton) pre-oxidation and MD to desalinate PW and to allow for an easy management of the final effluent. This sequence is applied to treat a synthetic PW that 137 138 mimics the effluent from primary treatment, which typically includes de-oiling and flotation or sedimentation. A thermal Fenton reaction is first applied as a potential PW oxidation step 139 and as a pre-treatment for MD desalination. The study gives insight into the relationship 140 between the content of organics and the performance of the membrane distillation step by 141 comparing a raw feed stream with the feed subject to Fenton oxidation. Furthermore, the 142 143 performance of traditional Fenton is compared with that of modified Fenton systems. To this purpose, non-toxic and biodegradable organic ligands, namely, citrate and EDDS, are added 144 145 in PW at unadjusted pH to assess the ability of iron-ligand complexes to act as effective 146 oxidation catalysts (Chen et al., 2019; Tandy et al., 2006; Van Devivere et al., 2001). The

safety of the final desalinated effluent from the coupled system is then fully evaluatedthrough acute toxicity measurements.

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150

151 2. Materials and Methods

152 2.1. Chemicals, membrane, and produced water preparation

All the organic contaminants, the iron ligands, i.e., sodium citrate and EDDS, ferrous 153 sulfate (FeSO₄), hydrogen peroxide (30% w/w), HCl, and NaOH, were purchased from 154 Sigma-Aldrich (Milan, Italy). Sodium chloride, sodium sulfate, and sodium bicarbonate were 155 acquired from Carlo Erba (Milan, Italy). All the solutions needed for the acute toxicity 156 analysis, namely, the reconstitution, the diluent, and the osmotic solutions were purchased 157 from Modern Water (London, UK). The freeze-dried Vibrio fischeri culture was purchased 158 159 from Ecotox LDS (Cornaredo (MI), Italy). Type I ultrapure water was used for the 160 experiments. A commercially available polytetrafluoroethylene (PTFE) membrane (Aquastill, Sittard, Netherlands) was deployed in MD filtration tests. 161

The composition of the synthetic PW was based on published values of real wastewaters 162 and is listed in **Table 1**, together with the resulting total organic carbon (TOC) and total 163 dissolved solids (TDS) values (Coha et al., 2021; Estrada and Bhamidimarri, 2016; Olsson et 164 al., 2013). Humic acids and a liquid petroleum jelly consisting of paraffins were used as 165 representative compounds for natural dissolved organic matter and oil & grease, respectively 166 167 (Lester et al., 2015). Xylenes, benzene, toluene, and methyl tert-butyl ether (MTBE) were 168 selected as representative volatile organic compounds (VOCs) (Coha et al., 2021). Cyclohexane was added as representative of the $< C_{10}$ hydrocarbon fraction (Estrada and 169 Bhamidimarri, 2016; Lester et al., 2015). Phenol was added as representative substance for 170

the common phenols content in PW. The TDS included sodium, calcium, and magnesium

172 chlorides. All the components were added into water and the matrix was sonicated at room

temperature for 1 h to enhance solubilization and mixing.

174

175 **Table 1** Composition of the synthetic produced water, compared with the reference real

176 streams. The matrix includes representative pollutants to mimic typical TOC and TDS values.

		Synthetic produced water		Real produced water
Parameter	Component	Concentration (ppm)	Equivalent TOC (ppm)	Concentration (ppm)
	Paraffins	200.0	Not dissolved	
	Humic acids	200.0	60.0	
	Cyclohexane	2.0	1.8	
	Phenol	2.5	1.9	
TOC	Xylenes	1.0	1.0	
	Benzene	12.0	11.3	
	Toluene	4.0	3.4	
	MTBE	260.0	178.0	
	TOT	681.5	257.4	Maximum ~500
	Sodium chloride	100,000		
тре	Calcium chloride	2,500		Typically, 35,000 – 240,000
103	Magnesium chloride	4,000		
	TOT	106,500		Average ~100,000
pН		5.5		

177

178 *2.2.Oxidation conditions*

All the oxidation reactions were performed at room temperature $(25 \pm 2 \,^{\circ}C)$ under gentle stirring for a total duration of 1 h, using different ratios of hydrogen peroxide and catalyst. The catalyst was iron(II) in the case of the traditional Fenton process, and a complex ligandiron(II) in the case of modified Fenton. To promote organics oxidation, three additions of hydrogen peroxide (0, 20, 40 min) were carried out, each one consisting of a 1/3 aliquot of the desired total amount. The dosages of iron sulfate and hydrogen peroxide are listed in **Table 2** for the traditional Fenton system; in these cases, the pH of the synthetic PW was adjusted to ~3 (HCl). At the end of the reaction, the pH was increased to ~10 by addition of NaOH and this step caused the precipitation of $Fe(OH)_3$. After the sedimentation of the precipitate at 4 °C overnight, the supernatant was collected and used for analysis and as a feed matrix for the following MD filtration tests.

190

191 **Table 2** Traditional Fenton dosages in tests operated at different oxidation conditions.

Entry	FeSO₄ (mM)	H ₂ O ₂ (mM)
MQ H ₂ O	-	-
Produced water	-	-
Ox 1	0.5	5
Ox 2	5.0	50
Ox 3	5.0	25
Ox 4	1.0	25
Ox 5	1.0	50
Ox 6	0.1	5
Ox 7	5.0	100

192

To perform the modified Fenton oxidations, the iron-ligand complexes, namely, Fe-EDDS and Fe-citrate were spiked in the synthetic PW from a stock solution containing 0.1 M of iron(II) and 0.1 M of individual ligand. The pH was not adjusted and was equivalent to ~4 upon addition of Fe-EDDS and to ~5 upon addition of Fe-citrate. No precipitate formation was observed in these systems. The resulting samples were used for analysis and as feed for the following MD filtration tests without further processing.

