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2	Diamondized carbon nanoarchitectures as electrocatalytic material for sulfate-based
3	oxidizing species electrogeneration
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26 Abstract

The introduction of nanotechnology seems to be an imperative factor to intensify the synergic 27 effects of electrocatalytic materials to produce strong oxidant species or to increase the active 28 sites on their surfaces as well as to enhance the conversion yield in a fuel cell, high-added 29 value products, electrolytic treatment for environmental protection or the detection limit in 30 electroanalysis. Recently, a new type of 3D-diamond electrodes was developed with boron-31 doped carbon nanowalls (B:CNW), which was manufactured using the microwave plasma-32 33 assisted chemical vapor deposition (CVD) process, improving the charge transfer and enhancing the electrochemical performance. The applicability of a BDD/boron-doped carbon 34 nanowalls (BDD/B:CNW) anodes to degrade organic pollutants has been already 35 investigated; however, no attempts at the electrosynthesis of oxidizing species using these 36 diamond-carbon nanostructures have been reported yet. Therefore, the electrosynthesis of 37 sulfate-based oxidizing species was studied here to answer relevant questions from both 38 39 fundamental and practical point-of-view. The results demonstrated that persulfate was 40 efficiently produced at the BBD plate, while that the ion-radical sulfate could be the most important oxidant at BDD/B:CNW anode when compared to other electrocatalytic materials, 41 including BDD surfaces. Persulfate concentrations ranged from 3 to 6 µM, depending on the 42 applied current density (2.5, 5.0, and 15 mA cm⁻²), at diamond electrodes. A dye-model 43 44 pollutant - methyl orange (MO) was degraded below the limit of detection within 45 min 45 using BDD/B:CNW when in-situ sulfate-based oxidizing species were electrogenerated. These kinds of 3D-diamond-carbon nanostructures are thus promising as novel 46 electrocatalyst for various catalytic applications in the environmental and energy fields. 47

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- 49 Keywords: boron-doped diamond electrode, sulfate-based oxidizing species, diamond-
- 50 carbon nanoarchitectures, electrocatalyst, electrogeneration.

51 **1. Introduction**

The scientific community has shown a preference to apply EO as an environment-friendly 52 electrochemical process because of its uncomplicatedness and acceptance among the 53 electrochemical advanced oxidation processes (EAOPs), in terms of electrolytic technology, 54 amenability, environmental compatibility, automation, no reagents use, renewable energies-55 driven and in-situ production of higher concentration of oxidants [1]. In the last decades, EO 56 has competently demonstrated its effectiveness to mineralize different organic compounds in 57 58 a great variety of water matrices or wastewaters [2]. Nevertheless, the extension of mineralization/degradation of these pollutants and their concentration as well as the type of 59 the electrogenerated oxidants strongly depends on the operating parameters and the nature of 60 the anode electrocatalytic material [3,4]. 61

Various electrodes have been used in EO, like Ir-, Pt-, Ru-, Ti-, Pb- and Sn- oxide-based 62 anodes as well as synthetic diamond films [5,6]. However, boron-doped diamond (BDD) 63 64 anodes are considered the most effective electrocatalytic materials for degrading/mineralizing refractory/priority pollutants and waterborne agents [1,2,7–9]. It is 65 due to their effective *in-situ* production of oxidizing agents, at higher concentrations, mainly 66 reactive oxygen species (ROS) (Eq. (1)-(5)) [9]. Also, peroxodisulfate $(S_2O_8^{2-})$, 67 peroxodicarbonate $(C_2O_6^{2-})$ and peroxodiphosphate $(P_2O_8^{4-})$ can be electrochemically 68 69 formed with ROS from the oxidation of sulfate or bisulfate by Eq. (6), bicarbonate by Eq. (7) 70 and phosphate by Eq. (8) at the BDD surface [10,11].

71 BDD + H₂O
$$\rightarrow$$
 BDD('OH) + H⁺ + e⁻ (1)

72
$$^{\circ}OH \rightarrow ^{\circ}O + H^{+} + e^{-}$$
 (2)

$$73 \qquad 2^{\bullet}O \rightarrow O_2 \tag{3}$$

$$74 \qquad 2^{\bullet}OH \rightarrow H_2O_2 \tag{4}$$

75
$$O_2 + O \rightarrow O_3$$
 (5)

76
$$2SO_4^{2-} \rightarrow 2SO_4^{-} \rightarrow S_2O_8^{2-} + 2e^-$$
 (6)

77
$$2\text{HCO}_3^- \rightarrow \text{C}_2\text{O}_6^{2-} + 2\text{H}^+ + 2\text{e}^-$$
 (7)

78
$$2PO_4^{3-} \rightarrow P_2O_8^{4-} + 2e^-$$
 (8)

79 As already scientifically stated, the effective electrosynthesis of oxidizing species requires an electrode with a large surface area, high electrocatalytic activities, long-term stability, as 80 well as a low cost of materials [5,12]. Thus, the introduction of nanoarchitectures seems to 81 be an essential factor to intensify the synergic effects of electrocatalytic materials to produce 82 strong oxidant species or to increase the active sites on their surfaces [13]. Novel 83 nanostructured materials can exhibit better electrochemical properties than their bulk 84 85 counterparts with a specific pore dimension distribution and well-defined structuremorphology, maximizing the exposed-surface area and minimizing the mass transport 86 phenomena [14-16]. Thus, the advances in the preparation of highly sophisticated 87 88 electrocatalytic nanomaterials may lead to further improvements in the mineralization of organic pollutants as well as the electrosynthesis of oxidants, making them suitable for smart 89 90 water solutions [17]. These technologies will play a key role in achieving the Sustainable 91 Development Goal 6 (SDG6) since these represent a substantial opportunity if their implementation is carried out to guarantee sustainability and increase competence in water 92 93 management (to treat and distribute water for human use) [18].

