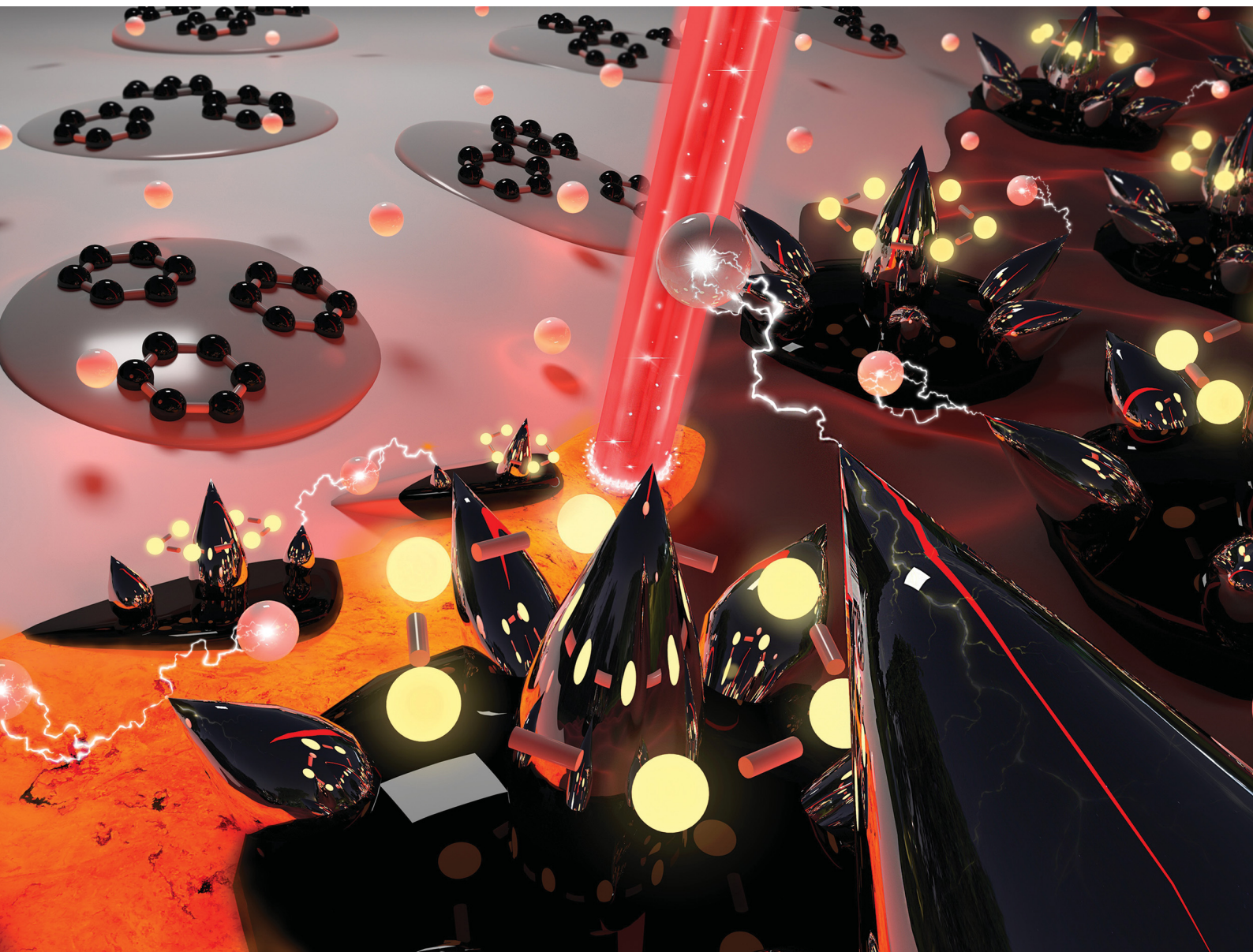


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**PAPER**

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Laser-induced highly oriented pyrolytic graphite  
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## Laser-induced highly oriented pyrolytic graphite for high-performance screen-printed electrodes†

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Screen-printed carbon electrodes (SPCEs) are enjoying increasing popularity in different electrochemistry areas, from electroanalysis to energy storage and power generation. Highly oriented pyrolytic graphite (HOPG), an ordered form of graphite, displays excellent electrochemical properties. However, its application in screen-printed electrodes has remained elusive. In this work, we present a straightforward laser-based process to selectively transform, in ambient conditions, the surface of conventional SPCEs into highly homogeneous HOPG. Energy densities between 6.8 and 7.7 mJ cm<sup>-2</sup> result in a binder-free, high-purity HOPG surface with very fast electron transfer rates. The electrode transformation to HOPG has been followed by SEM, Raman spectroscopy and XPS. Cyclic voltammetry of model systems ferrocyanide, ferrocenecarboxylic acid, dopamine and hydroquinone has been used to determine variations in electrode kinetics (from 50% increase for ferrocenecarboxylic acid, up to ca. 2 orders of magnitude for ferrocyanide and dopamine) and interfacial capacitance (from 40 up to 220 μF cm<sup>-2</sup>). Finally, differential pulse voltammetry (DPV) has been used to demonstrate the ability of these electrodes to detect dopamine in the presence of an excess amount of ascorbic acid.

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### Introduction

The mass fabrication of electrochemical sensors and biosensors, batteries and fuel cells has benefited enormously from screen-printing technologies.<sup>1–3</sup> Carbon-based materials, particularly graphite, have become dominant<sup>4</sup> due to their excellent balance between suitable electrochemical properties (chemical inertness, wide accessible potential window and low background currents, among others) and affordable cost. In spite of the wealth of existing carbon allotropes, screen-printed carbon electrodes (SPCE) are mainly based on graphite<sup>1</sup> and amorphous carbon.<sup>5</sup> Nanocarbon-based inks containing nanotubes or graphene are less common, but they are also available.<sup>6</sup> One important limitation of SPCE is that, due to their fabrication process, they display significantly lower electron transfer rates than their counterparts made of bulk carbon material.<sup>7</sup> This is evidenced by the large peak-to-peak separations displayed by the cyclic voltammetry of reversible redox

couples used in electrode characterization.<sup>7,8</sup> The reason is found in the heterogeneous composition of screen-printing inks and the electrode fabrication process. In addition to graphite, these screen-printing pastes contain a dielectric polymer and a solvent, and are thermally cured. Curing temperature is critical because it affects both the conductivity and the electrochemical activity of the printed structures.<sup>7</sup> Although high curing temperatures are preferred, only ceramic substrates withstand temperatures above 400 °C, able to eliminate both solvent and binder, leading to more conducting and electrochemically active electrodes. On the other hand, barring polyimides, polymeric substrates used in most flexible and printed electronics applications cannot cope with temperatures above 120–140 °C. As a result, electrodes printed on polymer or paper substrates seldom match their ceramic counterparts. Under these circumstances, SPCE are often modified through the addition of nanomaterials, *i.e.*: nanotubes,<sup>9</sup> nanoparticles,<sup>10</sup> or various graphene forms<sup>5</sup> to improve their electrochemical properties. While these modifications result in better sensors<sup>5</sup> and power sources,<sup>2</sup> the methods are often complex and difficult to up-scale.

