

# Journal of Materials Chemistry A

Materials for energy and sustainability

Accepted Manuscript

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# Elucidating Interfacial Parameters of Platinum–Palladium Bulk Alloy Single Crystals

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## ABSTRACT

The application of mixed catalysts in chemical to electrical energy transformation reactions can be a strategy for enhancing the catalysis of these reactions. Platinum and palladium exhibit significant catalytic activity in reactions of different natures, influenced not only by composition but also by structure. The present work presents a fundamental study of the characterization and electrochemical behavior of platinum-palladium bulk alloy single-crystal electrodes (Pt<sub>100-x</sub>Pd<sub>x</sub>), prepared via a modified Clavilier protocol. Electrodes with varying mass ratios of Pt and Pd, compositions ranging from 0 to 15% Pd, and crystallographic orientations of (111) and (100) were manufactured. X-ray photoelectron spectroscopy (XPS) and low-energy electron diffraction (LEED) reveal a well-correlated bulk and surface composition with ordered crystalline structures. Cyclic voltammetry in HClO<sub>4</sub> and H<sub>2</sub>SO<sub>4</sub> electrolytes indicates stronger anion interactions compared to pure platinum. Charge displacement experiments by CO adsorption estimate the effect of surface orientation and Pd content on the potential zero total charge (pztc). For Pt<sub>100-x</sub>Pd<sub>x</sub>(111) surfaces, the pztc is very close to that of Pt(111), while surfaces with (100) orientation show decreasing pztc values with higher Pd content, favoring OH adsorption at lower potentials. Finally, the CO oxidation reaction on these Pt-Pd bulk alloy electrodes is studied by cyclic voltammetry, providing a profoundly comparative discussion of this reaction with Pt(111), Pt(100), and Pt(111) modified with Pd sub-monolayers.

**Keywords:** Platinum–Palladium alloys; Single crystal electrodes; XPS; LEED; pztc; adsorption



## 32 1. Introduction

View Article Online  
DOI: 10.1039/D4TA01771D

33 The search for new electrode materials with better electrocatalytic activity and selectivity  
34 has led to the study of alloys and multicomponent systems, since in most cases, the activity of  
35 pure metals is not sufficient for practical applications. The electrochemical properties of a  
36 material are not only influenced by its composition but also by its surface structure. In this context,  
37 the use of catalysts with a well-defined surface, such as single-crystal electrodes, allows for the  
38 exploration of electrocatalytic reactions at the atomic level. Since the surface structure is well-  
39 defined, they facilitate the untangling of complex reaction mechanisms and set the basis for the  
40 development of advanced electrocatalysts.

41 Metallic catalysts of mixed composition may present better catalytic activity. The  
42 bifunctional <sup>1</sup>, third body <sup>2</sup>, and electronic effects <sup>3</sup> are the main phenomena that result in the  
43 synergistic interaction of different metals that make up an improved catalyst. These effects are  
44 widely explored in catalysts applied in transforming chemical energy into electrical energy in  
45 direct liquid fuel cells (DLFCs), powered by the oxidation of organic molecules at the anode and  
46 the reduction of oxygen at the cathode. Thermodynamically, organic molecules present attractive  
47 conditions for their use in DLFCs <sup>4</sup>. However, the oxidation of these molecules occurs through  
48 more than one reaction pathway <sup>5,6</sup>, leading to the formation of poisoning intermediates, and  
49 partially oxidized products. All these problems reduce the energetic efficiency of the DLFCs. CO  
50 and organic acids are one of the main undesirable by-products when it comes to the oxidation of  
51 organic molecules into platinum. Due to the strong interaction of these molecules with the surface,  
52 high overpotential values are necessary to oxidize them to CO<sub>2</sub> and thus restore the activity of the  
53 catalyst and maintain a continuous reaction.

54 Platinum and palladium possess similar physicochemical properties and display good  
55 catalytic activity for the reactions used in a DLFC. Although platinum is considered the  
56 electrocatalyst per excellence, in some cases, palladium performs better than platinum in the  
57 oxidation of organic molecules. A typical example of this is the formic acid oxidation reaction <sup>7,8</sup>.  
58 Contrary to what is observed for platinum, this reaction is not limited by surface poisoning by CO



59 on palladium. Voltammetry studies combined with density functional theory (DFT) show that the  
60 oxidation of formic acid in a Pd monolayer deposited on Pt(111) occurs due to the formation of a  
61 large formate coating, hindering the formation of CO on the surface<sup>9</sup>. However, one of the main  
62 problems with pure palladium is the absorption of hydrogen at low potentials, which may alter  
63 the properties of the electrode<sup>10,11</sup>.

64 The use of palladium monolayers on platinum basal planes as templates has been a way of  
65 studying the electrochemical behavior of Pd<sup>12–18</sup>. Under these conditions, hydrogen absorption is  
66 not observed, allowing studies to be carried out in a lower potential region than those used in pure  
67 palladium electrodes. However, in these studies, the growth of 3D islands easily occurs without  
68 the complete formation of monolayers, compromising the good structural definition of the  
69 surface. An alternative approach to studying the Pt-Pd bimetallic system consists of preparing  
70 alloys with well-ordered surface structures. In this sense, Pt<sub>100-x</sub>Pd<sub>x</sub> alloy single crystals can be  
71 used for fundamental electrochemical studies without the problem of hydrogen absorption and  
72 preserving their well-defined surface. In the case of Pt-Pd alloy single crystal surfaces, the amount  
73 of published work is small and many open questions about their interfacial properties have not  
74 been elucidated. The first reports of the use of a Pt<sub>100-x</sub>Pd<sub>x</sub> alloy single-crystal surface in  
75 electrochemical studies were presented by Attard et al.<sup>19–21</sup>. Through the combination of different  
76 spectroscopy and voltammetry techniques, it was shown that regardless of the mixed composition  
77 of the single-crystal electrodes, the alloyed surfaces are well-defined, and electrochemical studies  
78 can be conducted at the atomic level on these surfaces.

79 The specific adsorption of species plays a crucial role in surface electrochemistry, exerting  
80 a significant influence on reaction kinetics by either promoting or inhibiting the process. Charge  
81 displacement experiments allow the determination of the potential of zero total charge (pztc) of  
82 platinum group metals and enable the discrimination of the different adsorption processes taking  
83 place at the surface of the electrode<sup>22</sup> according to the sign of the associated total charge.