Recently, a versatile electrode was discovered, called the "3D-nanostructured BDD/B:CNW"
anode. This 3D-diamondized-carbon nanostructure possesses extraordinary electrochemical
properties (e.g., peculiar morphology, specific surface area and the larger extends of exposed

edges with a greater number of functional groups and chemisorbed heteroatoms, in particular, 97 oxygen), allowing a higher reactivity of the carbonaceous nanostructure for environmental, 98 fuel and sensors applications, compared to 2D-BDD electrode [19–21]. However, no 99 information concerning the effectiveness of this 3D-nanostructured BDD/B:CNW anode to 100 electrogeneration oxidants has been reported yet. Thus, for the first time, the electrosynthesis 101 of sulfate-based oxidants species has been studied by using a 3D-nanoarchitectured 102 BDD/B:CNW anode to answer relevant questions from both fundamental and practical point-103 104 of-view. An organic model compound was also chosen to prove the oxidative efficiency of *in-situ* $S_2O_8^{2-}$ electrogeneration when compared with other electrode materials. The results 105 are discussed considering the existing literature on nanostructured electrodes for 106 electrocatalytic advanced oxidation processes. 107

108

109 **2.** Materials and Methods

110 **2.1.** Materials

All reagents were of analytical grade and used without further purification. The water was obtained from purified water (Milli-Q system, with resistivity ≥ 18 M Ω cm at 25 °C). Methyl orange (MO) purity was 95% and it was used as received.

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2.2. Synthesis and characterization of BDD/B:CNW

BDD and BDD/BCNW were synthesized using the MWPECVD system (SEKI Technotron AX5400S, Japan). Thin films were grown on niobium substrates for a total of 5 h, resulting in 3 µm of thickness. The detailed parameters of the thin film synthesis can be found elsewhere, for BDD [ref1] and for the BDD:BCNW in Pierpaoli et al [ref2, ref3]BCNW in Siuzdak *et al.* [22], Dettlaff *et al.* [23], and Sobaszek *et al.* [24]. The surface characteristics

of the BDD/B:CNW was carried out using a Hitachi model TM 3000 top microscope with a 121 high sensitivity semiconductor backscattered electron detector to obtain scanning electron 122 microscopy (SEM) images, using a significant magnification and operating at 15 kV voltage 123 acceleration with a tungsten filament. The SEM images were taken under a vacuum at 124 controlled temperatures and the energy dispersive X-ray (EDS) spectrum was obtained to 125 analyze the concentrations of the elements in the diamond-carbon nanostructures. Raman 126 spectra were recorded in the 50–3200 cm^{-1} range, upon excitation by a 532 nm laser; the 127 spectral resolution was equal to 2.9 cm^{-1} , in the range of $50-2000 \text{ cm}^{-1}$ and 2.4 cm^{-1} in the 128 range of 2000–3200 cm⁻¹ with an integration time of 5 s (20 averages), with a diffraction 129 grating of 300 lines/mm². 130

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2.3. Electrochemical measurements

Experiments were carried out in a conventional three-electrode system in Pyrex material 132 with a capacity of 10 mL, and measurements were performed between 0.0 and +1.8 V at 10 133 mV s⁻¹ for quasi-steady polarization curves in 0.05 M of Na₂SO₄. BDD and BDD/B:CNW 134 anodes have significant areas in cm^2 , however, an exposed geometric area of ca. 0.075 cm^2 135 was used as the working electrode for electrochemical measurements, while this area was 136 increased for bulk electrolysis (see next subsection). A platinum wire and an Ag/AgCl (3 M) 137 were employed as the auxiliary and reference electrodes, respectively. The electroactive 138 139 surface area (Areal) and the differential capacitance (CDL) were experimentally estimated [25– 27], according to Eq. 1. To do that, a potential range was selected, for BDD and BDD/B:CNW 140 141 electrodes, where a non-Faradaic current response is registered. This range was ≈ 0.2 V potential window centered on the open-circuit potential of the system. Then, cyclic 142 143 voltammetry (CV) measurements were conducted in static solution by sweeping the potential across the non-Faradaic region and back at 9 different scan rates (0.02, 0.04, 0.06, 0.08, 0.10, 0.12, 0.14, 0.16, 0.18 and 0.20 V s⁻¹) at $25\pm1^{\circ}$ C in 0.05 M of Na₂SO₄. For each measurement, five voltammetric profiles were recorded, and the last curve was always selected to be used for plotting the graphics in this work.