Here, we describe a new selective method to transform the surface of graphite SPCE into a binder-free and homogeneous highly oriented pyrolytic graphite (HOPG) electrode using a CO<sub>2</sub> laser ( $\lambda = 10.6 \mu\text{m}$ ) in ambient conditions (in air). Laser has been used in the past to activate carbon electrodes,<sup>11–16</sup> including screen-printed ones,<sup>17,18</sup> and it has been reported to

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improve the crystallinity of polyacrylonitrile carbon fibers.<sup>19</sup> However, to the best of our knowledge, this is the first report on the laser-synthesis and characterization of HOPG from a graphite SPCE.

The application of laser treatments to carbon electrodes is not new. McCreery *et al.* studied the laser activation of conventional glassy carbon (GC) and highly ordered pyrolytic graphite (HOPG) using a Nd:YAG laser capable of producing 20 ns and 2 mJ pulses at 532 nm,<sup>20</sup> 10 ns, 300 J pulse at 532 and 1064 nm,<sup>21</sup> and even extremely high laser energies, in the MW cm<sup>-2</sup> range.<sup>11-13</sup> In contrast, Osborne *et al.*<sup>17</sup> applied 1500 pulses of 200 mJ using a 193 nm excimer laser on a carbon paste electrode and observed an improvement of its electrochemical response. The heterogeneous electron transfer rate constant for the ferro/ferricyanide couple increased from  $2.75 \times 10^{-6}$  cm s<sup>-1</sup> up to  $2.49 \times 10^{-3}$  cm s<sup>-1</sup>, as estimated from the corresponding peak-to-peak separation found in cyclic voltammetry experiments at 100 mV s<sup>-1</sup>. The increases in background capacitive currents reported in that work were explained on the grounds that the laser ablates the binder on the surface, leaving a larger carbon surface exposed to the solution. This work showed the changes induced by the laser on the electrode surface, including the appearance of “powdery deposits”, which they could not identify and attributed to charred binder debris. More recently, Hwang *et al.*<sup>18</sup> used a CO<sub>2</sub> laser, similar to the one used here, to create trenches on a screen-printed electrode, increasing surface area and producing capacitors with higher specific surfaces and energies. This work discussed the morphological changes induced at the micro-scale but not the laser effects on the material itself.

Thus, in the light of these and other recent reports of “laser-induced graphene”,<sup>22,23</sup> and laser-induced graphitization,<sup>19,24</sup> we set out to apply a CO<sub>2</sub> laser treatment to graphite screen-printed electrodes, aiming to improve their electrochemical response. Raman spectroscopy<sup>25,26</sup> ruled out the formation of laser induced graphene under the conditions reported here, but it strongly pointed to the formation of highly oriented pyrolytic graphite (HOPG), a very exciting electrode material.<sup>27</sup> Moreover, the process also removes the resin binder from the electrode surface, which area is increased and results in a better electrode–solution interface.

This finding is of great significance because hitherto the fabrication of low-cost, let alone screen-printed HOPG electrodes has not been possible. The conventional means to produce HOPG involves decomposition of gaseous hydrocarbons, often acetylene, at temperatures in the range 700–2400 °C, accompanied by a hot pressing process at high pressure.<sup>28</sup> Here we show that a mild CO<sub>2</sub>-laser irradiation can transform conventional graphite SPCEs into high-performance HOPG electrodes. This goes beyond the mere activation<sup>29-36</sup> of a carbon screen-printed electrode as it achieves a uniform surface of highly ordered graphite, with a high potential impact in all areas of Electrochemistry.

## Experimental

### Reagents and materials

All chemicals and materials were used as received without further purification. Potassium hexacyanoferrate(II) trihydrate,

potassium chloride, potassium nitrate, dopamine hydrochloride and L-ascorbic acid were purchased from Sigma-Aldrich (ES). RP-1 Agent (Mitsubishi Gas Company, JP) gas-barrier bags were used to preserve electrodes under dry and anoxic conditions.

Melinex ST506 PET substrates were sourced from SABIC Snij-Unie (The Netherlands). GST4500 (Sun Chemical-Servilan, ES) carbon paste for working and counter electrode, C21180423D2 (Sun Chemical-Servilan, ES) silver paste for reference electrode and EDAG PF 455B (LOCTITE) dielectric polymer were used for electrode material. Note that the graphite paste composition is not known at this stage, and Sun Chemical could not provide it due to confidentiality issues. Electrodes were designed using VectorWorks 2020 Student Edition (Techlimits, ES). Film-positives and screens were outsourced to Paymsar (ES). The used screens were fabricated using 90 threads per cm and 40 μm fiber diameter SEFAR 1500 fabric mounted on aluminum frames.

### Instrumentation

PET substrates and electrodes were cut and treated using a 30 W CO<sub>2</sub> Epilog Mini 18 laser engraver (Laser Project, ES). A Honle UV Technology UVACUBE 400 was used as irradiation chamber for dielectric film curing. Electrochemical measurements were performed using a SPELEC UV-vis spectroelectrochemistry instrument (Metrohm-Dropsens, ES) controlled by DropView SPELEC software (version 3.0), installed on a PC running Windows 10. The morphology of the working electrodes was assessed by scanning electron microscopy (FEG-SEM Hitachi S-4800) at 15 kV. Profilometry was performed on the working electrode for texture analysis. An Alpha-Step D100 mechanical profilometer with a strength of 1 mg was used to measure the step and the roughness of the obtained structures. Raman measurements were carried out using an InVia Raman of Renishaw equipped with a microscope Leica DMLM. A Modu-Laser brand argon ion laser with a wavelength of 514 nm was used. XPS measurements were performed with a SPECS system (Berlin, Germany) equipped with Phoibos 150 1D-DLD and monochromatic irradiation source Al K $\alpha$  (1486.7 eV). DSC and TGA were carried out in a nitrogen atmosphere using a Netzsch STA 449 F3 Jupiter analyser.

### Electrode fabrication

The fabrication of conventional 3-electrode screen-printed electrodes has been described elsewhere,<sup>35</sup> but a summary will be given here for convenience. First, silver tracks and pseudo-reference electrode were printed on a PET substrate. Secondly, working and auxiliary electrodes were printed, followed by a dielectric coating. Curing conditions were: 10 min at 100 °C in lab stove after silver and carbon paste printing. The dielectric film was cured by UV light for 45 seconds.

### Electrochemical measurements

In all cases, the open circuit potential (OCP) was determined prior to the start of any cyclic voltammetry or chronoamperometric experiments. The starting potential was then selected



within 50 mV of the OCP to ensure a near zero starting current. All measurements were carried out at room temperature. Solutions. 2 mM  $K_4Fe(CN)_6$  in 0.1 M  $KNO_3$ , 1.3 mM dopamine in 0.1 M  $H_2SO_4$ , 1.4 mM hydroquinone in 0.1 M acetate buffer pH 4.3 and 0.22 mM ferrocenecarboxylic acid in 0.2 M KCl.