84 In the present work, a fundamental study of the characterization and electrochemical  
85 behavior of platinum–palladium bulk alloy single crystals is presented. Previous works in  
86 electrocatalysis have shown that the electrocatalytic behavior of the material depends not only on



87 the surface structure and composition but also on the interfacial structure. Thus, surface charge<sup>24</sup>  
88 adsorbed species<sup>24,25</sup>, pH, and water structure modify<sup>26</sup> the activity of the electrode, and in some  
89 cases catalyze or deactivate reactions. A full characterization of the interfacial properties of these  
90 electrodes is required when their electrocatalytic behavior is to be understood. For this reason,  
91 Pt<sub>100-x</sub>Pd<sub>x</sub> alloy single crystal electrodes with (111) and (100) orientation of bulk compositions  
92 ranging from 0 to 15 % Pd were prepared and characterized by a combination of low-energy  
93 electron diffraction (LEED), X-ray photoelectron spectroscopy (XPS), and electrochemical  
94 techniques. The interaction of ion perchlorate and sulfate with Pt<sub>100-x</sub>Pd<sub>x</sub> surface, as well as the  
95 adsorption of hydrogen and (hydro)oxygenated species, were studied under a voltammetric  
96 regime using reaction media based on perchloric or sulfuric acid. Finally, the potential of zero  
97 total charge and the electrocatalytic oxidation of CO were investigated by considering the effect  
98 of the surface orientation and composition.

## 99 2. Experimental section

### 100 2.1 Electrode preparation.

101 Platinum single-crystal electrodes were manufactured following Clavilier's protocol<sup>27</sup> and a brief  
102 modification of this procedure was introduced to prepare the Pt-Pd alloy single-crystal electrodes  
103<sup>19,21,28</sup>. Briefly, the end of an ultrapure (Mateck, 99.99 %) Pt wire (0.5 mm diameter) is melted in  
104 a propane-oxygen flame to produce a single crystal bead of approximately 2-3 mm in diameter.  
105 By controlling the length of the melted Pt wire it is possible to control the amount of Pt in the  
106 single crystal bead. Then, a known amount of Pd wire (Good Fellow, 99.99 %) is added to the  
107 melted Pt bead, and both elements are allowed to blend for several minutes in the molten solution.  
108 After careful and slow cooling of the bulk alloy, a single crystal bead with (100) and (111)  
109 microfacets is formed. The single crystal is then placed on a goniometer head at one end of an  
110 optical bench, where it is oriented, cut, and polished following the procedure described elsewhere  
111<sup>19</sup>. As the final step, the crystal is annealed in a Bunsen flame for 1 h to eliminate the surface  
112 damage from the polishing procedure. Pt-Pd alloy single crystal electrodes with (111) and (100)  
113 orientations of nominal bulk compositions ranging from 0 to 15 % Pd were prepared following



114 this methodology. The Pt composition and the geometrical area of the electrodes measured using  
115 XPS spectra and an optical microscope, respectively, are given in Table 1.

116 Before each electrochemical experiment, the single-crystal electrodes were flame  
117 annealed in a Bunsen flame (propane-air) and cooled in a reductive atmosphere of an Ar and H<sub>2</sub>  
118 mixture (N-50, Air Liquide) with a 3:1 ratio. The electrode surface is then protected by a drop of  
119 H<sub>2</sub>/Ar-saturated high-purity water (Millipore, Milli-Q system, 18.2 MΩ cm) and transferred to  
120 the electrochemical cell.

121  
122 Table 1. Surface composition extracted from Pt 4f and Pd 3d XPS spectra, and electrode geometric  
123 area of the different electrodes used in this study.

<b>Pt(111) and Pt<sub>100-x</sub>Pd<sub>x</sub>(111) Surfaces</b>				
Geometric Area / cm <sup>2</sup>	0.0371	0.0468	0.0374	0.0505
%Pt (XPS)	100	97	96	88
<b>Pt(100) and Pt<sub>100-x</sub>Pd<sub>x</sub>(100) Surfaces</b>				
Geometric Area / cm <sup>2</sup>	0.0458	0.0469	0.0412	0.0472
%Pt (XPS)	100	99	93	85

124

## 125 2.2 Ultra-high vacuum surface characterization.

126 X-ray photoelectron spectroscopy (XPS) measurements were carried out using a sample-to-  
127 analyzer angle of 90°. The system employed was a commercial hemispherical analyzer (SPECS  
128 GmbH, Phoibos 150 equipped with a MCD-9 Detector) using a pass Energy of 15 eV and a  
129 monochromatic XRM X-ray source (SPECS GmbH) utilizing an Al anode (E = 1486.7 eV). A



130 source power (P) of 300W was used.

131 Low energy electron diffraction (LEED) measurements were conducted using a  
132 commercial system (SPECS GmbH, ErLEED 150). After the optimization of the measurement  
133 conditions, electron energies of 132.5 and 80-95 eV were used for Pt(111) and Pt(100),  
134 respectively. Sputtering was carried out using Ar-ions ( $E = 2$  keV) for 7 minutes at a chamber  
135 pressure of  $3 \times 10^{-5}$  mbar. Consecutive sample annealing was carried out using an electron source  
136 emission current (I) of 18 mA which equates to a sample temperature of roughly 850 °C in the  
137 employed system. Each sample was sputter-annealed three times.

138 Figure S1 shows XPS C1s spectra of the Pt(100) electrode with different pretreatments.  
139 A significant amount of carbon species is present at the surface when various distinct pre-  
140 treatments are applied to the electrode, which highlights the high affinity of Pt for C species under  
141 ambient conditions. Upon flame-annealing in air and quick transfer to the UHV chamber, this  
142 amount is significantly reduced, but adventitious carbon is still found, which is expected even for  
143 small exposures to the ambient atmosphere. Only when the single crystals are cleaned in UHV  
144 using cycles of Ar ion sputtering and consecutive annealing, a fully carbon-free surface is  
145 obtained. LEED patterns were only attainable on the UHV-cleaned surfaces, which accounted for  
146 the carbon contaminations with the other preparation methods in UHV.