199 2.3.Membrane distillation tests

The MD tests were performed in direct contact configuration using a lab-scale batch system (Ricceri et al., 2019). The feed and distillate streams were circulated counter-currently on their respective sides of the membrane. A constant crossflow rate of 1.66 L/min (0.278 m/s crossflow velocity) was maintained during the tests. The housing cell comprised a 250mm long, 50-mm wide, and 2-mm deep rectangular channel for a total active membrane area

of 125 cm². The flux across the membrane was computed by recording the change in weight 205 of the distillate tank in time through a computer-interfaced balance. Initial volumes of ~1.9 L 206 and 1 L were used for the feed and distillate streams, respectively, unless otherwise stated. 207 208 Water was used in the distillate side, with specific conductivity always below 20 µS/cm. The specific conductivity in the distillate tank was measured continuously during each test by a 209 conductivity meter (COND 7+, XS Instruments, Italy). The temperature of the feed and 210 211 distillate tanks were maintained constant throughout the experiments, at respective values of 50 ± 2 and 25 ± 1 °C, by means of a thermostatic water bath and a chiller. 212

213 2.4. Analytical methods

214 The TOC of the matrices was measured using a Shimadzu TOC-L analyzer (catalytic oxidation on Pt at 680 °C). The calibration was performed using standards of potassium 215 phthalate and of NaHCO₃/Na₂CO₃. The headspace, solid phase microextraction technique 216 (HS-SPME) was chosen as extraction method before carrying out the GC-MS analysis. 217 218 Following each reaction experiment, the vials were left in a thermostatic bath at 50 °C for 10 min to promote the transfer of all the relevant compounds into the gas-phase headspace. 219 Then, a SPME fiber (df 75 µm, fiber assembly carboxen/polydimethylsiloxane, Supelco) was 220 221 inserted through the septum of the cap and was left in the headspace for 10 min, before withdrawing it for the subsequent GC-MS analysis. Samples were analyzed on an Agilent 222 6890 GC system coupled with an Agilent 5973 mass selective detector (MSD). For the 223 224 chromatographic separation, a Zebron-5MS capillary column (30 m·× 0.25 mm·× 0.25 μ m) was used. The injection port temperature was 270 °C, and the oven temperature program was 225 226 set as follows: 35°C for 5 min, followed by an increase to 260 °C at a rate of 15 °C/min (total run time 25.33 min). Helium was used as carrier gas at a constant flow of 1 mL/min, and the 227 injector was held in splitless mode. The interface temperature was 270 °C and the ionization 228

energy was 70 eV. The molecular structures of the by-products were identified by means ofmass spectrum library.

The determination of the residual iron in solution was carried out by a spectrophotometric 231 procedure adapted from previous literature (Goncalves et al., 2020; Harvey et al., 1955). The 232 total iron was determined by reducing the Fe(III) to Fe(II) with ascorbic acid $(4 \times 10^{-4} \text{ M})$ and 233 complexing the Fe(II) with o-phenanthroline $(4 \times 10^{-3} \text{ M})$ under acidic conditions (buffer pH = 234 3: H₃PO₄ 1 mM, NaH₂PO₄ 3 mM). The Fe(II) was determined without performing the 235 236 reduction step, and Fe(III) was obtained as the difference between total iron and Fe(II). The calibration was obtained using a commercial standard solution of Fe(III) (1000 mg^{Fe}/L, 237 Sigma-Aldrich). The spectrophotometric analyses were performed using a Varian CARY 100 238 239 Scan double-beam UV-vis spectrophotometer, using quartz cuvettes with 10 mm path length and working at a wavelength of 510 nm. Surface tension measurements were performed with 240 241 a digital tensiometer (K10, Krüss): the measurements were repeated three times on each 242 sample.

243 2.5. Toxicity analysis

All the toxicity experiments were performed with a Microtox Model 500 analyzer (Milan, 244 Italy). The analysis was performed with the bioluminescence inhibition assay using the 245 246 marine bacterium Vibrio fischeri. Samples were tested in a medium containing 2% sodium chloride, and the luminescence was recorded after 5, 15, and 30 min of incubation at 15 °C. 247 The luminescence inhibition percentage was determined by comparison with a non-toxic 248 control. The toxicity curves and the values of EC_{50} were obtained from the software 249 (MicrotoxOmni). The pH of all the samples was adjusted in the range 6-8 before the analysis. 250 251 The method used for the toxicity analysis is the method APAT-IRSA 8030 (APAT, 2003). All the samples obtained at the end of the oxidation experiments were quenched with catalase 252 in order to avoid the detrimental effect of the residual hydrogen peroxide on the toxicity 253

254 measurements; see Figure S1 of the Supplementary Material (SM). Moreover, the samples 255 after the oxidation experiments were quantified for residual iron content and EDDS was 256 added in EDDS:Fe molar ratio of 1:1, to prevent the toxic effect of the residual iron in 257 solution (see Figure S2 in SM).

3. Results and Discussion

260 3.1.Efficacy of thermal Fenton oxidation on organics removal

Table 3 and Figure 1a summarize the results of the Fenton oxidations in terms of TOC 261 removal and surface tension (ST) values, as a function of the relative addition of iron(II) and 262 H₂O₂. The highest TOC removal rates were obtained with Fe/H₂O₂ ratios between 0.02 and 263 0.2. At low iron dosage (≤ 0.5 mM, Ox1, Ox6), insufficient catalyst was available in solution, 264 while at high reagent concentrations (Ox7), the reaction was possibly self-inhibited. Previous 265 266 studies highlighted that a Fe/H₂O₂ molar ratio around 0.02 should avoid self-inhibition reactions while providing high efficiency of oxidation (De Laat and Gallard, 1999; Voelker 267 268 and Sulzberger, 1996). Oxidation 3 (Ox3) reached the highest percentage of TOC removal, 269 coinciding with the largest dosage of FeSO₄ (5 mM) and a Fe/H₂O₂ ratio of 0.2. However, Ox4 also achieved a high percentage of TOC removal, but with a substantially lower amount 270 of iron(II), namely, 1 mM, corresponding to a Fe/H₂O₂ ratio of 0.04. These conditions also 271 allowed reaching the highest value of ST (69 \pm 3.1 dyn/cm), close to the ST measured for 272 pure water (72 \pm 1.8 dyn/cm) and substantially higher than that of the PW (50.8 \pm 2.7 273 dyn/cm). Ox4 was thus identified as the most promising oxidation and further tests were 274 conducted using the matrix oxidized under this condition. 275

Table 3 Resulting TOC removal rates, and surface tension values of the oxidized matrix in
tests operated at different oxidation conditions.

Entry	TOC removal (%)	Surface tension (dyn/cm)
Type I H ₂ O	-	72.0 ± 1.8
Produced water	-	50.8 ± 2.7
Ox 1	47.6 ± 0.5	67.8 ± 1.6
Ox 2	55.3 ± 1.3	63.5 ± 1.8
Ox 3	58.5 ± 2.2	60.1 ± 2.3
Ox 4	52.2 ± 2.4	69.5 ± 3.1
Ox 5	50.9 ± 1.7	54.2 ± 2.1
Ox 6	34.7 ± 2.1	68.6 ± 2.0
Ox 7	33.3 ± 2.3	59.3 ± 1.7



Figure 1. (a) Graphical representation of the relationship between TOC removal and reagent dosage. The green color represents higher values of TOC removal. (b) Percentage of degradation of the parent organic contaminants in the synthetic produced water after Fenton reaction (Ox4, 60 min). The percentage of degradation was obtained by computing the target peaks area detected by GC-MS. (c) Chemical structures of the residual by-products preconcentrated onto the fiber during the SPME extraction and detected by GC-MS at the end of the Fenton reaction (Ox4, 60 min).