148
$$A_{real} = \frac{C_{DL}}{C^*} = \frac{\mu F}{\frac{\mu F}{real \ cm^2}} = cm^2$$
(1)

In Eq. 1, C_{DL} is the differential capacitance of the electrode-electrolyte interface and C* is 149 approximately 60 μ F (real cm²)⁻¹ which is a reference value for porous materials, regardless of 150 Experiments 151 its composition [25,27]. were performed with an Autolab potentiostat/galvanostat model PGSTAT302N. 152

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2.4. Spectrophotometric analysis

For UV-vis spectrophotometric measurements to determine the production of persulfate, a Shimadzu spectrophotometer model 1800 was employed by using *in-situ* chemical oxidation (ISCO) method [28,29], operating from 190 to 800 nm. A single quartz cuvette was used with an optical path of 1 cm. It is important to remark that persulfate generation can be followed by ISCO method without H_2O_2 interference because the analysis of absorption spectra of a yellow color solution resulting from the reaction of persulfate and iodide in the presence of sodium bicarbonate reveals a clear absorbance spectrum at 352 nm.

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2.5. Electrochemical determination of methyl orange concentration

Electroanalytical measurements were carried out in a conventional three-electrode system in Pyrex material with a capacity of 25 mL for determining MO concentrations by differential pulse voltammetry (DPV) analysis. BDD anode (silicon substrate, 500 ppm of boron doping,

2-3 μ m of thickness), with an exposed geometric area of 0.28 cm², was used as the working 167 electrode, while a platinum wire and Ag/AgCl (KCl 3 M) were employed as the auxiliary 168 and reference electrodes, respectively. DPV experiments were performed with an Autolab 169 PGSTAT 320N model to establish an analytical curve by measuring different concentrations 170 of MO in 0.05 M Na₂SO₄. Other details of the procedure are as follows: equilibration time, 171 5 s; modulation time, 0.05 s; interval time, 0.5 s; initial potential, 0 V; end potential, +1.8 V; 172 step potential, 25 mV; and modulation amplitude, 100 mV. For determining MO 173 174 concentration at each predetermined time during EO tests, the samples were spiked with a 175 known quantity of a standard solution, and the standard addition method was implemented.

176

¹⁷⁷ **2.6.** Bulk experiments

178 Experiments were carried out using a bulk electrochemical cell under stirring agitation which 179 consisted of several electrodes (anodes such as BDD plate, BDD/B:CNW, TiPt, Pb/PbO₂, 180 Ti/TiO₂-nanotubes/PbO₂ and Ti as cathode, respectively) of 1.5 cm of diameter (geometrical 181 area $\approx 7.07 \text{ cm}^2$). Electrolysis of 50 mL of a synthetic solution containing 0.05 M Na₂SO₄ as 182 the supporting electrolyte was performed by applying 2.5, 10, and 15 mA cm^{-2} for 150 min 183 to prepare sulfate-based oxidizing species. Sulfate concentration was chosen based on the 184 existing literature to produce a significant concentration of persulfate [29,30]. During all 185 experiments, samples of the sulfate-based oxidizing solutions were collected at the end of the 186 electrolysis, and subsequently, these were analyzed by the ISCO-spectrophotometric method, 187 determining the final sulfate-based oxidants concentration produced. For the EO of an 188 organic target compound, 25 mL of 0.05 M of Na₂SO₄ containing 10 mg L⁻¹ of MO were 189 electrochemically treated by mixing with magnetic stirring for 40 min and aliquots were withdrawn at predetermined times to determine chemical oxygen demand (COD) and the
concentration of the target pollutant by the electroanalytic procedure. COD levels were
measured by using pre-dosed reagents (HANNA® vials) in 2 mL samples. Samples for COD
determination were digested in a thermal reactor (HANNA instrument) at 150°C for 2 h.
Subsequently, the samples initial and final, for all oxidation tests, were analyzed on a
spectrophotometer (Hanna HI 83099), at 25°C.

- 196
- ¹⁹⁷ **3. Results and Discussion**

198 **3.1.** Characterization of diamond electrode

The BDD/B:CNW electrode morphology presents microscale valleys with a diameter of a 199 200 few micrometers, as observed in the SEM images (Fig. 1a). As previously reported [13,14,31] and confirmed by SEM images; on the one hand, carbon grows vertically in form of multi-201 202 layer graphene walls covered by diamond clusters. Microcavities having an opening in the 203 range between tens and hundreds of nanometres are formed between the nanowall boundaries 204 (Fig. 1a). On the other hand, "nano-flaps" also grow perpendicularly as secondary 205 protuberances to the surface of the nanowalls due to the process of synthesis [13,14,31]. Meanwhile, EDS spectrum evidence that the concentrations of the elements in the diamond-206 carbon nanostructures are preferentially carbon and oxygen, confirming the BDD/B:CNW 207 208 composition (Fig. 1b).