## Results and discussion

### Transforming screen-printed graphite into HOPG

The electrodes discussed here were 2.5 mm diameter disk electrodes, screen-printed on polymeric substrates along with auxiliary and pseudo-reference electrodes in a conventional configuration to facilitate electrochemical experiments. Fig. 1 depicts the electrode fabrication process, including the laser raster treatment. Briefly, the electrodes were screen-printed in a three-layer process as described elsewhere.<sup>37</sup> Profilometric analysis of the carbon working and auxiliary electrodes reported here yielded a thickness of  $10 \pm 3 \mu\text{m}$ .

After printing, and before dicing, the electrodes were treated in air under ambient conditions using a 30 W Epilog Mini-18  $CO_2$  laser engraver. The laser system has several adjustable raster parameters, including image resolution, raster velocity, and laser power. The resolution was set at 600 dpi, and the raster velocity at approximately  $830 \text{ mm s}^{-1}$  (50% of the instrument maximum) in all experiments. This represents a pulse duration of approximately 51  $\mu\text{s}$ . Changing resolution and raster speed was ruled out to make the study more manageable.

The effect of increasing laser energies on the electrochemical properties of the electrodes was evaluated for a nominal laser power range between 1.8 W and 3.6 W. Considering the electrode size ( $r = 1.25 \text{ mm}$ ), it is estimated that energies of roughly  $5.1$  to  $10.2 \text{ mJ cm}^{-2}$  were applied. The minimum laser energy causing a visible effect on the electrochemical properties being  $5.1 \text{ mJ cm}^{-2}$ . Above  $10.2 \text{ mJ cm}^{-2}$ , on the other hand, all the printed carbon coating was ablated and there was no electrode material left. In the following, we will refer to the energy density applied to the electrodes normalized to

electrode area. Thus, the figures may be considered independent of laser raster velocity or resolution and they should facilitate comparison with other reports.<sup>38</sup> Table S1 (ESI<sup>†</sup>) provides the correspondence between laser settings and the reported energy densities.

The electrode material transformation has been evaluated as a function of applied energy, using a combination of microscopy and spectroscopy techniques including SEM, Raman and XPS. Profilometry, contact angle and four-probe sheet resistance measurements were also performed followed by a comprehensive electrochemical study.

### Surface characterization

**Changes in appearance.** The first observed variation on the electrodes during fabrication following laser treatment was their appearance. While electrode surface color is of little importance in most applications, it can have a major impact on spectroelectrochemical measurements, as it strongly influences material–light interaction, including absorption and/or scattering. ImageJ<sup>39</sup> analysis of the lasered electrodes showed a 20% increase in color intensity with applied power between the ready-printed electrode and the electrode exposed to  $7.7 \text{ mJ cm}^{-2}$ . Beyond this point, the electrode color intensity decreased again as larger proportion of the electrode was ablated and the printed structure became translucent (see Fig. S1, ESI<sup>†</sup>). This darker appearance of the lasered electrodes demonstrated surface transformation.

**Changes in surface morphology.** Fig. 2 shows representative SEM images of a ready-printed surface, and those treated with laser energies up to  $7.7 \text{ mJ cm}^{-2}$ . Note that SEM images of electrodes treated with more than  $7.7 \text{ mJ cm}^{-2}$  looked nearly identical up to  $10.2 \text{ mJ cm}^{-2}$ . The central images show the surface at the boundary of the lasered region. Laser raster tracks can be observed, particularly at lower laser energies, allowing us to determine the laser spot size in those conditions. Spot size increases with applied energy from *ca.*  $75 \mu\text{m}$  at  $5.1 \text{ mJ cm}^{-2}$ , up to over  $95 \mu\text{m}$  at  $7.7 \text{ mJ cm}^{-2}$ . At higher applied energies, adjacent raster tracks are too overlapped to measure the spot size. The images on both sides of the figure show the ready-printed surface (left) and representative images of the surface following laser treatment at  $7.7 \text{ mJ cm}^{-2}$ . The ready-printed electrode surface seems to consist of small graphite portions protruding through the binder, whereas the lasered surface shows no signs of binder, and a very homogeneous coverage of porous graphite. The graphite formations remind of the “powdery deposits” reported in the past.<sup>17</sup> As it will be shown next, these formations were probably HOPG crystallites.

**Laser effect on electrical properties.** Two sets of different measurements were carried out to evaluate the electrical conductivity of the lasered screen-printed structures. First, silver contacts were added on opposite sides of the electrodes to measure their electrical resistance. The results, provided in Fig. S2 (ESI<sup>†</sup>), show that the square resistance increased from roughly  $138 \pm 5 \Omega$  for the ready-printed electrode, up to roughly  $600 \pm 10 \Omega$  for electrodes receiving a treatment of  $7.7 \text{ mJ cm}^{-2}$ . Above this, electrical resistance soared to the k $\Omega$  range as the

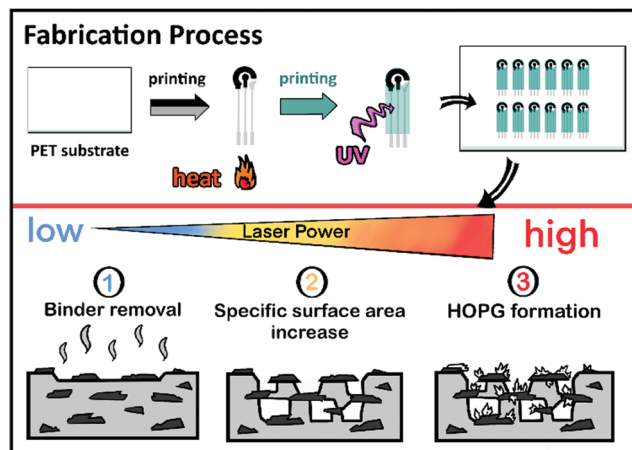


Fig. 1 Schematic representation of the HOPG-SPCE fabrication process.