The solution obtained after Fenton oxidation Ox4 was further characterized through GC-286 MS analysis. Figure S3 in the SM presents the chromatograms and the profile of target 287 substance degradation observed after 20, 40, and 60 min of oxidation. Figure 1b summarizes 288 289 the degradation efficiency after 60 min toward various organic contaminants. A near total degradation of phenol, xylenes, and cyclohexane was observed. The oxidation process 290 degraded benzene and toluene with a yield around 75%. However, only a portion of MTBE 291 292 was degraded, and this result may also explain the residual TOC after Ox4 (see Table 3). Indeed, MTBE contributes to a large part of the total TOC of the synthetic PW (see Table 1). 293 294 The major challenge for the high efficacy of the Fenton reaction in a hypersaline PW is 295 arguably the ability of humic acids and chloride to scavenge the hydroxyl radical (Goldstone et al., 2002; Kiwi et al., 2000), although humic acids may also favor the Fe(III)-Fe(II) 296 297 recycling (Vione et al., 2004). Nevertheless, the observed TOC removal rates and the yield of 298 degradation of target parent substances suggest the high potential of the Fenton reaction to reach suitable levels of decontamination, also in the presence of a significant amount of 299 300 scavengers.

The GC-MS also allowed detection of the main volatile by-products of the Fenton process (Figure 1c). Based on the molecular structure, the compounds labeled as 1, 2 and 3 in Figure 1c reasonably derived from MTBE, while 4 and 5 likely derived from toluene and o-xylene, respectively. Note that the identified by-products were more hydrophilic than the starting contaminants, thus explaining the observed increase in ST. The relatively high percentage of TOC removal together with the formation of more hydrophilic by-products are promising conditions to obtain an improved feed solution of an MD step.

310 Organic compounds in the feed matrix may affect the MD step by fouling and by wetting the membrane, hence lowering the desalination efficiency, or by freely passing through the 311 hydrophobic membrane material (PTFE) (Pasternak and Kolwzan, 2013; Vesterkvist et al., 312 313 2012). Wetting is theoretically described by eq. 1 and experimentally observed with an 314 increase of conductivity in the distillate stream of the MD step (Donaldson et al., 1969; Rezaei et al., 2017). The water flux values presented in Figure 2a suggest that the Fenton 315 316 pre-treatment did not have an effect on the productivity or on the achievable recovery of the MD step under laboratory filtration conditions: the recovery was roughly 75%, upon which 317 the water flux went to zero due to scaling (namely, the deposition of crystals on the 318 319 membrane or within its pores) and pore blockage. Note that similar productivity was also observed when synthetic PW matrices containing solely salts or solely organics were used as 320 321 the feed solutions. In the latter case the flux did not go to zero, due to the absence of salt 322 precipitation, but it steadily decreased during the test possibly due to fouling phenomena. The fouling phenomena in MD are mostly governed by hydrophobic interactions between organic 323 324 compounds and the MD membrane material (PTFE). These interactions cause the deposition and attachment of macromolecules on the membrane surface, thus impairing the MD process 325 326 by influencing the behavior of water at the membrane/solution interface and by partly covering the surface pores (Horseman et al., 2021b). Increased conductivity of the distillate 327 328 solution was measured beginning roughly at 50% recovery for the various feed streams, 329 except that containing only organic compounds; see Figure S4 of the SM. This result suggests 330 that in our study salt passage and possibly wetting were mostly imputable to high salt 331 concentrations, with organics only associated with fouling mechanisms. Salts can crystallize 332 within the pores of the membrane, enlarging them, hence lowering the LEP. Since wetting is observed when the transmembrane pressure overcomes the LEP, the lower is the LEP, the 333

lower is the transmembrane pressure needed to let unpurified liquid water freely pass across

the membrane.



Figure 2 (a) Results of MD filtration tests with different feed solutions: (red circles) synthetic produced water, (blue squares) only the organic content of the synthetic produced water, (orange triangles) only the TDS content of the synthetic produced water, and (green diamonds) the resulting feed after the thermal Fenton oxidation. (b) TOC concentration in the distillate tank (initial volume 1 L) as a function of cumulative permeated volume; the lines connecting the data points are only intended as guides for the eye.

The only organic compounds in the synthetic PW which may induce wetting were humic 343 acids (HA) and the water-miscible compounds (WMC), namely, BTX, MTBE, phenol, and 344 345 cyclohexane. HA are amphiphilic and may partly act as surfactants, which tend to create liquid water channels through the pores of the membrane (Horseman et al., 2021b); however, 346 they do not present a clear separation between the hydrophobic and hydrophilic portion of the 347 molecule, and can virtually maintain a repulsive behavior for liquid water after the interaction 348 349 with the hydrophobic membrane. This can result in fouling, but not necessarily in wetting (Horseman et al., 2021a; Klavins and Purmalis, 2010; Wang et al., 2018). This observation 350 351 has been rationalized with the generally high aromatic fraction of typical HA molecules, with 352 low-density hydrophilic functional groups. On the other hand, WMC may induce wetting by lowering the ST of the solution. A solution containing hydrophobic WMC may be 353 354 characterized by a lower ST than pure water (see Table 3). As LEP is correlated to the ST of the solution (see Eq. 2), it decreases as ST decreases, with higher chances of wetting. 355 However, since no wetting from organic compounds was detected in this study, it is 356 357 reasonable to assume that the ST threshold needed to observe wetting under the condition of this study was lower than the ST value of the synthetic PW (50.8 \pm 2.7 dyn/cm). The fact that 358 organic fouling did not cause wetting may be rationalized with the short duration of the lab 359 experiments, which were run for approximately 8 h before the observed drop in water flux 360 due to scaling. The slow kinetics of fouling phenomena in such a system may require longer 361 362 filtration times to show wetting effects and should become important at real scale during 363 operation. Note that paraffins form a different phase, hence they cannot lower the ST of the solution or create water bridges within the membrane pores, but only freely pass through the 364 365 PTFE membrane by virtue of their hydrophobic nature.