### 147 **2.3 Electrochemical Characterization.**

148 The electrochemical experiments were conducted at room temperature in a custom-made glass  
149 cell with a three-electrode configuration. The counter electrode (CE) consists of a platinum wire  
150 whose area is much larger than the working electrode (WE). The electrode potential was measured  
151 against a reversible hydrogen electrode (RHE) connected to the cell through a Lugging capillary.  
152 All potentials shown in this study are given on the RHE scale. The single crystal electrodes were  
153 characterized by cyclic voltammetry in meniscus configuration in argon-purged 0.1 M HClO<sub>4</sub>  
154 (Sigma-Aldrich ACS reagent – 70-77 %) and 0.1 M H<sub>2</sub>SO<sub>4</sub> (Merck KGeA Suprapur – 96%)  
155 electrolytes prepared with ultrapure water (ElgaPurelab Ultra Analytic system, 18.2 MΩ cm). The  
156 voltammetric profiles of Pt(111) and Pt(100) were recorded at the beginning of each working



157 session to ensure the cleanliness of the electrochemical system <sup>21,29,30</sup>.

158 CO oxidation experiments were carried out to determine the properties of the electrodes for  
159 this process. CO adsorption on the electrode surface was performed under a potentiostatic regime,  
160 where the electrode was polarized at 0.1 V, and a CO flow was bubbled through the electrolyte  
161 until complete blockage of the surface (monitored by cycling the electrode between 0.05 and  
162 0.3 V). Throughout this procedure, the electrode current was monitored to ensure that the  
163 electrode surface was fully saturated with adsorbed CO, according to the procedure described in  
164 reference <sup>31</sup>. Before the oxidation of the adsorbed CO, the CO present in the supporting electrolyte  
165 and the cell atmosphere were removed by argon flow for 15 minutes, maintaining the electrode  
166 potential in the potentiostatic regime. Then, two cyclic voltammograms at 0.02 V·s<sup>-1</sup> were  
167 recorded to register the electrochemical oxidation of adsorbed CO in the first one and the recovery  
168 of the initial profile in the second one.

169 Surface charge displacement experiments were performed by adsorbing CO on the surface  
170 in a potentiostatic regime, following the protocol previously used by our group <sup>32</sup>. Before CO  
171 adsorption on the surface, a voltammetric profile of the electrode surface was recorded to ensure  
172 the cleanliness of the supporting electrolyte and the condition of the electrode surface. The  
173 electrode was polarized in potential where the oxidation of adsorbed CO does not occur (this  
174 information was obtained using the results of oxidation of adsorbed CO, previously described),  
175 being: 0.1, 0.2, or 0.3 V, and started measuring the current over time. A stream of CO was injected  
176 into the cell atmosphere until the system current remained at zero for a few seconds. With the  
177 integration of the current/time curve, the nature of the species adsorbed on the electrode was  
178 estimated at each studied potential, as well as its charge. All CO adsorption, as well as adsorbed  
179 CO oxidation experiments, were performed at least three times, and the results are presented in  
180 terms of their mean values with their associated error.

181

### 182 3. Results and discussion

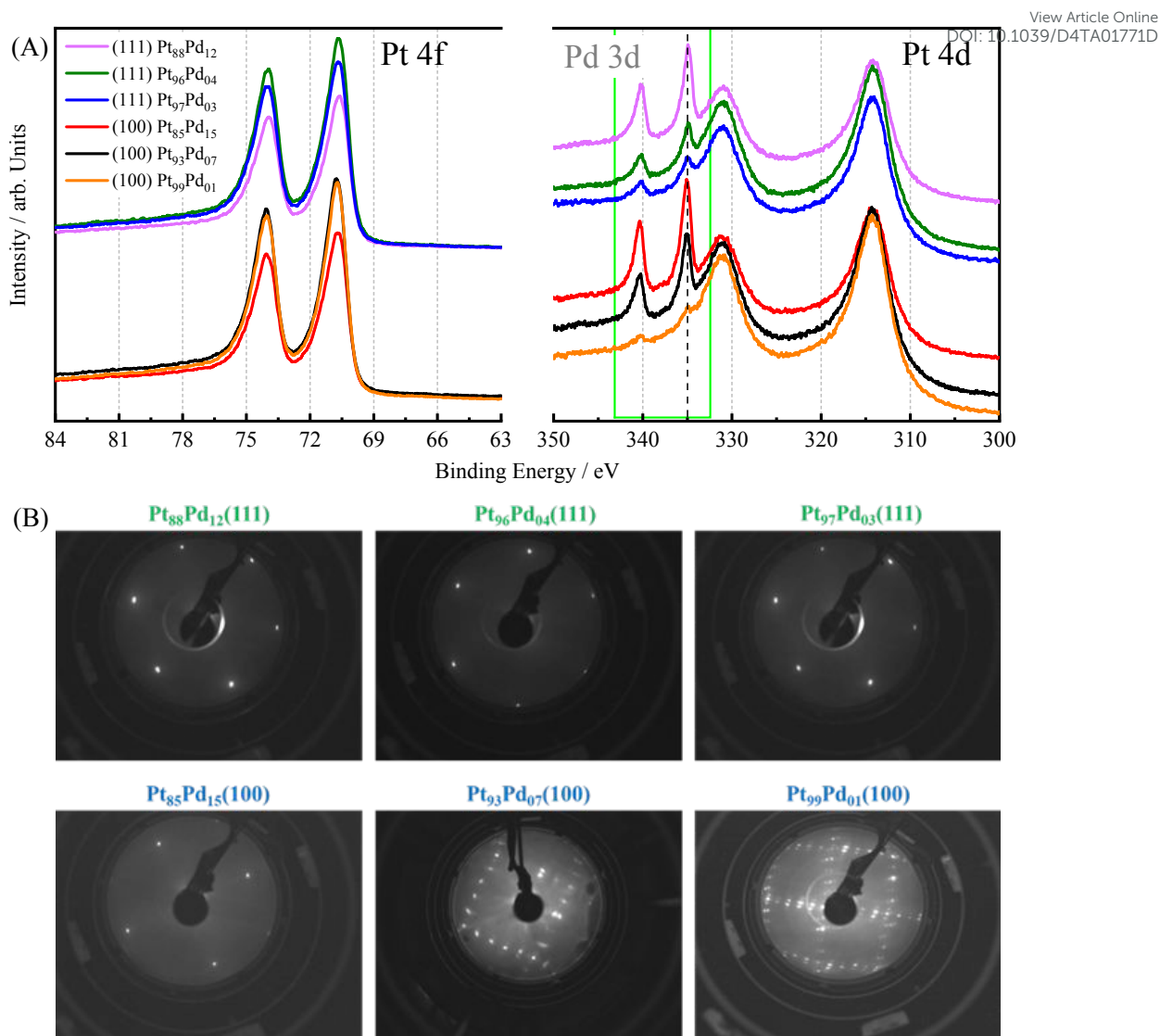
#### 183 3.1. UHV characterization of the Pt<sub>100-x</sub>Pd<sub>x</sub> single-crystal electrodes