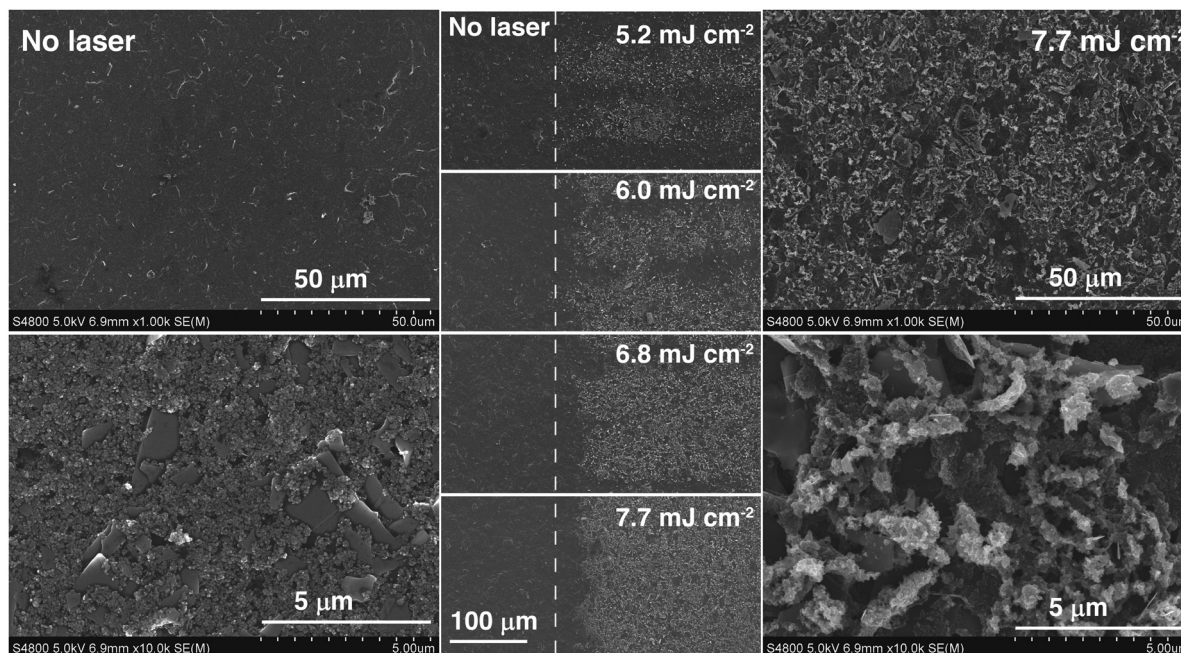


Fig. 2 SEM images of different screen-printed electrode surfaces treated with different laser powers. The dotted line crossing the central images separates the ready-printed area (left) from the laser-treated area (right).

electrodes were thinned below  $5\ \mu\text{m}$ . However, the sudden and abrupt change was not attributed only to a thinning of the printed layer, but to the appearance of discontinuities in the layer. These discontinuities were attributed to the faster ablation rates of the binder, a cellulosic material, and smaller graphite particles when compared to larger graphite particles in the paste, leading to the formation of gaps and voids and the corresponding decrease of the overall electrical conductivity of the defective layer.

In addition to this, *ca.*  $30\ \mu\text{m}$ -thick carbon paste layers were prepared by doctor blade on PET substrates. Following the same curing treatment as the electrodes,  $1 \times 1\ \text{cm}^2$  samples were prepared. The square resistance was measured using a four point probe technique. No significant changes were observed in the electrical conductivity of the printed structures, which suggests that the transformation induced by the laser on the material may have been only superficial. Indeed, the electrical resistivity of the bulk material seemed unchanged after laser treatment. Four-point probe measurements yielded very similar electrical resistivity for untreated and SPCEs exposed to different laser powers. It was concluded that film electrical resistance increased as a result of the laser ablating too deep into the original coating. As shown in Fig. S3 (ESI<sup>†</sup>), the ablated material thickness was directly proportional to laser power, at a rate between  $1.9$  and  $2.2\ \mu\text{m}\ \text{W}^{-1}$ .

**Raman demonstrates the transformation from graphite to HOPG.** Fig. 3 shows the Raman spectra of the ready-printed coating and those of electrodes treated with  $\text{CO}_2$  laser up to  $10.2\ \text{mJ}\ \text{cm}^{-2}$  energies, as well as spectra from basal-plane HOPG and graphene. Raman spectroscopy is commonly used to provide a structural fingerprint that facilitates the identification of carbon allotropes.<sup>40</sup> As Fig. 3 illustrates, the Raman

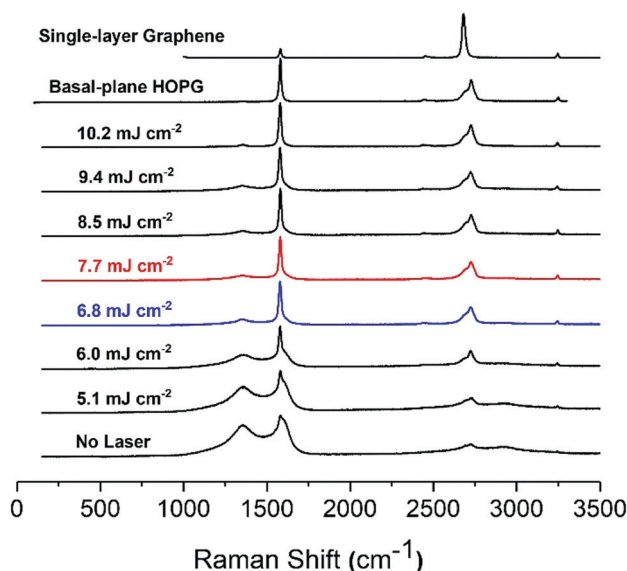


Fig. 3 Raman spectra of ready-printed (bottom) and lasered SPCE. The figure also shows the spectra of basal-plane HOPG and of single-layer graphene (top) for comparison. The best conditions are highlighted in blue ( $6.8\ \text{mJ}\ \text{cm}^{-2}$ ) and red ( $7.7\ \text{mJ}\ \text{cm}^{-2}$ ).

spectra of the laser-treated electrodes is nearly identical to that of HOPG, and very different from that of graphene.

The Raman spectra of the carbon paste show the presence of the characteristic D ( $1360\ \text{cm}^{-1}$ ), G ( $1600\ \text{cm}^{-1}$ ) and 2D or G' ( $2730\ \text{cm}^{-1}$ ) bands, typical of graphitic materials.<sup>26</sup> The D band is originated by the distortion of the  $\text{sp}^2$  crystal structure.<sup>41,42</sup> It is a defect-induced Raman feature, so its removal upon laser



irradiation suggests that a more ordered material has emerged. As previously shown in Fig. 2, laser treatment removes binder and brings about crystallite formation along the surface with no preferential path. The higher the laser energy applied, the larger the resulting crystallite structures. These crystals can be attributed to purification and arrangement of graphite layers by elimination of amorphous graphite and distortions in the  $sp^2$  graphitic structure. In fact, it has been reported that the intensity decrease in  $1360\text{ cm}^{-1}$  band of the Raman spectrum is correlated with increasing graphite micro crystallite size and with graphitic edge plane density.<sup>43</sup> This decrease of the  $1360\text{ cm}^{-1}$  band concurs with the formation of microcrystals, as shown in SEM images of the lasered SPCE.