While the Fenton pre-treatment did not provide specific advantages in terms of productivity, which was governed by salt concentration and partly affected by fouling, or of

prevention of wetting, which was not observed even for the use of untreated PW as feed 368 stream, oxidation had beneficial effect in terms of MD effluent quality and toxicity. The 369 370 results in **Figure 2b** suggest a clear reduction of the TOC in the distillate when desalinating 371 the feed matrix subject to Fenton reaction; see Figure S5 of the SM for the chromatograms of the final effluent treated only in MD and by the coupled Fenton-MD system. As expected, the 372 MD process did not separate water from volatile WMC or non-aqueous oils. The first data 373 374 points in Figure 2b showed a high TOC passage, justifiable considering an instantaneous passage of a fraction of the organic content, specifically, paraffins and WMC. Subsequently, 375 376 the TOC in the distillate tank decreased steadily by dilution with the nearly pure water vapor 377 permeating the membrane. The lower amount of TOC measured upon oxidation of the feed stream with Fenton is imputable to both the mineralization of a fraction of toxic compounds 378 379 (~52% of mineralization, Table 3) and the transformation of organic substances to more 380 hydrophilic compounds, which are less prone to pass through the hydrophobic membrane. Fenton oxidation may also provide beneficial effects in terms of MD performance at real 381 382 scale by thwarting fouling phenomena that would occur at longer time scales, but this effect could not be observed in this study (Abdel-Karim et al., 2021; Siyal et al., 2017). 383

384 *3.3.Effect of the coupled system on the toxicity of the final effluent*

385 A general index of the safety of an effluent is its toxicity. Toxicity is also a legislated parameter, allowing or denying the discharge of an effluent in the sewage system. According 386 to the Italian regulations (D.Lgs. 152/2006), the acute toxicity limit to discharge an effluent 387 388 in the sewage system is 80% of the inhibition of the target microorganism (in this case, 389 Vibrio fischeri). The synthetic PW of this study presented an acute toxicity around 100% (Figure S6a of the SM), a value also expected for most of the real PW due to the wide variety 390 391 and large concentrations of contaminants typically present. Both organics and concentrated 392 salts may present large toxic effects, hence they both require a specific treatment. Figure 3 393 presents the residual acute toxicity of the effluent treated with only MD and with the coupled 394 Fenton-MD system, expressed in term of dilution factor of the original samples. The EC_{50} value was used as a comparative parameter of the quality of different effluents, and it is 395 396 defined as the half maximal effective concentration, namely, the concentration required to 397 obtain 50% of microorganism inhibition. Therefore, the higher the EC_{50} value, the lower the toxicity. EC₅₀ increased significantly (from a dilution factor of 0.094 ± 0.009 to 0.25 ± 0.03) 398 399 when the effluent was oxidized and then desalinated, compared to a stream that was not pretreated, i.e., the coupled system gave lower toxicity. However, when tested as is, the residual 400 401 toxicity of the effluent was still high and over the limit of 80% after 30 min of contact time with the bacteria. 402



Figure 3. Residual toxicity of the effluent expressed in term of dilution factor of the original sample (a) after MD treatment only, and (b) upon treatment by the coupled Fenton-MD system (solid data points) with and (empty data points) without Fe sequestration. The toxicity was measured after 5, 15, and 30 minutes of contact (in red, blue, and orange, respectively) with the *Vibrio fischeri* culture. The dash line indicates the point where 50% of acute toxicity is reached.

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411 To understand the relative contribution to toxicity of the different contaminants, toxicity experiments were carried out with individual components (Figures S6 and S7 of the SM). 412 Low toxicity was associated with HA and paraffins, while higher toxic effects were related to 413 414 parent WMC, which however were degraded effectively by Fenton. CaCl₂ and MgCl₂ also showed negligible toxicity, while NaCl showed some toxicity only at concentrations >50 g/L, 415 that is, far above the concentration measured after MD desalination. On the other hand, 416 Fe^{II}SO₄ showed significant toxicity: residual iron may thus be responsible for the toxicity 417 observed for the effluent from Fenton and MD treatments. However, by addition of EDDS in 418 419 molar ratio 1:1 to residual iron (0.26 mM), the acute toxicity of iron was shut down (see Figure S2 and S7 in SM for the toxicity of Fe-EDDS and iron, respectively) and that of the 420 effluent markedly decreased (from ~80% to ~40%), with a substantial increase in EC_{50} (from 421 a dilution factor of 0.25 \pm 0.03 to ~2.5). Remaining toxicity after this post-treatment step may 422 be reasonably attributed to the residual WMC and to the by-products of Fenton oxidation. 423 Note that in this study EDDS was added to mask the toxic behavior of residual iron, which 424 confirms that residual metal concentration causes toxicity in the effluent. Different strategies 425 426 to remove residual iron may be implemented in real plants, for example, its precipitation at 427 basic pH. Cleaner and more novel steps may involve enhanced ion exchange resins and the use of adsorbents, such as magnetic nanoparticles (Khatri et al., 2017). 428

429 3.4. Comparison between traditional Fenton and Fenton process with iron ligands

Iron ligands keep iron in solution without the need for pH adjustment to 3 and limit the production of sludge, thus potentially allowing for a more streamlined operation compared to the traditional Fenton process. Citrate and EDDS were chosen as non-toxic (Figure S2 of the SM) ligands and the Fe-citrate and Fe-EDDS systems were applied as catalysts (Zhang et al., 2016). The optimized condition in terms of Fe-ligand to H_2O_2 molar ratio corresponded to the dosages relative to Ox7 in **Table 2**; each ligand was dosed equimolarly with iron. The results summarized in Figure 4a suggest higher degradation efficiency of the modified Fentonsystems toward WMC with respect to the traditional thermal Fenton.



Figure 4 (a) Comparison of the organics removal rate by traditional Fenton and modified Fenton reactions with citrate and EDDS as iron ligands. Residual toxicity of the final effluent treated by the coupled system comprising modified-Fenton and MD using (b) Fe-citrate and (c) Fe-EDDS as catalyst. The open symbols are related to the effluent upon iron sequestration by EDDS. The toxicity was measured after 5, 15, and 30 minutes of contact with the *Vibrio fischeri* culture. The dash horizontal line indicates the point where 50% of acute toxicity is

reached; the lines connecting the data points are only intended as guides for the eye.

Specifically, the modified Fenton processes achieved near complete degradation of all
WMC, including MTBE that was instead not removed by the classic Fenton reaction; see also
the chromatograms and percentage of substrates removal at different oxidation times in
Figures S8, S9, and S10 (SM).