In this context, the CVD process developed by Bogdanowicz's group [14] is an innovative approach to obtaining a carbon nanostructure with concave and convex curvatures in a simple way [13,14,31]. Thus, it was possible to understand that the electrode surface and their composition, as well as the density of electronic states of electrode materials [XXX], could influence during the production of oxidant species. From the Raman spectra (Fig. 1c), the

intense peak at 1332 cm-1 sharply observed for the BDD sample is assigned to sp3-214 hybridized carbon, which decreases in intensity and downshift to 1310 cm-1 for the 215 BDD:BCNW sample, which is typical for typical for heavily boron-doped diamond [ref], 216 217 since the diborance concentration inside the reactor increases as well. Moreover, it is possible to observe the G-band at around 1580 cm-1, arising from the presence of graphitic material 218 (i.e., sp2-C), and the D-band which indicates defects in graphitic structures. Finally, D' is 219 220 another second-order Raman band, appearing as a shoulder peak of the G band, originating 221 from a second-order Raman scattering at the K point, involving one iTO phonon and one 222 defect state [ref].





Fig. 1. (a) SEM images of the BDD/BCNW grown on Nb, (b) EDS spectrum, confirming the
composition of BDD/BCNW electrode: C (94.42 wt.%, 97.40 at.%), O (2.90 wt.%, 2.25 at.%)
and Nb (2.68 wt.%, 0.36 at.%). And (c) Raman spectra for the BDD and BDD:BCNW
samples.

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3.2. Electrochemical measurements

CV curves in a non-faradaic potential range were recorded at different scan rates (0.04, 0.06, 231 0.08, 0.10, 0.12, 0.14, 0.16, 0.18 and 0.20 V s^{-1}) to estimate the electro-active surface of BDD 232 and BDD/B:CNW electrodes. By plotting the current, which was measured in the middle of 233 234 the double-layer region and recorded at different scan rates, versus the scan rate (e.g.: BDD/B:CNW, Fig. 2); a straight line was obtained, which allowed determining the double 235 layer capacitance values for these electrocatalytic materials (Insets in Fig. 2). Then, by using 236 the Eq. (1), the geometric area of the electrodes ($\approx 0.075 \text{ cm}^2$) and the C_{DL} values obtained 237 (5.40 µF and 41.52 µF for BDD and BDD/B:CNW, respectively); the real surface areas were 238 estimated, obtaining 0.216 cm² and 0.692 cm² for BDD and BDD/B:CNW, respectively. It 239 240 is important to remark that, based on existing literature data [26], reference values (C*) for compact and porous materials should be considered to estimate the electroactive surface area, 241 regardless of its composition. Then, 60 μ F (real cm²)⁻¹ was the reference value for 242 BDD/B:CNW, while, 25 μ F (real cm²)⁻¹ for BDD plate due to the most homogeneous surface 243 [26,27,32]. 244



Fig. 2. Cyclic voltammetric (CV) analysis (from 0.04 to 0.20 V s⁻¹) to estimate the capacitance for BDD/B:CNW electrode in 0.05 M of Na₂SO₄ by plotting the current values in the double-layer regions as a function of scan rate. Inset: double-layer capacitance measurements by using CV technique at different scan rates.

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251 Although there are several methods to experimentally determine the real surface area such as 252 hydrogen adsorption, double layer capacitance, surface oxide reduction, underpotential deposition of metals, and adsorbed carbon monoxide stripping [27]; for the case of BDD 253 electrodes, some approaches could be efficiently applied. However, the double-layer 254 capacitance method allows to obtain accurate surface measurements, avoiding significant 255 surface changes due to the oxygen/hydrogen evolution, formation/reduction of oxides, and 256 257 modification of carbon materials [27,32,33]. The double-layer capacitance also allows measuring the total surface area accessible to the solution, and it is not destructive [27]. In 258 259 fact, in this case, the real electrode areas are superior to the geometrical value; it could be 260 due to the character of diamond surfaces, allowing for proper contact of the solution with all active sites and consequently, a significant increase in the real area was achieved respect to 261 0.075 cm^2 . 262

For BDD/B:CNW electrode, an increase in the porous properties of the material was achieved due to the 3D nanostructures (see SEM images, Fig. 1) [20,21,24]; for this reason, an increase in the solution contact (solution diffusion between/into the porous) could be expected [14]. Meanwhile, this behavior is reduced in the BDD plate due to the homogeneous surface. In this context, the roughness factor ($RF = A_{real}/A_{geometric}$) was also determined for BDD and

BDD/B:CNW electrodes, considering the geometric surface and the estimated electroactive area for each one of the electrodes, achieving values of about 2.88 and 9.22, respectively. From these results, it is possible to deduce that, the new type of 3D-diamond electrodes with boron-doped carbon nanowalls (B:CNW) increased its roughness by about 3.2-folds in comparison to the BDD plate surface [14,15,21,34,35]. This result evidenced that, the roughness controls the contact of the active sites with the solution and thus, increasing its active area [14,21].

On the other hand, it is important to characterize the potential window and anodic 275 oxygen evolution reaction by electrochemical measurements at both carbon materials. Fig. 3 276 shows the polarization curves registered at BDD and BDD/B:CNW anodes in a sulfate 277 278 solution. A slight variation in the potential window is achieved, evidencing an important change in the potential of the oxygen evolution reaction (OER). This behavior is mainly 279 280 associated with the different carbon constituents on the BDD surfaces as well as the electrochemical decomposition of the supporting electrolyte [30,36,37]. Linear polarization 281 curves for BDD/B:CNW electrode showed that the potential for the OER was shifted to a less 282 positive potential (≈1.43 V vs Ag/AgCl). Based on previous works [36,38], it was 283 demonstrated that graphite electrode presents lower oxygen overpotential than BDD 284 electrodes, similar to the behavior achieved at BDD/B:CNW. Conversely, BDD plate showed 285 to be a poor electrocatalyst for OER, shifting this reaction to more positive potentials (Fig. 286 287 3).