The G band is the graphitic band. It arises from the C–C  $sp^2$  bond and is common in  $sp^2$  carbon systems.<sup>5</sup> It is the sharpest band in our series of spectra. In turn, the 2D ( $G'$ ) band consists of two components for the bulk graphite and one sharp for graphene.<sup>44</sup> The intensity of G and 2D bands, including the presence of two peaks at 2D evidence the presence of multi-layered graphite in the carbon inks. The untreated SPCE paste shows a high intensity for D band, indicative of distortions in the graphitic crystal structure. A wide 2D band reveals the presence of amorphous carbon. As the laser energy applied increases from  $5.1$  to  $6.8\text{ mJ cm}^{-2}$ , the D band decreases while the intensity of the G band remains stable, and the 2D becomes more defined, suggesting a higher quality graphite. Between  $6.8$  and  $9.4\text{ mJ cm}^{-2}$ , the D band is stable. The  $I_D/I_G$  ratio informs of the structure disorder, and lower ratios indicate less disordered structures.<sup>5,45</sup> For an energy density of  $10.2\text{ mJ cm}^{-2}$  the D band is at a minimum, and  $I_D/I_G$  is around  $0.07$ . Also, the 2D band is well defined with its two components showing the presence of high-quality graphite.

Highly ordered pyrolytic graphite, HOPG, displays D and G bands at  $ca. 1600\text{ cm}^{-1}$  and  $2650\text{ cm}^{-1}$ , respectively. HOPG is turbostratic, meaning that the graphitic planes have their  $c$ -axes (perpendicular to the main plane) parallel aligned in the crystal but are rotationally disordered relative to adjacent crystals.<sup>4</sup> Conventional HOPG is usually composed of large crystals. In contrast, the SEM images clearly show the polycrystalline nature of our printed electrodes, which nevertheless present Raman spectra consistent with basal plane HOPG.<sup>25</sup> The inter-crystallite size parameter  $L_a$  can be estimated from Raman spectra through eqn (1):<sup>44</sup>

$$L_a(\text{nm}) = (2.4 \times 10^{-10}) \lambda_{\text{laser}}^4 \left( \frac{I_D}{I_G} \right)^{-1} \quad (1)$$

where  $\lambda_{\text{laser}}$  corresponds to the Raman laser wavelength ( $514\text{ nm}$ ) used, and  $I_D$  and  $I_G$  represent the D and G peak intensity, respectively. In the present case, laser treatment resulted in an increase of  $L_a$  from  $ca. 19\text{ nm}$  for the ready-printed electrode up to  $ca. 115\text{ nm}$  after application of  $7.7\text{ mJ cm}^{-2}$ .  $L_a$  continued to rise with applied power as the band at  $1360\text{ cm}^{-1}$  decreased. At  $10.2\text{ mJ cm}^{-2}$ , an  $L_a$  of  $243\text{ nm}$  was found. As shown in Fig. 3, increasing laser power gradually improved the crystallinity of the resulting electrode surface. At energy densities of  $7.7\text{ mJ cm}^{-2}$  and above, the material

Raman spectrum was almost identical to that of HOPG,<sup>28</sup> and clearly different from that of graphene. Discriminating between edge-plane pyrolytic graphite (EPPG) and basal-plane pyrolytic graphite (BPPG) can be difficult. Katagiri<sup>25</sup> showed how the Raman D and G band ratio differs between BPPG and EPPG, with EPPG displaying larger ratios due to the relatively higher structural disorder introduced by the edges. While Raman spectroscopy of our materials may suggest a significant formation of basal plane, the small size of the graphite particles in the original paste results in a material where edge sites are abundant. However, the  $I_D/I_G$  ratio of  $0.07$  obtained after application of  $10.2\text{ mJ cm}^{-2}$  leads to  $L_a$  values near  $243\text{ nm}$ , consistent with BPPG. Table S1 (ESI†) presents the  $L_a$  value as a function of applied laser energy.

**XPS surface analysis confirms an increase in  $sp^2$  carbon.** The surface composition of several treated electrodes was investigated by XPS.<sup>46</sup> Fig. 4 shows the XPS spectra in the energy region where carbon moieties appear, between  $280$  and  $300\text{ eV}$ . The figure displays the spectrum of the ready-printed material, and that of the material following a laser treatment at  $7.7\text{ mJ cm}^{-2}$ .

While the increase in  $sp^2$  carbon found by XPS is supported by Raman spectroscopy, the significant  $sp^3$  carbon peak is harder to explain from the SEM images and the Raman spectra. The relative concentration of surface  $sp^2$  carbon increases from  $ca. 15\%$  to above  $26\%$  after irradiation with laser energies  $7.7\text{ mJ cm}^{-2}$  and higher, in line with Raman spectra showing a more orderly graphite structure as laser energy increases. At the same time, the  $sp^3$  content drops from about  $40\%$  down to about  $35\%$ , which is still relatively high and seemingly in contradiction with Raman spectra. The increase in graphite crystallinity may be explained on the grounds that because laser is applied in air, the high local temperatures, combined with oxygen lead to the consumption of defective graphite, which leaves behind the more thermally stable graphite. Temperatures achieved by the laser could not be accurately measured, but thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) of a carbon paste sample (Fig. S4, ESI†) revealed some interesting clues. After solvent evaporated around  $200\text{ }^\circ\text{C}$ , the binder decomposed between  $250\text{ }^\circ\text{C}$  and  $400\text{ }^\circ\text{C}$ . Next, graphite was decomposed in at least three distinct steps. It is likely that amorphous graphite will decompose at lower temperatures, and more crystalline material could withstand harsher conditions. In light of this, it may be reasonable to argue that the laser can achieve temperatures above  $400\text{ }^\circ\text{C}$  in all cases.

Regarding the  $sp^3$  signal, on the other hand, the apparent disagreement between Raman and XPS may be due to the presence of binder polymer and ablation residues that may not give a significant Raman signal, while being clearly visible by XPS.

**Water-in-air contact angle changes.** As shown above, laser treatment brings about significant changes both in surface morphology and chemical structure. As a result, treated SPCEs became increasingly hydrophilic with applied laser energy. The contact angle for water in air at the ready-printed electrode was  $71^\circ \pm 2^\circ$ , laser energies further resulted in decreasing contact angles down to  $47^\circ \pm 5^\circ$ , as shown in Fig. S5 (ESI†). This is a



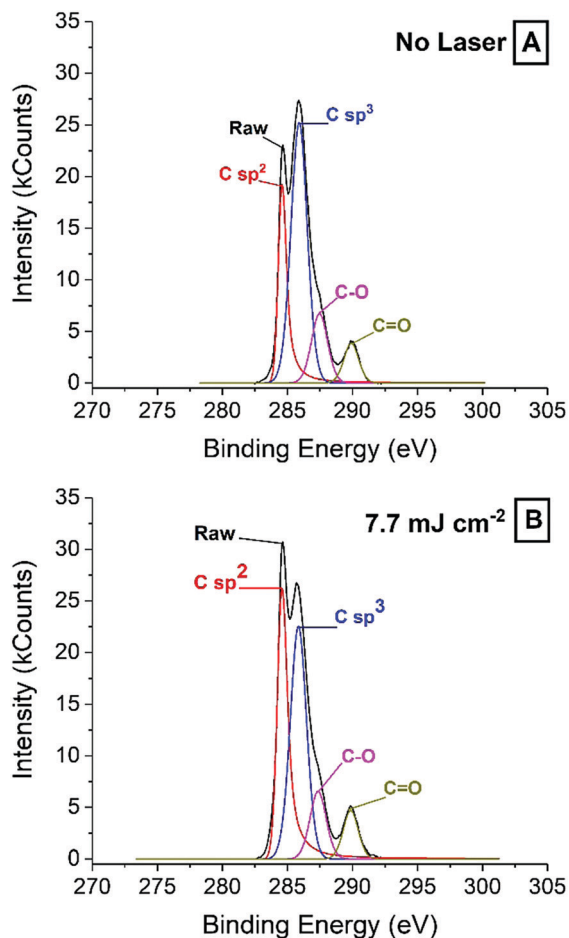


Fig. 4 XPS spectra of a ready-printed electrode surface (A) and an electrode surface treated with  $7.7 \text{ mJ cm}^{-2}$  laser energy (B).