184 To validate the preparation of the  $\text{Pt}_{100-x}\text{Pd}_x$  alloy single crystals, the surface composition  
185 and structure of the electrodes were assessed using XPS and LEED, respectively. Although  
186 complete miscibility of Pt and Pd has been reported elsewhere<sup>33</sup> and no surface segregation was  
187 calculated for this alloy<sup>34</sup>, earlier works showed that Pd can segregate and that this separation is  
188 structure-dependent<sup>30</sup>. However, our XPS data reveal a good correlation between bulk and  
189 surface compositions. To evaluate the surface composition, Pt 4f and Pd 3d XPS spectra were  
190 acquired on the UHV-prepared single-crystal surfaces, Figure 1(A). Both metals were found to  
191 be in their metallic state ( $E_{\text{B,Pt}} = 70.76$  eV and  $E_{\text{B,Pd}} = 335.04$  eV)<sup>35</sup>, irrespective of the amount of  
192 Pd in the alloy and the surface orientation. The measured surface composition, determined by the  
193 relative integrated peak area ratio of the Pt 4d to Pd 3d core level region, fits the expected relative  
194 nominal values and is summarized in Table 1. It should be noted that although XPS is a surface-  
195 sensitive technique, the signal acquired typically probes up to 5-10 nm below the surface.  
196 Nonetheless, the contribution of the different layers to the spectra diminishes with the distance to  
197 the surface. Although enrichment of Pd has been proposed for the topmost layer, the values  
198 obtained here are clearly an averaged value of all the sampled layers and therefore, do not only  
199 reflect the very last surface layer.





200

201 Figure 1 (A) Pt 4f and Pd 3d XPS spectra for  $\text{Pt}_{100-x}\text{Pd}_x(111)$ , and  $\text{Pt}_{100-x}\text{Pd}_x(100)$  single crystal  
 202 surfaces. (B) LEED patterns acquired on  $\text{Pt}_{100-x}\text{Pd}_x(111)$ , and  $\text{Pt}_{100-x}\text{Pd}_x(100)$  single crystalline  
 203 surfaces after UHV sputter-annealing treatment. The electron energies were 132.5 and 80-95 eV  
 204 (111) and (100) structures, respectively.

205

206 Figure 1(B) shows the corresponding LEED patterns of the  $\text{Pt}_{100-x}\text{Pd}_x(111)$ , and  $\text{Pt}_{100-x}\text{Pd}_x(100)$   
 207  $\text{Pd}_x(100)$  surfaces. The low background and sharpness of the diffraction spots point out the well-  
 208 ordered surface structure of all samples. All of the (111) crystals yield a  $(1\times 1)$  structure with no  
 209 indication of any other ordered overlayer phase present, independently of the amount of Pd in the  
 210 electrode <sup>36</sup>. For the (100) surfaces containing a low amount of Pd, a Pt-hex-R0.7° structure is

9



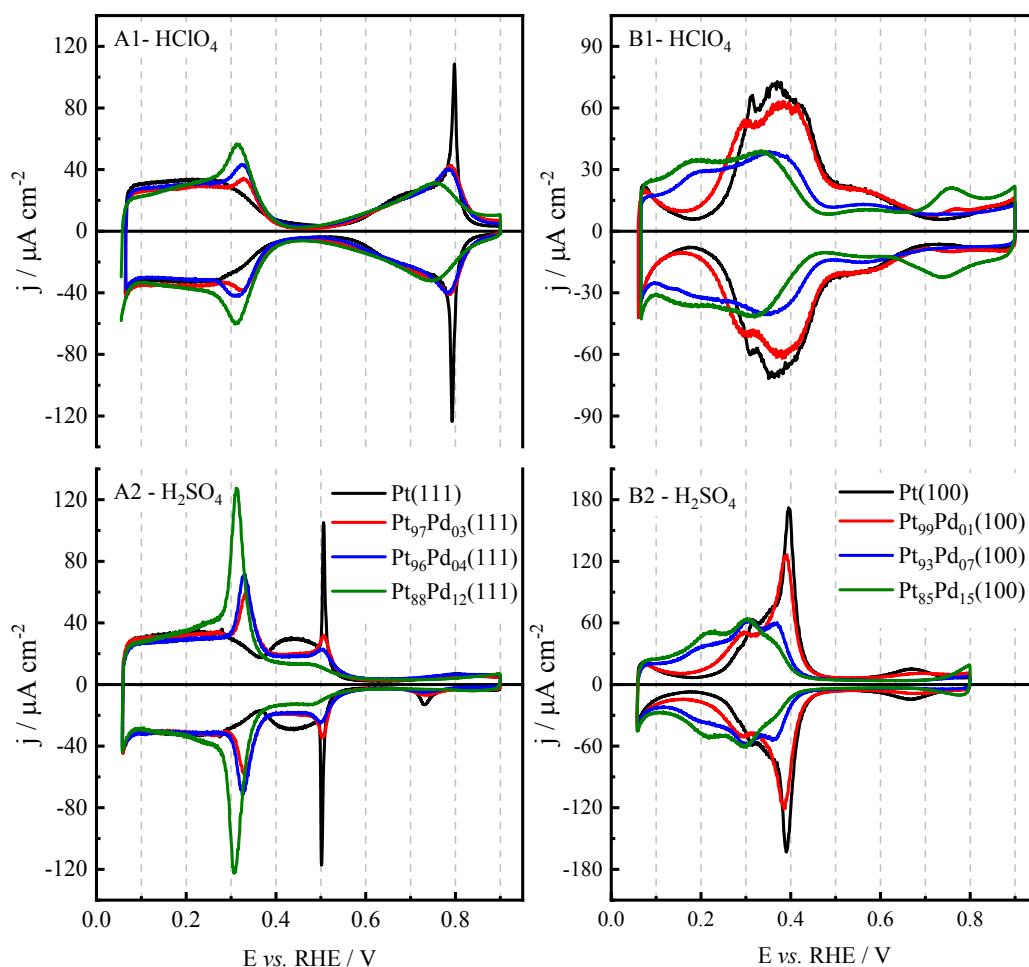
211 observed, which is consistent with previous studies on pure Pt(100) and Pt(100) with small  
212 amounts of foreign adatoms<sup>30,37,38</sup>. When increasing the Pd content up to 15 %, as obtained from  
213 the integrated XPS areas (Table 1), the LEED shows a (1×1) structure pattern. It is known that  
214 Pd(100) does not reconstruct<sup>39</sup>, which leads to the suppression of the hexagonal (hex)  
215 reconstruction once the Pd content is sufficiently high<sup>30</sup>.