288

Fig. 3. Polarization curves registered at BDD and BDD/B:CNW electrodes in 0.05 M of
Na₂SO₄. Inset: data in terms of Tafel plots at lower and higher overpotential regions from the
potential of hydroxyl radicals' formation.

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Meanwhile, additional effects could be observed when the analysis of the Tafel plots 293 294 is executed. E/log *j* plots at both BDD electrodes are shown in the inset of Fig. 3, considering 295 the real surface area for each one of them. As can be observed, the Tafel plots change their 296 slope according to the lower overpotential region (<1.6 V) or high overpotential region 297 (>1.62 V). At BDD/B:CNW, the OER seems to be the main process than that registered at 298 BDD plate. However, no unique slope can be attributed to the Tafel plots at both BDD 299 electrodes due to the complex set of the electrochemical reactions taking place involving most probably the OER and persulfate production. Tafel slopes about 0.6117 V decade⁻¹ and 300 0.5363 V decade⁻¹ for BDD plate and BDD/B:CNW electrode were estimated, respectively, 301 at lower overpotential region. Meanwhile, 0.3810 V decade⁻¹ and 0.3740 V decade⁻¹ in the 302

high overpotential region were obtained for BDD plate and BDD/B:CNW electrodes, 303 respectively. Two distinct linearity regions can be attributed to the electrochemical reactions 304 on the Nernst layer at BDD surface, associated with the sulfate ions in solution and water 305 discharge, which are transformed to persulfate and hydroxyl radicals ('OH) as well as oxygen, 306 respectively [39,40]. From Tafel plots data as well as the current and potentials (inset of Fig. 307 3), it is possible to infer that an important electrochemical reaction at the lower overpotential 308 region is attained at both diamond electrodes, which could be associated with the specific 309 310 interaction of sulfate ions with each one of the carbon surfaces [37,40,41].

Based on the existing literature [3,4,8], in the case of diamond electrodes, the 311 mechanisms followed depend strongly on (i) extension of the oxidation of SO_4^{2-} ions at the 312 313 diamond surface, via direct electron transfer, to produce persulfate (Eq. 2), (ii) the production of 'OH at anode surface in the reaction cage (Eq. 1), (iii) the interaction of 'OH with SO_4^{2-} 314 ions, in the Nernst layer, to favor the production of SO_4^{-} (principal intermediate) and 315 subsequently, persulfate (Eq. 3), as well as, in some cases, the association between the sp²-316 impurities on its surface and the adsorption of SO_4^{2-} ions on these active sites to also form 317 $SO_4^{-\bullet}$ and then, producing persulfate (Eqs. 4 and 5), 318

$$319 \quad 2SO_4^{2-} \to S_2O_8^{2-} + 2e^- \tag{2}$$

$$320 \quad SO_4^{2-} + {}^{\bullet}OH \to SO_4^{-\bullet} + H^+ + e^-$$
(3)

$$321 \quad BDD(SO_4^{2-}) \to BDD(SO_4^{-\bullet}) + e^-$$
(4)

$$322 \quad BDD(SO_4^{-\bullet}) + SO_4^{-\bullet} \to S_2O_8^{2-} \tag{5}$$

In the case of the BDD plate, the electrochemical reactions, which occurs at the lower overpotential region (<1.6 V), a more strong interaction is attained than that at BDD/B:CNW.

The Tafel slope is higher than the other one $(0.6117 \text{ V decade}^{-1} \text{ and } 0.5363 \text{ V decade}^{-1}$, for 325 plate BDD and BDD/B:CNW electrode, respectively). These behaviors appear to be 326 associated to the production of $S_2O_8^{2-}$ or SO_4^{--} at BDD surface [41], evidencing that, a direct 327 electron transfer is attained at BDD plate, in the former. While a fast-adsorptive interaction 328 of SO_4^{2-} ions on B:CNW-active sites (sp²-configuration) is achieved to form SO_4^{-} at 329 BDD/B:CNW electrode (Eq. 8) [36]. Afterward, at both electrodes, similar surface-layer-330 based approaches are accomplished at a higher overpotential region (>1.6 V), which are 331 332 related to the production of 'OH (Eq. 1) and O₂ (Eq. 6) as well as the formation of sulfatebased oxidizing species (such as SO_4^{-1} (Eq. 3) or $S_2O_8^{2-1}$ (Eq. 3 and 7)) by the participation 333 of free heterogeneous 'OH (Eq. 3). 334

$$335 \quad 2^{\bullet} OH \to O_2 + 2H^+ + 2e^- \tag{6}$$

336
$$SO_4^{-\bullet} + SO_4^{2-} \to S_2O_8^{2-} + e^-$$
 (7)

It is important to remark that, the surface-layer-based processes are achieved closer to the electrode surface, in the Nernst layer (from Armstrong to nanometers), which mainly depend on the diffusion distance of the free heterogeneous 'OH, the thickness of the diffusion layer, the convection (agitation and flow rate) and the diffusion coefficient of SO_4^{2-} ions in the solution [8]. However, the limiting steps to produce $S_2O_8^{2-}$ could be regarded as the diffusion of sulfate in the diffusion layer to react with the free heterogeneous 'OH (Eq. 3) as well as the subsequent reactions at BDD (SO_4^{-*})-sites on diamond surface (Eq. 5).