surprising result because we expected that the increase in  $L_a$  value resulting from laser irradiation would lead to a more hydrophobic surface. In this case, although surface chemistry effects cannot be entirely ruled out, we believe that the effect is predominantly due to the electrode porosity caused by surface binder ablation. The hydrophilicity of the laser-treated sample is related to the surface roughness and its chemical state. Wettability of hydrophobic surfaces is qualitatively explained in terms of two extreme models, namely Wenzel's model for wetted states and the Cassie–Baxter model for unwetted states.<sup>47</sup> According to Wenzel's model, a hydrophobic surface with adequate roughness can lead to a fully wetted situation. In the present case, laser action results in improved wetting mainly through the enhancement of surface roughness, which is overall beneficial to electrochemical performance in aqueous solutions. At the same time, XPS data presented in Fig. 4B shows the presence of C–O and C=O bonds on the lasered surface, which may facilitate water access into the pores.

### Electrochemical characterization

Carbon material structure determines electrochemical properties.<sup>4,28</sup> Edge plane sites provide SPCEs with remarkable traits, as edge sites

are considered to be more reactive than basal planes.<sup>27,48</sup> Fig. 5 shows cyclic voltammograms of  $2.2 \text{ mM}$  ferrocyanide in  $0.1 \text{ M KNO}_3$  at  $100 \text{ mV s}^{-1}$  for electrodes treated with different laser energies. The ready-printed electrode (Fig. 5A) displays peak-to-peak separation in excess of  $400 \text{ mV}$ . While this is a very large value, it is common for screen-printed electrodes cured at low temperatures.<sup>7</sup> It is important to note how the oxidation and reduction peaks become better defined, with higher currents and narrower peak-to-peak separations, as the laser energy increases from  $5.1 \text{ mJ cm}^{-2}$  up to  $7.7 \text{ mJ cm}^{-2}$  (Fig. 5B–E). Application of higher energies has a deleterious effect on electrode performance, as shown in Fig. 5F and G. The explanation is simple: higher laser energies result in a deeper layer ablation, which increases the film resistance and precludes an effective electrode polarization.

Fig. 5H shows the Randles–Ševčík representation of the oxidation peak current *vs.* square root of the scan rate. The plot shows the expected diffusion control for the current from the Randles–Ševčík equation substituting for the experimental conditions and electrode geometry described in the Experimental section, as well as the current from a ready-printed electrode and that of the same electrode after exposure to  $7.7 \text{ mJ cm}^{-2}$  laser energy. The latter dataset deviates much less from the theoretically expected behaviour than the ready-printed electrode. This is due to a more homogeneous electrode surface displaying faster electron transfer rates.

Last, Fig. 5I shows the increase in interfacial capacitance as a function of laser energy. This capacitance has been determined from charging currents measured from cyclic voltammetry at scan rates below  $200 \text{ mV s}^{-1}$  according to the method described before.<sup>49</sup> In short, the potential was scanned between  $-0.2$  and  $1 \text{ V vs. Ag/AgCl}$ , in  $0.1 \text{ M H}_2\text{SO}_4$ . The current at  $0.5 \text{ V vs. Ag}$  was plotted against the scan rate and the capacitance was calculated from the slope of the linear region. Note that the capacitance measured at other potentials may differ, and that we are only providing this value to show the changes that follow laser treatment. The differential capacitance measured this way increased in direct proportion to the laser energy applied, from  $38 \mu\text{F cm}^{-2}$  at the ready-printed electrode,  $45 \mu\text{F cm}^{-2}$  at  $5.1 \text{ mJ cm}^{-2}$ , and up to  $220 \mu\text{F cm}^{-2}$  following treatment with  $8.5 \text{ mJ cm}^{-2}$ . This capacitance value is in line with previous reports by McCreery on the laser activation of glassy carbon electrodes.<sup>18</sup>

Ferrocyanide is typically used as model system due to its reversibility and wide availability. However, it is also sensitive to electrode surface quality.<sup>50</sup> Moreover, we next explored the response of the laser-treated electrodes towards other benchmark redox pairs: dopamine, hydroquinone and ferrocenecarboxylic acid. Fig. 6 shows typical cyclic voltammograms ( $10 \text{ mV s}^{-1}$ ) obtained for these pairs at ready-printed and at  $7.7 \text{ mJ cm}^{-2}$ , which we consider close to the optimum laser treatment. Direct  $\text{CO}_2$  laser activation resulted in the greatest electron transfer enhancements for ferrocyanide (Fig. 6A) and dopamine (Fig. 6B), which became noticeably more reversible according to the lower peak-to-peak separation, lower onset potentials, and higher currents. Dopamine undergoes inner-sphere electron transfer and is affected by surface oxides, so