451 Despite the higher percentage of degradation of the target contaminants, the modified 452 Fenton reaction did not show further beneficial effect on the MD process in terms of productivity and effluent quality compared to traditional Fenton; see Figure S11 of the SM. 453 454 With regard to EC_{50} , the quality of the effluent treated with the coupled system including modified Fenton and MD clearly increased compared to the PW treated only with MD, 455 specifically, from a dilution factor of 0.094 \pm 0.009 to 0.13 \pm 0.02 and to 0.14 \pm 0.04 in the 456 case of Fe-citrate and Fe-EDDS, respectively (see Figure 4b and c). However, the EC₅₀ was 457 lower compared with the residual toxicity observed in the system that included the traditional 458 459 Fenton oxidation (0.25 ± 0.03), possibly due to the formation of more toxic by-products, e.g., 460 N-(2-aminoethyl) aspartic acid (Zhang et al., 2016). When the contribution to toxicity of the residual iron (residual iron was 0.27 and 1.30 mM for Fe-citrate and Fe-EDDS, respectively) 461 462 was prevented by EDDS addition, the acute toxicity of the effluent clearly decreased (from ~90% to ~70% in both cases) and the EC₅₀ further increased (to 0.4 and 0.5 for Fe-citrate and 463 Fe-EDDS, respectively). At the end of the process, the effluent treated with the Fe-EDDS-464 based Fenton system presented a slightly better quality in terms of toxicity than that treated 465 466 with the Fe-citrate system, consistently with the slightly higher degradation efficiency of the 467 former compared to the latter.

Figure 5 offers a final evaluation of the toxicity parameters from the various treatments. The traditional Fenton coupled with the MD desalination was the best in terms of effluent toxicity. However, the modified Fenton-MD coupled systems were able to overcome some of the practical limitations of the traditional Fenton while still providing an effluent with suitable quality for safe discharge as sewage (toxicity <80%). The modified Fenton oxidation
may be more advisable for applications whereby easier operational tasks and lower sludge
production are important, such as offshore.



475

476 **Figure 5** Summary of the residual toxicity and EC_{50} values of the various effluents treated 477 with both MD and oxidation processes after 30 minutes of contact with the *Vibrio fischeri* 478 culture. The dash line is relative to the value of 80% of acute toxicity, namely, the regulated 479 legislative limit for safe discharge into the sewage system in Italy.

480

481 **4. Concluding remarks, challenges, and implications**

482 This work evaluated the impact of traditional and modified Fenton oxidations for the abatement of highly toxic organic contaminants and as pre-treatment options for the 483 484 subsequent desalination of hypersaline produced waters by membrane distillation. Fe-citrate and Fe-EDDS were used as inexpensive, easy to handle, environmentally friendly and 485 486 biodegradable systems in the modified Fenton processes. All the oxidative processes 487 provided relatively high degradation efficiency toward target contaminants also in the presence of typical scavengers of the Fenton reaction, namely, chloride and humic acids. The 488 observable beneficial effects of an oxidative pre-treatment were not evident in terms of MD 489

490 productivity, fouling, or wetting, but directly translated into lower permeation of organics 491 during distillation and in a significantly lower toxicity of the desalinated effluent. 492 Specifically, the EC_{50} and the acute toxicity (inhibition % of target organisms) were used as 493 indexes for the evaluation of the quality of the final effluent.

494 In conclusion, the coupled oxidation-MD systems to treat PW allow a much less toxic 495 effluent compared to the initial PW toxicity. All the final effluents obtained in this study may 496 be safely discharged in the sewage system and treated within the civil wastewater treatment 497 trains, according to the Italian legislation. The oxidation processes are promising for PW treatments since they are able to degrade the toxic initial target contaminants almost 498 completely. Moreover, the modified Fenton process is able to effectively treat PW while 499 overcoming the practical limitations of traditional Fenton (sludge production and acidic pH). 500 501 The Fenton processes add iron in the effluent environment, this metal being associated with 502 intrinsic toxicity; thus, the toxicity associated with the residual iron needs to be properly 503 addressed and managed in real plants. A wide range of options are available to remove iron 504 and the best-fitting one should be selected case-by-case. Moreover, the oxidations struggle in 505 achieving complete mineralization of all the organics in PW, thus some potentially toxic byproducts may be formed. Therefore, an accurate monitoring of the by-products may be 506 507 necessary, possibly also enforcing some control on the reaction pathway, to evaluate in each case the possibility to safely discharge the final effluent to a civil wastewater treatment plant 508 509 and in case to specifically target the most troublesome by-products in a tertiary treatment 510 step.

512 CRediT authorship contribution statement

Giulio Farinelli: Conceptualization, Data curation, Formal analysis, Investigation, 513 Methodology, Visualization, Writing - original draft. Marco Coha: Data Curation, Formal 514 analysis, Investigation, Methodology, Visualization, Writing - review & editing. Marco 515 Minella: Data curation, Validation, Methodology, Writing - review & editing. Debora 516 Fabbri: Resources, Supervision, Writing - review & editing. Marco Pazzi: Formal analysis, 517 518 Methodology. Davide Vione: Resources, Supervision, Writing - review & editing. Alberto 519 Tiraferri: Funding acquisition, Project administration, Resources, Supervision, Visualization, Writing - review & editing. 520

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522 Declaration of Competing Interest

- 523 The authors declare no competing financial interest.
- 524

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529 Appendix A. Supplementary data.