216

### 217 **3.2. Voltammetric characterization of the Pt<sub>100-x</sub>Pd<sub>x</sub> single crystal surfaces.**

218 Cyclic voltammograms of a Pt-based surface on a given electrolyte provide important  
219 correlations between surface structure and composition since the voltammetric profiles serve as  
220 surface fingerprints. Also, information on the adsorption processes that take place on the electrode  
221 surface can be achieved. This information is very relevant when electrocatalytic processes are  
222 studied since adsorbed molecules can activate or hinder the electrochemical reactions<sup>24,40,41</sup>.  
223 Figure 2(A) shows the voltammetric profiles of Pt(111), and Pt<sub>100-x</sub>Pd<sub>x</sub>(111) surfaces, in both  
224 0.1 mol L<sup>-1</sup> H<sub>2</sub>SO<sub>4</sub> and 0.1 mol L<sup>-1</sup> HClO<sub>4</sub>. The potential range was set in both electrolytes  
225 between 0.065 and 0.900 V for all surfaces except for those (100)-oriented and measured in  
226 sulfuric acid medium. For these electrodes the profile was recorded up to 0.80 V, avoiding the  
227 formation of the soluble palladium sulfate salt<sup>42-45</sup>. Unlike pure Pd electrodes, Pt<sub>100-x</sub>Pd<sub>x</sub> surfaces  
228 do not absorb hydrogen, which allows reaching potentials lower than 0.2 V<sup>30</sup>.





229

230 Figure 2 Voltammetric profiles of (A1-A2) Pt(111) and Pt<sub>100-x</sub>Pd<sub>x</sub>(111) surfaces in (A1) 0.1 mol  
 231 L<sup>-1</sup> HClO<sub>4</sub> and (A2) 0.1 mol L<sup>-1</sup> H<sub>2</sub>SO<sub>4</sub>, registered at 0.05 V s<sup>-1</sup>. Voltammetric profiles of (B1-B2)  
 232 Pt(100) and Pt<sub>100-x</sub>Pd<sub>x</sub>(100) surfaces in (B1) 0.1 mol L<sup>-1</sup> HClO<sub>4</sub> and (B2) 0.1 mol L<sup>-1</sup> H<sub>2</sub>SO<sub>4</sub>,  
 233 registered at 0.05 V s<sup>-1</sup>.

234

235 Pt<sub>100-x</sub>Pd<sub>x</sub>(111) surfaces in perchloric acid share common features with Pt(111) electrode  
 236 (Figure 2 (A1)). Thus, there is a clear distinction between hydrogen and anion adsorption regions.  
 237 However, the addition of Pd brings about some changes to the voltammetric profile. Currents in  
 238 the hydrogen region nicely overlap below 0.2 V for all the electrodes, although a slightly smaller  
 239 current is observed for the surfaces with a higher content of Pd. This behavior is also reported for  
 240 studies carried out in different acid, neutral, and alkaline electrolytes<sup>46-48</sup> because this potential

11



241 region is a consequence of the predominant reversible reaction of hydrogen adsorption. When Pd  
242 is added, a pair of reversible peaks appears in the potential region between 0.2 and 0.4 V, and  
243 their peak potential moves from 0.335 V (red curve) to 0.320 V (green curve) with increasing Pd  
244 loading. Also, the charge under the peak increases. On the other hand, some modifications also  
245 appear in the OH adsorption region. First, the sharp spike observed at 0.80 V for the Pt(111)  
246 disappears as the Pd concentration increases, and the onset for OH adsorption seems to shift to  
247 lower potential values. However, the general shape of the process is mainly preserved.

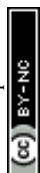
248 These voltammograms can also be compared to those obtained for Pt(111) electrodes  
249 modified with Pd sub-monolayers ( $\text{Pd}_{\text{sub}}\text{-Pt(111)}$ )<sup>22</sup>. Although the qualitative behavior is similar,  
250 that is, additional peaks appear in the hydrogen adsorption region, and the OH adsorption region  
251 is displaced to lower potential values, quantitative differences can be observed. First, for the Pd  
252 submonolayers, the new peak appears at 0.28 V for very low Pd coverages and splits into two as  
253 the coverage increases. On the other hand, the shape in the OH region is also significantly  
254 different. For very low coverages, the sharp peak for Pt(111) disappears and a new peak appears  
255 at 0.71 V. All these facts clearly indicate that the energetics and probably the distribution of the  
256 Pd atoms on the surface are significantly different on both surfaces:  $\text{Pt}_{100-x}\text{Pd}_x(111)$  and  $\text{Pd}_{\text{sub}}\text{-Pt(111)}$ .  
257

258 Changes in the cyclic voltammograms of  $\text{Pt}_{100-x}\text{Pd}_x(111)$  are more obvious in sulfuric acid  
259 solutions (Figure 2 (A2)). The voltammetric profiles in perchloric and sulfuric acid overlap for  
260 the same Pd concentration below 0.3 V, which necessarily implies that hydrogen  
261 adsorption/desorption processes are responsible for the voltammetric signals in this region (Figure  
262 S2). However, the signals appearing between 0.25 and 0.4 V significantly increase their intensity  
263 as compared to perchloric acid solutions. This fact means that the sulfate adsorption/desorption  
264 process is responsible for the current increase in this region. In parallel, the signals at  $E > 0.4$  V  
265 diminish. Thus, sulfate adsorption, which on the Pt(111) takes place between 0.35 and 0.60 V, is  
266 displaced at significantly lower potentials as the Pd content in the electrode increases. One  
267 important feature that disappears is the sharp peak associated with the order/disorder transition  
268 for the sulfate adlayer on Pt(111) at 0.5 V. Scanning tunneling microscopy (STM) studies have