For this reason, it is necessary to control the current density to stabilize the $S_2O_8^{2-}$ electrogenerated, determining the extension of the production of $S_2O_8^{2-}$ at both electrodes.



347 Fig xx – Schematic representation of the persulfate reaction pathway on the 348 BDD/BCNW electrode

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3.3. Electrochemical production of persulfate

By applying different current densities (2.5, 5.0 and 15.0 mA cm⁻²) at 25 °C, the S₂O₈^{2–} concentration produced at both electrodes was determined by ISCO method (Fig. 4). Based on the existing literature, the maximum $S_2O_8^{2-}$ concentration is achieved between 120 and 150 min of electrolysis (inset in Fig. 4), then, the final persulfate concentration was plotted for all electrocatalytic materials at different applied current densities (Figure 4).



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Fig. 4. Electrochemically persulfate generated at different anode materials. Batch cell with
50 mL of 0.05 M Na₂SO₄ at 25°C.

359 As can be observed, an increase in the concentration of persulfate was attained when 360 the applied current density was increased (Fig. 4). For BDD plate, a significant enhancement in the production of persulfate was attained concerning those achieved at BDD/B:CNW 361 362 electrode. This behavior is related to the favored mechanism to electrogenerated persulfate [8]. According to the Tafel plot analysis, a stronger interaction is attained at a lower 363 overpotential region (<1.6 V) which indicated that the electrochemical production of $S_2O_8^{2-}$ 364 via direct electron transfer could be attained at BDD plate [3,8]. Meanwhile, at higher 365 366 potentials, the production of 'OH radicals favors an enhancement of the persulfate concentration [29]. Conversely, at BDD/B:CNW anode, the production of persulfate depends 367

on the formation of SO₄^{-•} at BDD surface (which seems to be the main intermediate), and the 368 subsequent surface-layer-based reactions of SO_4^{-1} [10,37,42,42]. In this sense, the results 369 related to the quantification of persulfate concentration indicated that the extension of the 370 persulfate mechanism depends on the nature of the electrode and the j (which is directly 371 associated with the cell potential). In fact, at lower i, the increase in the persulfate 372 concentration is less significant than those achieved when higher *j*. For example, at BDD 373 plate, from 2.5 to 5.0 mA cm⁻², the persulfate concentration increases by 1.33-folds, while 374 from 5.0 to 15.0 mA cm⁻², it increases by 1.64-folds. Similar behavior is observed at 375 376 BDD/B:CNW anode, where increases of about 1.21-folds and 1.34-folds were estimated, passing from 2.5 to 5.0 mA cm⁻², and from 5.0 to 15.0 mA cm⁻², respectively. However, there 377 are significant differences in the persulfate concentrations electrochemically generated. 378

Another feature that should be indicated is that a successive intensification on the *j* 379 could favor an increase in the production of 'OH radicals as well as an increase in the oxygen 380 evolution reaction [43–45]. In the former, these can react with the SO_4^{2-} ions in the diffusion 381 layer to form SO_4^{-1} (Eq. 3); and subsequently, produce $S_2O_8^{2-1}$ [42,46]. Meanwhile, when j 382 increases significantly, these can be also wasted favoring the production of O₂ (Eq. 6), in the 383 384 latter. These assertions are in agreement with the behavior reported by Pires et al. using BDD electrode [36], by applying *j* values higher than 15 mA cm⁻². On the contrary, at 385 BDD/B:CNW anode, more active BDD (SO4^{-•})-sites on the diamond surface can be formed 386 by increasing the *i* values (Eq. 3), favoring an increase in the production of persulfate via 387 Eqs. (4) and (5). 388

389 To understand the fundamentals of the mechanism related to the nature of 390 electrocatalytic material, the electrogeneration of persulfate at diamond electrodes was also compared to TiPt, Pb/PbO₂ and Ti/TiO₂-nanotubes/PbO₂. As can be observed at Fig. 4, it is evident that $S_2O_8^{2-}$ is efficiently produced at both diamond electrodes, by applying different *j*, with respect to the other anodic materials. This enhancement in the persulfate electrosynthesis is due to the mechanisms associated with the anodic surfaces used as well as the extension of each one of them.

Primarily, it is important to remark that no similar production of 'OH radicals, via water discharge, is attained at all electrocatalytic materials due to their active and non-active nature. Then, following this statement [1,35], higher production of 'OH radicals implies higher production of persulfate. For example, Ti/Pt produced lower concentrations of $S_2O_8^{2-}$ (0.36, 0.42 and 0.5 µM at 2.5, 5.0 and 15.0 mA cm⁻², respectively) because lower electrogeneration of 'OH radicals was reached at its surface, limiting the electrosynthesis mechanism via the participation of free heterogeneous 'OH Eqs. (3), (7) and (8).