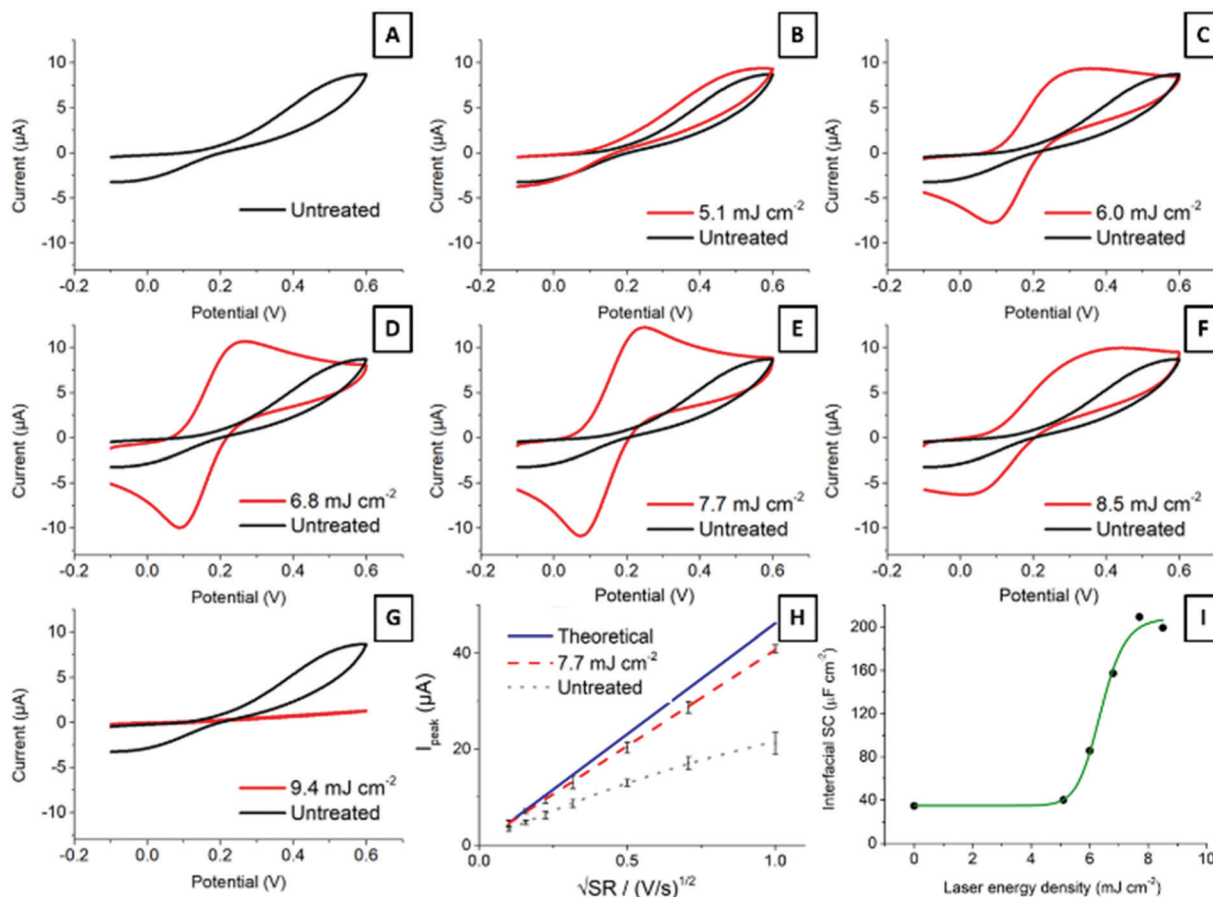


Fig. 5 Electrochemical response evolution with increasing laser energies (A–G), current peaks comparison between theoretical, ready-printed SPCE and 7.7 mJ cm<sup>-2</sup> activated SPCE (H) and variation in interfacial specific capacitance as a function of laser energy applied (I). The data correspond to different electrodes produced in the same batch, and treated with different laser powers.

certain interaction with the electrode surface is expected.<sup>51</sup> While ferrocyanide, on the other hand, is not rigorously “outer-sphere”, it is much more reversible than dopamine and reportedly surface oxide independent.<sup>4</sup>

Two laser effects can be responsible for the enhancement in electron transfer. First, phys- and chemisorbed particles and impurities may be removed by laser ablation. Second, active sites may be generated throughout HOPG microcrystalline planes. In the case of HOPG, the electron transfer rates for ferrocyanide and dopamine systems increase substantially when microstructural edge plane defects are formed by laser treatment. The high temperatures brought about by the laser action likely not only consume the binder, but also any amorphous or highly defective graphite particles too, leaving only the most crystalline and hence temperature resistant graphite. The edges of these highly crystalline regions are most likely responsible for the fast electron transfer rates observed.<sup>15</sup>

However, hydroquinone (Fig. 6C) and ferrocenecarboxylic acid (Fig. 6D) did not show such significant improvement in electron transfer. Both hydroquinone and ferrocenecarboxylic acid are considered outer-sphere redox systems, relatively insensitive to electrode surface composition.

The apparent heterogeneous electron transfer rate constant,  $k_s$ , for these systems, particularly dopamine and hydroquinone,

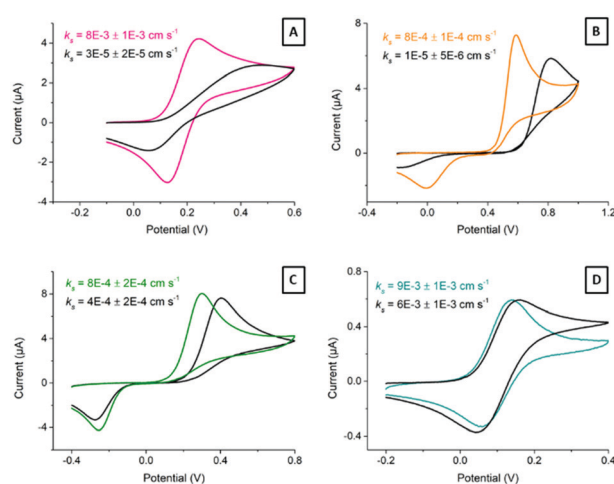


Fig. 6 Comparison between an ordinary SPCE (black lines) and laser activated (6.8 mJ cm<sup>-2</sup>) SPCE (coloured lines). Cyclic voltammograms at 10 mV s<sup>-1</sup> of 2 mM K<sub>4</sub>Fe(CN)<sub>6</sub> in 0.1 M KNO<sub>3</sub> (A), 1.3 mM dopamine in 0.1 M H<sub>2</sub>SO<sub>4</sub> (B), 1.4 mM hydroquinone in acetate buffer 0.1 M pH 4.3 (C) and background subtracted 0.22 mM ferrocenecarboxylic acid in 0.2 M KCl (D).

was estimated through simulation using KISSA software<sup>52,53</sup> over the scan rate range between 10 mV s<sup>-1</sup> and 1 V s<sup>-1</sup>. Note