269 revealed that sulfate forms an ordered adlayer with a  $(\sqrt{7}\times\sqrt{3})R19.1^\circ$  on several fcc (111) facets  
270 <sup>49–54</sup>. The phase transition from the disordered to the ordered structure is marked by the presence  
271 of sharp spikes, as those observed at the completion of the adsorption process on Pt(111) <sup>49</sup>,  
272 Au(111) <sup>50,51</sup>, and Pd(111) <sup>52</sup>. The initial adsorption process leads to the formation of very small,  
273 ordered domains, which grow progressively as the adsorption proceeds. This results in the  
274 formation of a large number of rotational and translational domain boundaries with a sulfate  
275 coverage lower than that expected for a perfectly ordered adlayer. At a certain potential, the  
276 different ordered domains coalesce to form large, ordered domains. In this process, the coverage  
277 increases abruptly, giving rise to the observed spikes in the voltammogram. For this process to  
278 take place, the adsorbed sulfate molecules should be able to move freely across the surface, so  
279 that adsorbed species at a domain boundary move from one domain to the neighboring one, so  
280 that an ordered domain grows at the expense of the second one. Furthermore, the mobility of  
281 sulfate requires that the adsorption energy of all the different surface sites should be the same.  
282 Because of this, the spike is very sensitive to the presence of defects or contaminants on the  
283 surface since they alter adsorption energies and restrict the movements of species on the surface.  
284 In fact, for Pt(111) and Au(111) their sharpness is indicative of the surface order (quality of the  
285 electrode and surface cleanliness). The disappearance of the spike is clearly a consequence of the  
286 different adsorption energies of sulfate on the different sites. The comparison between Pt(111),  
287 Pt<sub>100-x</sub>Pd<sub>x</sub>(111), and Pd(111) electrodes indicates that sulfate is more strongly adsorbed on Pd  
288 sites. The adsorption of sulfate will involve several atoms, and owing to the low concentration of  
289 Pd in the prepared alloys, sulfate will interact mainly with two types of ensembles: pure Pt  
290 ensembles and ensembles containing one Pd atom. In this latter type of ensembles, sulfate  
291 adsorption will be stronger. Then, the movement of sulfate across the surface is hindered because  
292 of this energy difference between the adsorption sites, which prevents the disorder/order  
293 transition.

294 When these voltammograms are compared to those of Pd<sub>sub</sub>-Pt(111), significant  
295 differences are also observed. On Pd<sub>sub</sub>-Pt(111), there is a sharp spike at ca. 0.24 V, whose full



296 width at half maximum (fwhm) is smaller than 3 mV and the spike corresponding to the phase  
297 transition of sulfate on Pt(111) is still visible (although significantly smaller) for coverages close  
298 to 1<sup>55</sup>. These facts indicate that the Pd adlayers on the Pt(111) have an island growth mode. Thus,  
299 the sharp spike is due to the competitive adsorption of hydrogen and sulfate on the Pd island,  
300 whereas the size of the free-Pd domains is large enough to give rise to a phase transition. Clearly,  
301 the situation for Pt<sub>100-x</sub>Pd<sub>x</sub>(111) is different. The disappearance of the peak related to the phase  
302 transition even for low Pd concentrations indicates that the distribution of Pd on the surface is  
303 homogeneous. Moreover, the wide peaks related to the competitive adsorption of hydrogen and  
304 sulfate at  $E > 0.3$  V suggest the presence of different Pd-Pt environments.

305 Pt(100) is a more complex surface than Pt(111) in terms of electrochemical  
306 characterization since different voltammetric profiles can be obtained for Pt(100) depending on  
307 the cooling conditions after the flame-annealing treatment<sup>56</sup>. In this work, reductive cooling  
308 conditions have been used for all surfaces to avoid oxygen adsorption and obtain two-  
309 dimensionally ordered terraces. The reconstruction of the Pt(100) surface, which is eliminated  
310 upon the immersion of the electrode in the electrolyte, gives rise to the formation of mesas on the  
311 electrode surface with the excess of atoms in the hex reconstruction with respect to the (1×1)  
312 structure. The absence of the reconstruction on the Pt<sub>100-x</sub>Pd<sub>x</sub>(100) electrodes with moderate Pd  
313 content (Figure 1 (B)) suggests that the surface order of these electrodes should be higher than  
314 that of the pure Pt(100) electrode. It should be noted that, in the case of the reconstructed surfaces,  
315 the reconstruction is lifted upon immersion in water, and the resulting surface structure is stable  
316 between 0.0 and 0.9 V.

317 Figure 2 (B) displays the voltammetric profiles of Pt(100) and Pt<sub>100-x</sub>Pd<sub>x</sub>(100) surfaces in  
318 perchloric and sulfuric acid solutions, illustrating a different and characteristic profile depending  
319 on the Pd content of the alloy and the electrolyte used. On Pt(100) electrodes, the signals below  
320 0.4 V are assigned mainly to hydrogen adsorption, whereas the wave between 0.45 and 0.6 V  
321 corresponds to adsorbed OH. As the Pd content increases, there is a shift of the hydrogen  
322 adsorption to lower potentials, as can be seen in Figure 2 (B). On the other hand, the signals in  
323 the OH adsorption region also diminish. In this case, two different scenarios can be considered:



324 either OH adsorption is disfavored on the  $\text{Pt}_{100-x}\text{Pd}_x(100)$  alloy electrodes or OH adsorption takes  
325 place at lower potentials overlapping with the final stages of hydrogen adsorption and thus, the  
326 signals in the voltammogram between 0.3 and 0.4 V can also contain contributions from OH  
327 adsorption. This issue will be revisited when CO displacement experiments are analyzed. Again,  
328 this profile is different from that obtained for  $\text{Pd}_{\text{sub}}\text{-Pt}(100)$  in this media <sup>13</sup>.