530 Supplementary material related to this article can be found, in the online version.

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533 **References**

- Abdel-Karim, A., Leaper S., Skuse C., Zaragoza G., Gryta M., Gorgojo P., 2021. Membrane cleaning
 and pretreatments in membrane distillation a review. Chem. Eng. J. 422.
 https://doi.org/10.1016/j.cej.2021.129696
- Adewumi, M. A., Erb J. E., Watson R. W. Initial Design Considerations for a Cost-Effective
 Treatment of Stripper Oil-Well Produced Water. In: P. RJ, R. EF, editors. Produced Water.
 46. Springer, Botsom, MA, 1992, pp. 511-522.
- Ahmadun, F. R., Pendashteh A., Abdullah L. C., Biak D. R. A., Madaeni S. S., Abidin Z. Z., 2009.
 Review of technologies for oil and gas produced water treatment. J. Hazard. Mater. 170, 530551. https://doi.org/10.1016/j.jhazmat.2009.05.044
- 543 Al-Ghouti, M. A., Al-Kaabi M. A., Ashfaq M. Y., Da'na D. A., 2019. Produced water characteristics, 544 treatment and reuse: Α review. J. Water Process Eng. 28, 222-239. 545 https://doi.org/10.1016/j.jwpe.2019.02.001
- 546 APAT, I., CNR. Metodi Ecotossicologici. Manuali e linee guida 29/2003 Serie 8000, 2003.
- Aquiliina, P. Impairment of Gas Well Productivity by Salt Plugging: A Review of Mechanisms,
 Modeling, Monitoring Methods, and Remediation Techniques. SPE Annual Technical
 Conference and Exhibition, San Antonio, Texas, USA, 2012.
- Ariono, D., Purwasasmita M., Wenten I. G., 2016. Brine Effluents: Characteristics, Environmental Impacts, and Their Handling. J. Eng. Technol. Sci. 48, 367-387. https://doi.org/10.5614/j.eng.technol.sci.2016.48.4.1
- Ayed, L., Asses N., Chammem N., Ben Othman N., Hamdi M., 2017. Advanced oxidation process
 and biological treatments for table olive processing wastewaters: constraints and a novel
 approach to integrated recycling process: a review. Biodegradation 28, 125-138.
 https://doi.org/10.1007/s10532-017-9782-0
- Bessa, E., Sant Anna G. L., Dezotti M., 2001. Photocatalytic/H2O2 treatment of oil field produced
 waters. Appl. Catal. B-Environ. 29, 125-134. https://doi.org/10.1016/S0926-3373(00)00199-5
- 559 Canedo-Arguelles, M., Kefford B., Schafer R., 2019. Salt in freshwaters: causes, effects and prospects 560 issue. introduction to the theme Philos. Τ. R. Soc. В 374. 561 https://doi.org/10.1098/Rstb.2018.0002
- Chahbane, N., Popescu D. L., Mitchell D. A., Chanda A., Lenoir D., Ryabov A. D., et al., 2007. Fe III-TAML-catalyzed green oxidative degradation of the azo dye Orange II by H2O2 and
 organic peroxides: products, toxicity, kinetics, and mechanisms. Green Chem. 9, 49-57.
 https://doi.org/10.1039/b604990g
- Chang, H. Q., Li T., Liu B. C., Vidic R. D., Elimelech M., Crittenden J. C., 2019a. Potential and
 implemented membrane-based technologies for the treatment and reuse of flowback and
 produced water from shale gas and oil plays: A review. Desalination 455, 34-57.
 https://doi.org/10.1016/j.desal.2019.01.001
- 570 Chang, H. Q., Liu B. C., Wang H. Z., Zhang S. Y., Chen S., Tiraferri A., et al., 2019b. Evaluating the
 571 performance of gravity-driven membrane filtration as desalination pretreatment of shale gas
 572 flowback and produced water. J. Membr. Sci. 587.
 573 https://doi.org/10.1016/J.Memsci.2019.117187
- 574 Chen, L., Wang D., Long C., Cui Z. X., 2019. Effect of biodegradable chelators on induced
 575 phytoextraction of uranium- and cadmium-contaminated soil by Zebrina pendula Schnizl. Sci.
 576 Rep. 9. https://doi.org/10.1038/S41598-019-56262-9
- 577 Chen, W., Zhang F. F., Hong J. L., Shi W. X., Feng S. T., Tan X. F., et al., 2016. Life cycle toxicity
 578 assessment on deep-brine well drilling. J. Clean. Prod. 112, 326-332.
 579 https://doi.org/10.1016/j.jclepro.2015.07.062
- Chen, Y. M. L., Wang Z. X., Jennings G. K., Lin S. H., 2017. Probing Pore Wetting in Membrane
 Distillation Using Impedance: Early Detection and Mechanism of Surfactant-Induced
 Wetting. Environ. Sci. Tech. Let. 4, 505-510. https://doi.org/10.1021/acs.estlett.7b00372
- Coha, M., Farinelli G., Tiraferri A., Minella M., Vione D., 2021. Advanced oxidation processes in the
 removal of organic substances from produced water: Potential, configurations, and research
 needs. Chem. Eng. J. 414. https://doi.org/10.1016/j.cej.2021.128668

- da Silva, S. S., Chiavone-Filho O., de Barros Neto E. L., Foletto E. L., 2015. Oil removal from
 produced water by conjugation of flotation and photo-Fenton processes. J. Environ. Manage.
 147, 257-263. https://doi.org/10.1016/j.jenvman.2014.08.021
- Dalmacija, B., Karlovic E., Tamas Z., Miskovic D., 1996. Purification of high-salinity wastewater by
 activated sludge process. Water Res. 30, 295-298. https://doi.org/10.1016/0043 1354(95)00170-0
- 592 De Laat, J., Gallard H., 1999. Catalytic decomposition of hydrogen peroxide by Fe(III) in
 593 homogeneous aqueous solution: Mechanism and kinetic modeling. Environ. Sci. Technol. 33,
 594 2726-2732. https://doi.org/10.1021/Es981171v
- 595 Diya'uddeen, B. H., Aziz A. R. A., Daud W. M. A. W., 2012. On the Limitation of Fenton Oxidation
 596 Operational Parameters: A Review. Int. J. Chem. React. Eng. 10.
 597 https://doi.org/10.1515/1542-6580.R2
- Donaldson, E. C., Thomas R. D., Lorenz P. B., 1969. Wettability Determination and Its Effect on Recovery Efficiency. Soc. Pet. Eng. J. 9, 13-20. https://doi.org/10.2118/2338-PA
- Estrada, J. M., Bhamidimarri R., 2016. A review of the issues and treatment options for wastewater
 from shale gas extraction by hydraulic fracturing. Fuel 182, 292-303.
 https://doi.org/10.1016/j.fuel.2016.05.051
- Farinelli, G., Minella M., Pazzi M., Giannakis S., Pulgarin C., Vione D., et al., 2020. Natural iron
 ligands promote a metal-based oxidation mechanism for the Fenton reaction in water
 environments. J. Hazard. Mater. 393. https://doi.org/10.1016/j.jhazmat.2020.122413
- Farinelli, G., Minella M., Sordello F., Vione D., Tiraferri A., 2019. Metabisulfite as an Unconventional Reagent for Green Oxidation of Emerging Contaminants Using an Iron-Based Catalyst. ACS Omega 4, 20732-20741. https://doi.org/10.1021/acsomega.9b03088
- Franken, A. C. M., Nolten J. A. M., Mulder M. H. V., Bargeman D., Smolders C. A., 1987. Wetting
 Criteria for the Applicability of Membrane Distillation. J. Membr. Sci. 33, 315-328.
 https://doi.org/10.1016/S0376-7388(00)80288-4
- Goldstone, J. V., Pullin M. J., Bertilsson S., Voelker B. M., 2002. Reactions of hydroxyl radical with
 humic substances: Bleaching, mineralization, and production of bioavailable carbon
 substrates. Environ. Sci. Technol. 36, 364-372. https://doi.org/10.1021/es0109646
- Goncalves, N. P. F., Minella M., Fabbri D., Calza P., Malitesta C., Mazzotta E., et al., 2020. Humic
 acid coated magnetic particles as highly efficient heterogeneous photo-Fenton materials for
 wastewater treatments. Chem. Eng. J. 390. https://doi.org/10.1016/J.Cej.2020.124619
- Gonzalez, D., Amigo J., Suarez F., 2017. Membrane distillation: Perspectives for sustainable and
 improved desalination. Renew. Sust. Energ. Rev. 80, 238-259.
 https://doi.org/10.1016/j.rser.2017.05.078
- Haber, F., Weiss J., Pope W. J., 1934. The catalytic decomposition of hydrogen peroxide by iron salts.
 Proc. R. Soc. Lond. A Math. Phys. Sci. 147, 332-351.
 https://doi.org/10.1098/rspa.1934.0221
- Han, L., Tan Y. Z., Netke T., Fane A. G., Chew J. W., 2017. Understanding oily wastewater treatment
 via membrane distillation. J. Membr. Sci. 539, 284-294.
 https://doi.org/10.1016/j.memsci.2017.06.012
- Harvey, A. E., Smart J. A., Amis E. S., 1955. Simultaneous Spectrophotometric Determination of Iron(II) and Total Iron with 1,10-Phenanthroline. Anal. Chem. 27, 26-29. https://doi.org/10.1021/ac60097a009
- Horseman, T., Yin Y., Christie K. S., Wang Z., Tong T., Lin S., 2021a. Wetting, Scaling, and Fouling
 in Membrane Distillation: State-of-the-Art Insights on Fundamental Mechanisms and
 Mitigation Strategies. ACS ES&T Eng. 1, 117-140.
 https://doi.org/10.1021/acsestengg.0c00025
- 634Horseman, T., Yin Y., Christie K. S. S., Wang Z., Tong T., Lin S., 2021b. Wetting, Scaling, and635Fouling in Membrane Distillation: State-of-the-Art Insights on Fundamental Mechanisms and636MitigationStrategies.ACSES&TEng.1,117-140.637https://doi.org/10.1021/acsestengg.0c00025
- Howell, J. A., 2004. Future of membranes and membrane reactors in green technologies and for water
 reuse. Desalination 162, 1-11. https://doi.org/10.1016/S0011-9164(04)00021-9