403 Conversely, at the diamond electrode (e.g.: BDD plate), the efficient production of 404 'OH radicals is attained, favoring the generation of $S_2O_8^{2-}$ (such as 3.65, 4.87 and 7.98 μ M 405 at 2.5, 5.0 and 15.0 mA cm⁻², respectively) via the formation of $SO_4^{-\bullet}$.

$$SO_4^{-\bullet} + SO_4^{-\bullet} \to S_2O_8^{2-}$$
(8)

Secondly, the formation of $S_2O_8^{2-}$ via direct electron transfer and/or via (SO₄⁻⁺)-active sites on the electrode surfaces depends on the interaction of SO₄²⁻ ions and the electrocatalytic activity of each anode. In this frame, although the extension of the mechanisms is not completely clear, it seems that the electrogeneration of $S_2O_8^{2-}$, via direct oxidation of SO₄²⁻, is attained at Ti/Pt surface, as the main approach, confirming that a 412 complex set of electrochemical and chemical reactions can be attained at non-active anodes 413 and consequently, improving the electrosynthesis yield of $S_2O_8^{2-}$ concentration.

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3.4. Electrochemical oxidation of a target organic compound

To evaluate the performance of the persulfate electrochemically generated at diamond 416 surfaces and the other materials, oxidation in-situ experiments were carried out. Solutions of 417 25 mL of 0.05 M of Na₂SO₄ containing 10 mg L⁻¹ of MO were electrochemically treated by 418 applying 15 mA cm⁻², mixing with magnetic stirring during 40 min and aliquots were 419 withdrawn at predetermined times to determine the color removal, COD decay, and the 420 421 concentration of the target pollutant by an electroanalytic procedure. As can be observed in Fig. 5, EO process with BDD/B:CNW anode removed more than 95% of solution color in 5 422 min of electrolysis. Meanwhile, 93.8%, 95.7%, 98.6% and 85.8% were removed in 7, 10, 15 423 and 25 min at BDD plate, Ti/TiO2-nanotubes/PbO2, Pb/PbO2 and TiPt, respectively. 424 425 According to the spectrophotometric measurements, the intensity of the visible band at 465 426 nm continuously diminished until it disappears as a function of time during the galvanostatic electrolysis, leading to complete solution discoloration, in all cases. However, the absorbance 427 changes were reasonably rapid at BDD/B:CNW, BDD plate, Ti/TiO₂-nanotubes/PbO₂ and 428 Pb/PbO₂ anodes, indicating that the fragmentation of the MO chromophore group is rapidly 429 attained [29], promoting the elimination of color's solution and after that, the formation of 430 431 many other intermediates.



Fig. 5. Color removal effectiveness, as a function of time, at different anodic materials
electrogenerating *in-situ* sulfate-based oxidizing species by applying 15 mA cm⁻² at 25°C.
Inset: MO concentration decay, as a function of time, under similar experimental conditions.

In fact, when MO concentration was followed during the EO tests, it gradually 436 437 decreased and it was dependent on the nature of the material used. Again, non-active anodes were more efficient to remove MO from the solution (inset in Fig. 5), following an efficacy 438 order as BDD/B:CNW>BDD plate>Ti/TiO2-nanotubes/PbO2>Pb/PbO2>Ti/Pt. Only Ti/Pt 439 anode was less efficient, achieving 85% of removal from the MO initial concentration. 440 Analyzing the results obtained, BDD/B:CNW seems to be the most efficient anodic material 441 for the elimination of organic compounds from aqueous solution via SO₄²⁻/SO₄^{•-}/S₂O₈²⁻ 442 system in concomitance with free heterogeneous 'OH. However, the elimination of color and 443 MO is not an indicator of the effective removal of organic matter. Therefore, COD removal 444 was evaluated, at the end of the electrolysis time (40 min), by applying low and high *j* values 445

446 (2.5 and 15 mA cm⁻²). As observed in Fig. 6, different residual values of COD were
447 determined indicating that the organic matter was oxidized in all cases.



Fig. 6. Residual COD values, as a function of j (2.5 and 15.0 mA cm⁻²), after 40 min of
electrolysis at different anodic materials in EO tests at 25°C.

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On the one hand, residual COD concentrations in solution of about 15.1 and 9.0 mg 452 L⁻¹ were determined at 2.5 and 15 mA cm⁻² with BDD/B:CNW anode corresponding to 453 81.80% and 89.15% of COD removals, considering 83.0 mg L^{-1} as initial COD. On the other 454 hand, lower COD removals (4.76% and 25.51% at 2.5 and 15 mA cm⁻², respectively) were 455 achieved using Ti/Pt, as electrocatalytic material, that still represents a significant amount of 456 dissolved organic matter in solution. Based on these results, it is clear that the elimination of 457 organic matter occurred by the reactions attained in the reaction cage as surface-layer-based 458 approaches via $SO_4^{\bullet-}$, $S_2O_8^{2-}$ and free heterogeneous 'OH. But, the intensification of the 459

460 process is attained when a synergic effect is reached via the participation of all oxidant 461 species, which depends on the extension of the electrogeneration of each one of the oxidants, 462 mainly at non-active anodes.