329 The presence of sulfuric acid in the electrolyte brings about significant changes in the  
330 voltammogram due to the specific adsorption of sulfate. Below 0.3 V, the voltammograms for  
331  $\text{Pt}(100)$  in sulfuric and perchloric acid solutions overlap (Figure S2), clearly indicating that the  
332 signals up to 0.3 V are related to the adsorption/desorption of hydrogen. However, above this  
333 value, a sharp peak is observed in sulfuric acid. This peak is due to the competitive adsorption of  
334 hydrogen and sulfate on the surface. A similar peak is also observed for  $\text{Pd}(100)$  surfaces <sup>45</sup>. It  
335 has been shown that the replacement of one species by the other gives rise to sharp peaks <sup>57</sup>. As  
336 the Pd content increases, this sharp peak diminishes and finally disappears. From the comparison  
337 between the voltammograms in perchloric and sulfuric acid media (Figure S2), it can be seen that  
338 although the voltammograms overlap at the low potential limit, there is a significant increase in  
339 the currents between 0.2 and 0.4 V in the latter. This increase should be ascribed to the adsorption  
340 of sulfate. The absence of the sharp peak should be related to the heterogeneity of the surface sites  
341 and the different site requirements for hydrogen and sulfate adsorption. Whereas hydrogen  
342 adsorption occupies only one site, adsorbed sulfate is coordinated to several sites. Thus, changes  
343 in the surface energy induced by the presence of Pd affect more importantly hydrogen adsorption,  
344 because it depends only on the specific site composition. For sulfate adsorption, due to the small  
345 Pd content, this energy change is less important due to the involvement of several sites.

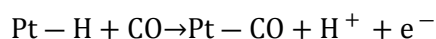
### 347 3.3. CO displacement experiments on the $\text{Pt}_{100-x}\text{Pd}_x$ single crystal surfaces.

348 To obtain more information on the nature of adsorbed species on the single-crystal  $\text{Pt}_{100-}$   
349  $x\text{Pd}_x$  surfaces, CO displacement experiments were carried out and the results were compared with  
350 those obtained for pure Pt electrodes with the same surface orientation. Due to the strong  
351 interaction of the sulfate ion with Pt and  $\text{Pt}_{100-x}\text{Pd}_x$  surfaces, the experiments were performed in





352 HClO<sub>4</sub> solution by dosing gaseous CO in the cell atmosphere at a fixed potential. Figure S3 shows  
353 the current transients for the CO displacement experiments on Pt<sub>100-x</sub>Pd<sub>x</sub>(111) surfaces, at 0.1, 0.2,  
354 and 0.3 V. The experimentally measured displacement charges correspond to oxidation processes  
355 associated with the oxidative displacement of adsorbed hydrogen, according to:



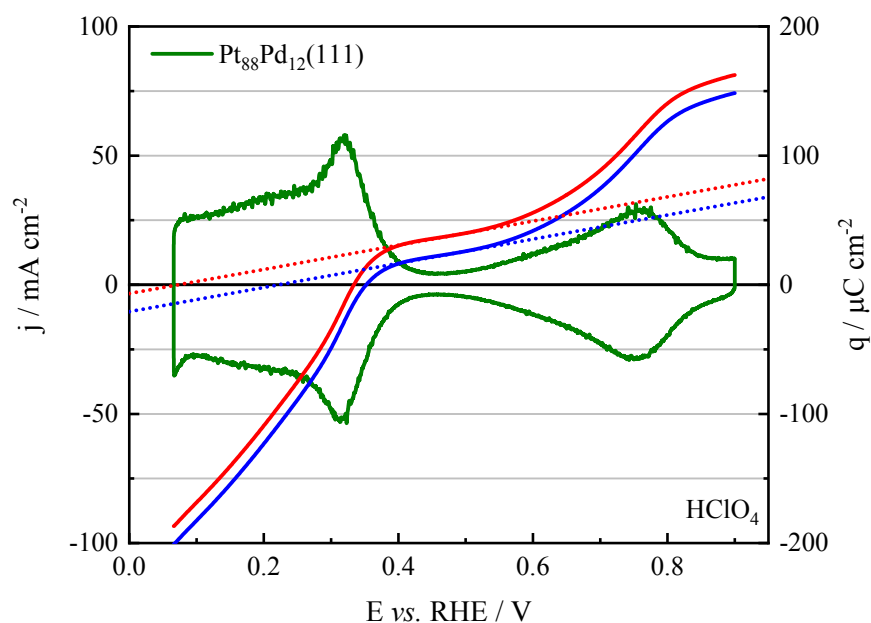
356  
357 Charge density-potential curves can be represented by combining the displaced charge and the  
358 charges calculated from the voltammetric profile in Figure 2 (A), using the equation<sup>58,59</sup>:

$$q_E = \int_{E^*}^E \frac{|j|}{\nu} - q_{dis, E^*}$$

359  
360 where  $q_E$  is the total charge at the applied potential  $E$ ,  $j$  is the average value of the cathodic or  
361 anodic current density,  $\nu$  is the scan rate (in V·s<sup>-1</sup>), and  $q_{dis, E^*}$  is the charge displaced by CO at  
362 the potential  $E^*$ , which is used as the integration constant.

363 Figure 3 shows the voltammetric profile of the Pt<sub>88</sub>Pd<sub>12</sub>(111) electrode together with the  
364 charge density-potential curves. The displaced charge density measured at 0.3 V has been used  
365 as the integration constant to obtain this curve. The experimental charges displaced at 0.1 and  
366 0.2 V nicely overlap with the integrated curve (see Table S1), validating the use of the CO  
367 displacement experiments to measure the total charge on the electrode<sup>60</sup>. As expected, increasing  
368 the electrode potential results in a rise in the charge density.