- Igunnu, E. T., Chen G. Z., 2014. Produced water treatment technologies. Int. J. Low-Carbon Technol.
 9, 157-177. https://doi.org/10.1093/ijlct/cts049
- Jiménez, S., Andreozzi M., Micó M. M., Álvarez M. G., Contreras S., 2019. Produced water treatment
 by advanced oxidation processes. Chemosphere 666, 12-21.
 https://doi.org/10.1016/j.scitotenv.2019.02.128
- Jiménez, S., Micó M. M., Arnaldos M., Ferrero E., Malfeito J. J., Medina F., et al., 2017. Integrated
 processes for produced water polishing: Enhanced flotation/sedimentation combined with
 advanced oxidation processes. Sci. Total Environ. 168, 309-317.
 https://doi.org/10.1016/j.chemosphere.2016.10.055
- 649Jimenez, S., Mico M. M., Arnaldos M., Medina F., Contreras S., 2018. State of the art of produced650watertreatment.651https://doi.org/10.1016/j.chemosphere.2017.10.139
- Jin, Y., Davarpanah A., 2020. Using Photo-Fenton and Floatation Techniques for the Sustainable
 Management of Flow-Back Produced Water Reuse in Shale Reservoirs Exploration. Water
 Air Soil Pollut. 231. https://doi.org/10.1007/s11270-020-04812-7
- Kabyl, A., Yang M., Abbassi R., Li S. H., 2020. A risk -based approach to produced water
 management in offshore oil and gas operations. Process Saf. Environ. 139, 341-361.
 https://doi.org/10.1016/j.psep.2020.04.021
- Kargbo, D. M., Wilhelm R. G., Campbell D. J., 2010. Natural Gas Plays in the Marcellus Shale:
 Challenges and Potential Opportunities. Environ. Sci. Technol. 44, 5679-5684. https://doi.org/10.1021/es903811p
- Khatri, N., Tyagi S., Rawtani D., 2017. Recent strategies for the removal of iron from water: A review. J. Water. Process Eng. 19, 291-304. https://doi.org/10.1016/j.jwpe.2017.08.015
- Kiwi, J., Lopez A., Nadtochenko V., 2000. Mechanism and kinetics of the OH-radical intervention
 during fenton oxidation in the presence of a significant amount of radical scavenger (Cl-).
 Environ. Sci. Technol. 34, 2162-2168. https://doi.org/10.1021/Es991406i
- Klavins, M., Purmalis O., 2010. Humic substances as surfactants. Environ. Chem. Lett. 8, 349-354.
 https://doi.org/10.1007/s10311-009-0232-z
- Kleinitz, W., Koehler M., Dietzsch G. The Precipitation of Salt in Gas Producing Wells. SPE
 European Formation Damage Conference, The Hague, Netherlands, 2001.
- Lester, Y., Ferrer I., Thurman E. M., Sitterley K. A., Korak J. A., Aiken G., et al., 2015.
 Characterization of hydraulic fracturing flowback water in Colorado: Implications for water
 treatment. Sci. Total Environ. 512, 637-644. https://doi.org/10.1016/j.scitotenv.2015.01.043
- Li, X. M., Zhao B., Wang Z., Xie M., Song J., Nghiem L. D., et al., 2014. Water reclamation from
 shale gas drilling flow-back fluid using a novel forward osmosis-vacuum membrane
 distillation hybrid system. Water Sci. Technol. 69, 1036-1044.
 https://doi.org/10.2166/wst.2014.003
- Lin, S. H., Nejati S., Boo C., Hu Y. X., Osuji C. O., Ehmelech M., 2014. Omniphobic Membrane for
 Robust Membrane Distillation. Environ. Sci. Tech. Let. 1, 443-447.
 https://doi.org/10.1021/ez500267p
- Liu, Y., Lu H., Li Y., Xu H., Pan Z., Dai P., et al., 2021. A review of treatment technologies for
 produced water in offshore oil and gas fields. Sci. Total Environ. 775.
 https://doi.org/10.1016/j.scitotenv.2021.145485
- Ma, H. Z., Wang B., 2006. Electrochemical pilot-scale plant for oil field produced wastewater by
 M/C/Fe electrodes for injection. J. Hazard. Mater. 132, 237-243.
 https://doi.org/10.1016/j.jhazmat.2005.09.043
- McCormack, P., Jones P., Hetheridge M. J., Rowland S. J., 2001. Analysis of oilfield produced waters
 and production chemicals by electrospray ionisation multi-stage mass spectrometry (ESI MSn). Water Res. 35, 3567-3578. https://doi.org/10.1016/S0043-1354(01)00070-7
- Messele, S. A., Bengoa C., Stuber F. E., Giralt J., Fortuny A., Fabregat A., et al., 2019. Enhanced
 Degradation of Phenol by a Fenton-Like System (Fe/EDTA/H2O2) at Circumneutral pH.
 Catalysts 9. https://doi.org/10.3390/Catal9050474
- Miklos, D. B., Remy C., Jekel M., Linden K. G., Drewes J. E., Hubner U., 2018. Evaluation of advanced oxidation processes for water and wastewater treatment - A critical review. Water Res 139, 118-131. https://doi.org/10.1016/j.watres.2018.03.042