The results obtained at diamond electrodes, such as the real surface area, Tafel plots, 463 464 persulfate production, and electrochemical degradation, evidenced that the concentration of SO_4 , $S_2O_8^{2-}$ and free heterogeneous 'OH could be different, under the experimental 465 conditions exposed here. This hypothesis is mainly related to the efficacy to produce 466 persulfate, discoloration level, and MO decay as well as COD removal comparing both BDD 467 plate and BDD/B:CNW anodes. In the former, all parameters were efficiently achieved and 468 superior to BDD/B:CNW anode. Meanwhile, COD was significantly removed from synthetic 469 470 effluent at BDD/B:CNW anode than the other electrodes. This behavior could be associated with the electrochemical generation of prior species that reacts in Nernst layer with the 471 organic compound before generating $S_2O_8^{2-}$. According to the literature [8,29], SO_4^{-} radicals 472 are the potential candidate, as already experimental [41] and theoretically [47] established; 473 in the case of diamond electrodes, the formation of BDD-(SO₄⁻⁻) sites is a pre-requisite step 474 to produce $S_2O_8^{2-}$ by recombining two SO_4^{--} species (Eq. (13)). However, as already proved 475 by electrochemical measurements, this behavior is strongly dependent on the carbon sp²-476 impurities content on the BDD surface. Graphitic sp^2 carbon act as the adsorption site for 477 sulfate-based species (e.g., SO_4^{2-} , HSO_4^{-}) at the diamond electrode, favoring the generation 478 of SO₄^{-•}, thus producing $S_2O_8^{2-}$. In the case of BDD/B:CNW electrode, its surface was 479 meaningly enlarged due to the 3D-diamond-carbon nanostructures. These nanostructures 480 (nano-flaps, nanowalls, as well as microcavities and micro-valleys) contain a key quantity of 481 482 hot-spot active sites, which could participate as adsorption sulfate-based species.

483 3D-diamond-carbon nanostructures + $SO_4^{2-}(aq) \rightarrow 3D$ -diamond-carbon nanostructures-(484 $SO_4^{2-})_{ads}$ (9)

485 3D-diamond-carbon nanostructures-
$$(SO_4^{-\bullet})_{ads} + SO_4^{-\bullet} \rightarrow S_2O_8^{2-}$$
 (10)

In this context, the formation of active-(SO₄-•) sites is mainly attained at BDD/B:CNW (Eq. 486 9), producing higher concentrations of SO_4^{-} at its surface when compared to the production 487 of free heterogeneous 'OH. Although similar oxidation-reduction potentials are registered 488 between SO₄^{-•} (2.5–3.1 V vs. NHE) and [•]OH (2.74 V vs. NHE), different extinction lifetimes 489 are attained in solution, 30–40 µs for SO4^{-•} and less than 1 µs for •OH [8], which allows to 490 substantially improve the SO₄^{-•} mass transfer performances and subsequently, its contact 491 chance and effectiveness with the pollutants in solution, as surface-layer- or as volume-492 solution- oxidation approaches [3,8,9,29,48]. Additionally, a wide pH window (from 2.0 to 493 8.0), due to the pH adjustment, can be used to favor the efficacy of the SO_4^{-} with the organic 494 pollutants. Consequently, BDD/B:CNW produces a higher concentration of SO₄^{-•} that, which 495 is available at active sites on its surface with a substantial lifetime, enhances the organic 496 497 matter removal (see Fig. 6) concerning other electrodes, mainly BDD plate.

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499 Conclusions

In summary, we can conclude that,

(i) BDD/B:CNW electrode is constituted by 3D-diamond-carbon nanostructures
which increase significantly its electroactive surface area, as confirmed by the
double layer capacitance determination obtaining an expansion of about 3.2folds respect to the area of a BDD plate.

505 (ii) The roughness factor indicated that the 3D-nanomaterial increased its porous
506 properties which were also confirmed by SEM images for BDD/B:CNW
507 surface.

- (iii) In both diamond materials, electrochemical measurements have indicated that 508 the complex set of the electrochemical reactions takes at lower and higher 509 over potential regions. However, it mainly seems that the persulfate 510 electrosynthesis depends on the stability/reactivity of 'OH and diamond 511 512 surface active sites to favor the production of SO₄⁻⁻. In the case of BDD plate, it produces rapidly SO_4^{-1} , via reaction between 'OH and SO_4^{2-1} , which form 513 $S_2O_8^{2-}$, but it is a less reactive oxidizing species. Meanwhile, SO_4^{--} species 514 are adsorbed at sp2-active sites at BDD/B:CNW favoring their participation 515 in the oxidation of MO. 516
- 517 (iv) In the case of PbO₂-based electrodes, indirect oxidation of SO_4^{2-} via 518 heterogeneous free 'OH can be hypnotized where a possible formation of the 519 active-(SO_4^{-*}) sites can be also feasible at α - PbO₂ or β - PbO₂ sites, which 520 should be still completely elucidated.

521 These conclusions open a new landscape associated with the real mechanism to 522 produce sulfate-based oxidizing species at BDD anodes as well as the electrochemical 523 generation of these oxidants at other non-active and active anodes.

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