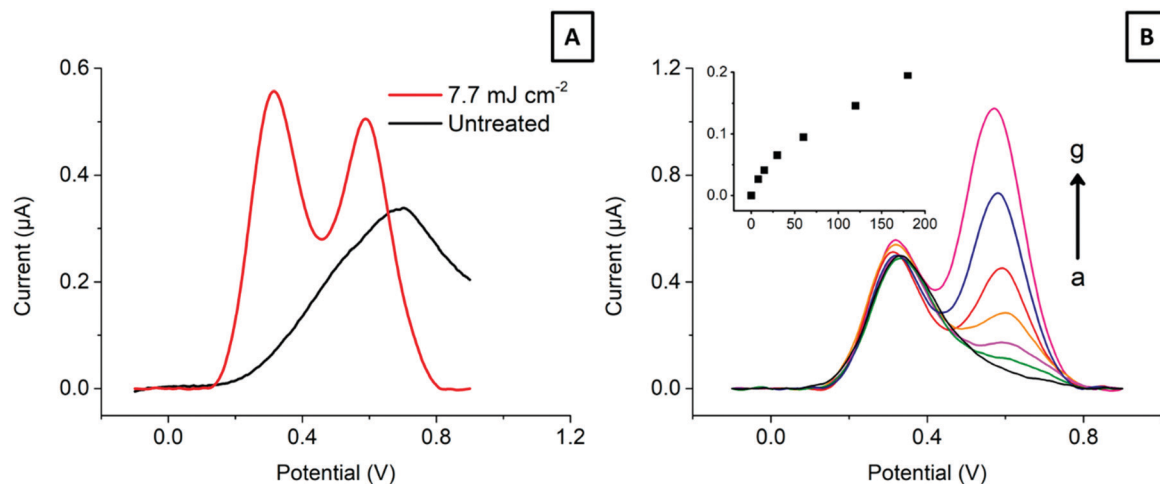


Fig. 7 (A) DPV of 150 μM ascorbic acid (AA) and 60 μM dopamine (DA) in pH 4 0.1 M PBS at a laser activated (7.7 mJ cm<sup>-2</sup>) SPCE (red) and ordinary SPCE (black). (B) From a to g: 0, 8, 15, 30, 60, 120, 180 μM dopamine in the presence of 150 μM AA. DPV parameters:  $E_{\text{pulse}} = 50$  mV;  $t_{\text{pulse}} = 50$  ms;  $E_{\text{step}} = 4$  mV; scan rate = 10 mV s<sup>-1</sup>. The inset shows the peak charge (μC) as a function of dopamine concentration (μM).

that Nicholson's<sup>54</sup> and related methods are valid only for systems displaying  $\Delta E_p \leq 200$  mV. On the other hand, Matsuda and Ayabe's method,<sup>55</sup> can be applied to systems displaying wider voltammetric peak separations, but the issue here is that the voltammograms display too large  $\Delta E_p$  even for very slow scan-rates, making it very difficult to choose the scan-rate meeting the condition  $\Lambda = 1$ . Here,  $\Lambda = k_s / \sqrt{(nFD\nu)/(RT)}$ , where  $n$  is the number of electrons,  $F$  is Faraday constant,  $D$  is the diffusion coefficient of the electroactive species,  $\nu$  is the scan rate,  $R$  is the gas constant and  $T$  the absolute temperature. In the cases of dopamine and hydroquinone, where protons are also involved in their respective mechanisms, only simulation provides a reliable way to assess the electron transfer steps.<sup>56</sup>

Regarding the time-stability of the laser treatment, we studied the electrode behaviour over time (data not shown but available on reasonable request). Activity losses around 50% were observed through ferrocyanide voltammetry a week after laser ablation. However, these could be almost completely prevented by storing the electrodes in an inert atmosphere, which suggests that reaction with ambient oxygen is likely to be detrimental, in line with observations by other workers.<sup>28</sup>

Electroanalysis of dopamine in the presence of ascorbic acid. Finally, to show the benefits of the laser treatment in an electroanalytical application, we have addressed the detection of dopamine, a neurotransmitter, in the presence of a large excess of ascorbic acid, a common interfering agent. Fig. 7 shows typical differential pulse voltammograms (DPV) of 0–180 μM dopamine in the presence of 150 μM ascorbic acid at an electrode lasered with 7.7 mJ cm<sup>-2</sup>.

The DPV parameters consider the electrode interfacial capacitance (see Fig. 5 above) and the current response. Fig. 7A clearly shows the ability of the treated HOPG-SPCE to resolve both signals, compared to the conventional SPCE. Moreover, as Fig. 7B displays, the two signals are resolved over a wide dopamine concentration range, making it possible to detect dopamine in the low μM range, with ascorbic acid present in a

near 100-fold excess. This is attributed to the ability of HOPG to discriminate between electroactive species with close redox potentials based on their different surface reactivity, in line with other reports.<sup>9,57,58</sup>

## Conclusions

We have shown that the application of relatively mild laser energies on graphite SPCEs, using a CO<sub>2</sub> laser in air, can bring three important advantages, namely, (i) the removal of surface binder, (ii) an increase in surface porosity and (iii) the purification of amorphous graphite and its transformation into HOPG. The former two benefits have been demonstrated by SEM images and contact angle measurements, while Raman spectroscopy supported the latter one. Surface binder removal is of great importance in the fabrication of sensors, energy storage and generation devices, and electrode interfaces, as it prevents contamination and ensures a biocompatible and chemically homogeneous interface. The increase in surface area can be more useful in power generation devices, such as supercapacitors, but it may also be exploited in electroanalysis. Last, the HOPG produced by laser irradiation of graphite microparticles in the printed electrodes display very high electrochemical activity. This can equally benefit sensors, actuators, and energy storage devices. In all cases studied, both inner- and outer-sphere systems, we have observed higher electron transfer rates after laser irradiation of the SPCE. The higher rates of electron transfer are evident from the voltammetric peak-to-peak separations over a range of scan rates. These benefits have been demonstrated through the detection of dopamine, a neurotransmitter, in the presence of ascorbic acid.

Raman and XPS analysis have shown that the described laser treatment results in a relative increase of sp<sup>2</sup> carbon, indicating a more ordered material. Indeed, crystallite size has been calculated according to Pimenta *et al.*<sup>44</sup> We have found that laser treatment increases crystallite size lattice parameter



$L_a$  from 20 nm for the untreated electrodes up to 240 nm for the laser-treated case ( $10.2 \text{ mJ cm}^{-2}$ ). The possibility to tune the crystallite size and thus obtain graphite closer to either EPPG or BPPG reproducibly is one of the greatest advantages of the reported process.

Another significant advantage is the process selectivity, which also makes it extremely energy efficient. Because the transformation only occurs in the irradiated areas and the  $\text{CO}_2$  laser spot size is in the order of  $100 \mu\text{m}$ , it is possible to treat only the desired electrode regions, leaving the rest of the surface untouched. Patterning of electrodes with areas of different reactivity can lead to the simultaneous detection of several analytes on a single electrode or the production of arrays of different electrodes in a single process. Laser processing is much more efficient in terms of time, energy and resources than other reported SPCE activation methods,<sup>30,32,33,36,59</sup> leading to comparable if not superior results.

In summary, the process and the new electrode material reported here are of great significance. The resulting surfaces are free from binder impurities, the exposed graphite is of high purity and crystallinity, and the surface area increases compared to readily printed structures. The process allows some flexibility, as it is possible to tune the graphite form between EPPG and BPPG. The resulting electrodes display excellent electrochemical properties and can generally benefit electrochemical applications.

## Author contributions

Alejandro Fidel Alba: investigation, formal analysis, visualization, writing-original draft. Joseba Totoricaguena-Gorriño: formal analysis, visualization, writing-original draft. Lia Campos-Arias: investigation. Nikola Peřinka: investigation, writing-review and editing. Leire Ruiz-Rubio: writing-review and editing. José Luis Vilas-Vilela: writing-review and editing, funding acquisition. Senentxu Lanceros-Méndez: writing-review and editing, funding acquisition. Francisco Javier del Campo: conceptualization, methodology, supervision, writing-original draft.

## Conflicts of interest

There are no conflicts to declare.

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