- Mohammad-Pajooh, E., Weichgrebe D., Cuff G., Tosarkani B. M., Rosenwinkel K. H., 2018. On-site
 treatment of flowback and produced water from shale gas hydraulic fracturing: A review and
 economic evaluation. Chemosphere 212, 898-914.
 https://doi.org/10.1016/j.chemosphere.2018.08.145
- Neff, J. M., Sauer T. C., Maciolek N. Composition, Fate and Effects of Produced Water Discharges to
 Nearshore Marine Waters. In: P. RJ, R. EF, editors. Produced Water. 46. Springer, Boston,
 MA, 1992, pp. 371-385.
- Olsson, O., Weichgrebe D., Rosenwinkel K. H., 2013. Hydraulic fracturing wastewater in Germany:
 composition, treatment, concerns. Environ. Earth Sci. 70, 3895-3906.
 https://doi.org/10.1007/s12665-013-2535-4
- Pasternak, G., Kolwzan B., 2013. Surface tension and toxicity changes during biodegradation of
 carbazole by newly isolated methylotrophic strain Methylobacterium sp GPE1. Int. Biodeter.
 Biodegr. 84, 143-149. https://doi.org/10.1016/j.ibiod.2012.07.021
- Rezaei, M., Warsinger D. M., Lienhard V. J. H., Samhaber W. M., 2017. Wetting prevention in membrane distillation through superhydrophobicity and recharging an air layer on the membrane surface. J. Membr. Sci. 530, 42-52. https://doi.org/10.1016/j.memsci.2017.02.013
- Ricceri, F., Giagnorio M., Farinelli G., Blandini G., Minella M., Vione D., et al., 2019. Desalination
 of Produced Water by Membrane Distillation: Effect of the Feed Components and of a Pretreatment by Fenton Oxidation. Sci. Rep. 9, 14964. https://doi.org/10.1038/s41598-01951167-z
- Ruiz-Aguirre, A., Polo-López M. I., Fernández-Ibáñez P., Zaragoza G., 2017. Integration of
 Membrane Distillation with solar photo-Fenton for purification of water contaminated with
 Bacillus sp. and Clostridium sp. spores. Sci. Total Environ. 595, 110-118.
 https://doi.org/10.1016/j.scitotenv.2017.03.238
- Shaffer, D. L., Chavez L. H. A., Ben-Sasson M., Castrillon S. R. V., Yip N. Y., Elimelech M., 2013.
 Desalination and Reuse of High-Salinity Shale Gas Produced Water: Drivers, Technologies,
 and Future Directions. Environ. Sci. Technol. 47, 9569-9583.
 https://doi.org/10.1021/es401966e
- Shang, W., Tiraferri A., He Q. P., Li N. W., Chang H. Q., Liu C., et al., 2019. Reuse of shale gas
 flowback and produced water: Effects of coagulation and adsorption on ultrafiltration, reverse
 osmosis combined process. Sci. Total Environ. 689, 47-56.
 https://doi.org/10.1016/j.scitotenv.2019.06.365
- Shokrollahzadeh, S., Golmohammad F., Naseri N., Shokouhi H., Arman-mehr M., 2012. Chemical
 oxidation for removal of hydrocarbons from gas-field produced water. Procedia Engineer 42,
 942-947. https://doi.org/10.1016/j.proeng.2012.07.487
- Siyal, M. I., Lee C., Kim J. O., 2017. Degradation of humic acid by ZVI based sono-Fenton as a
 pretreatment for reducing organic fouling likely in membrane distillation. Desalin. Water
 Treat. 99, 167-171. https://doi.org/10.5004/dwt.2017.21652
- Tandy, S., Ammann A., Schulin R., Nowack B., 2006. Biodegradation and speciation of residual SS ethylenediaminedisuccinic acid (EDDS) in soil solution left after soil washing. Environ.
 Pollut. 142, 191-199. https://doi.org/10.1016/j.envpol.2005.10.013
- Tang, P., Li J. L., Li T., Tian L., Sun Y., Xie W. C., et al., 2021. Efficient integrated module of gravity driven membrane filtration, solar aeration and GAC adsorption for pretreatment of shale gas wastewater. J. Hazard. Mater. 405. https://doi.org/10.1016/j.jhazmat.2020.124166
- Van Devivere, P. C., Saveyn H., Verstraete W., Feijtel T. C. J., Schowanek D. R., 2001.
 Biodegradation of metal-[S,S]-EDDS complexes. Environ. Sci. Technol. 35, 1765-1770.
 https://doi.org/10.1021/es0001153
- Vesterkvist, P. S. M., Misiorek J. O., Spoof L. E. M., Toivola D. M., Meriluoto J. A. O., 2012.
 Comparative Cellular Toxicity of Hydrophilic and Hydrophobic Microcystins on Caco-2
 Cells. Toxins 4, 1008-1023. https://doi.org/10.3390/toxins4111008
- Vione, D., Merlo F., Maurino V., Minero C., 2004. Effect of humic acids on the Fenton degradation
 of phenol. Environ. Chem. Lett. 2, 129-133. https://doi.org/10.1007/s10311-004-0086-3
- Voelker, B. M., Sulzberger B., 1996. Effects of fulvic acid on Fe(II) oxidation by hydrogen peroxide.
 Environ. Sci. Technol. 30, 1106-1114. https://doi.org/10.1021/Es9502132

- Wang, Z. X., Chen Y. M. L., Sun X. M., Duddu R., Lin S. H., 2018. Mechanism of pore wetting in membrane distillation with alcohol vs. surfactant. J Membrane Sci 559, 183-195. https://doi.org/10.1016/j.memsci.2018.04.045
- Zhai, J., Ma H., Liao J., Rahaman M. H., Yang Z., Chen Z., 2018. Comparison of Fenton, ultraviolet–
 Fenton and ultrasonic–Fenton processes on organics and colour removal from pre-treated
 natural gas produced water. Int. J. Environ. Sci. Technol. 15, 2411-2422.
 https://doi.org/10.1007/s13762-017-1604-8
- Zhang, Y., Klamerth N., Chelme-Ayala P., Gamal El-Din M., 2016. Comparison of Nitrilotriacetic
 Acid and [S,S]-Ethylenediamine-N,N'-disuccinic Acid in UV–Fenton for the Treatment of Oil
 Sands Process-Affected Water at Natural pH. Environ. Sci. Technol. 50, 10535-10544.
 https://doi.org/10.1021/acs.est.6b03050