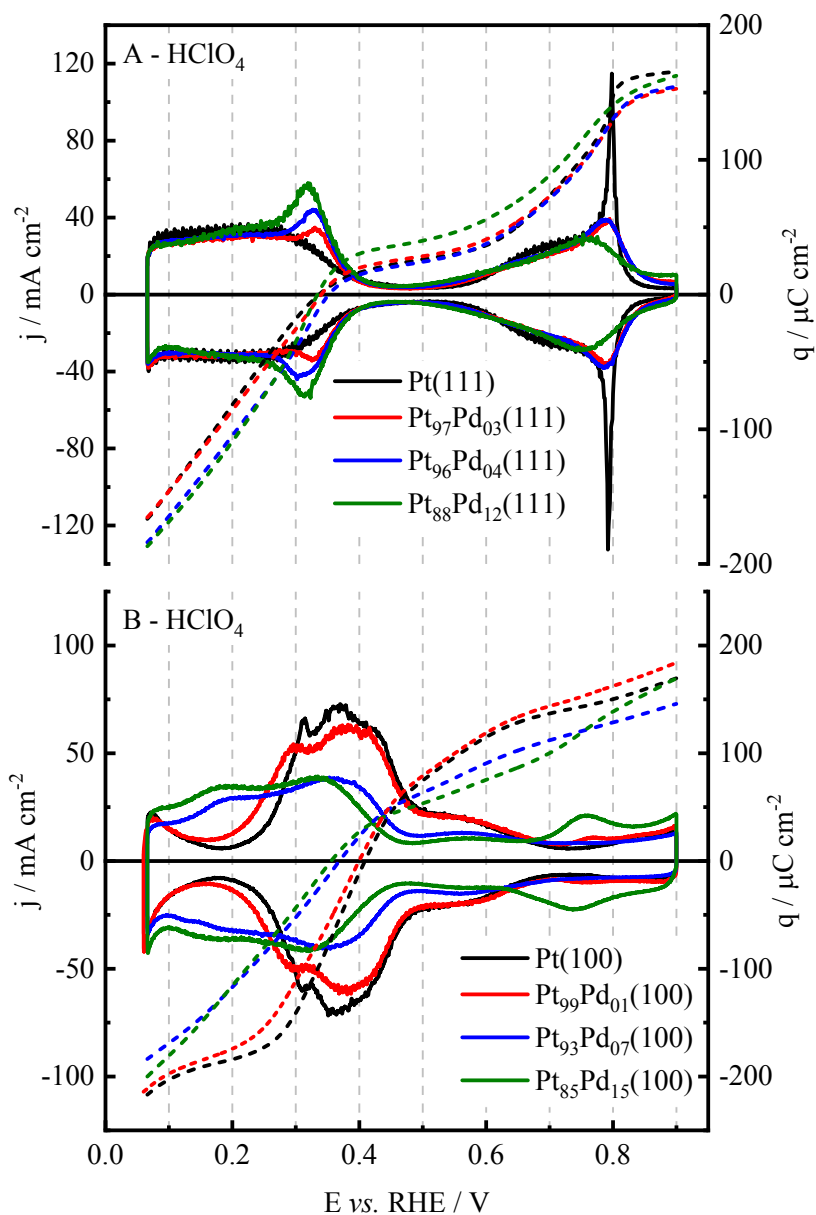
369  
 370 Figure 3 Cyclic voltammetric profile (green line) and total charge curve (red line) for the  
 371  $\text{Pt}_{88}\text{Pd}_{12}(111)$  surface in  $0.1 \text{ mol L}^{-1} \text{ HClO}_4$  as a function of the potential recorded at a scan rate  
 372 of  $50 \text{ mV s}^{-1}$ . The blue line is a corrected charge density taking into account the remaining charge  
 373 on the CO-covered surface (see ref. <sup>47</sup>).

374  
 375 The charge curves for Pt(111), and all the  $\text{Pt}_{100-x}\text{Pd}_x(111)$  surfaces are shown in Figure  
 376 4 (A). Several conclusions can be drawn from these measurements. First, the charge density at  
 377  $0.1 \text{ V}$  increases in absolute value with the Pd content. The same trend is observed at  $0.2 \text{ V}$ ,  
 378 showing that, for a given potential, the hydrogen coverage increases with the Pd content. For the  
 379 Pt(111) electrode, the charge measurements indicate that the hydrogen coverage at  $0.06 \text{ V}$ , the  
 380 onset of hydrogen evolution, is ca.  $2/3$ . In this case, the increase of the charge at  $0.06 \text{ V}$  for  
 381  $\text{Pt}_{88}\text{Pd}_{12}(111)$  allows the estimation of a coverage value of ca.  $0.75$ . It should be noted that, due  
 382 to the small difference in the lattice parameters of Pt and Pd (ca.  $1\%$ ), the expected difference in  
 383 the charge of a process exchanging  $1 \text{ e}$  per surface atom on Pt(111) and Pd(111) is ca.  $2\%$ , which  
 384 is significantly smaller than the value measured for the charge increase at the onset of hydrogen  
 385 evolution. For this reason, the increase in the charge value should be assigned to a change in the



386 hydrogen coverage related to the energetics of the hydrogen adsorption process. On the other  
387 hand, the comparison at 0.06 V with the Pd(111) electrode is not possible due to the hydrogen  
388 absorption on Pd which takes place at  $E < 0.2$  V. However, the measured charge density for the  
389 Pd(111) electrode in perchloric acid solution at 0.225 V is ca.  $-125 \mu\text{C cm}^{-2}$ , which is higher  
390 (in absolute value) than that measured for the  $\text{Pt}_{100-x}\text{Pd}_x(111)$  surface. Thus, as the Pd content  
391 increases, the hydrogen coverages at a given potential increase with the Pd content for (111)  
392 surfaces.





393

394 Figure 4 Cyclic voltammetric profile (full lines) and the total charge curve (dotted lines) for (A)  
 395  $\text{Pt}_{100-x}\text{Pd}_x(111)$ , and (B)  $\text{Pt}_{100-x}\text{Pd}_x(100)$  surfaces in  $0.1 \text{ mol L}^{-1} \text{ HClO}_4$  as a function of the potential  
 396 recorded at a scan rate of  $50 \text{ mV s}^{-1}$ .

397

398

399

The charge curves allow for obtaining the pztc of the different electrodes, which is the  
 potential at which the total charge displayed in Figure 4 (A) is zero. A more accurate value is



400 obtained when the charge curves are corrected by the residual charge of the CO-covered electrode  
401 as shown in <sup>47,61</sup>. For that, it will be assumed that the correction is the same one as that used for  
402 the Pt(111) electrode. The charge curve after correction is shown in Figure 3. The values of the  
403 pztc for the different electrodes are summarized in Table 2 and the comparison of the corrected  
404 charge curves for all the electrodes is shown in Figure S3. The values for the Pt(111) electrode  
405 are the same as those reported in the literature for this electrolyte solution <sup>47</sup>. Regarding the  
406 evolution of the potential of zero total charge for the different (111) surfaces (Table 2), for very  
407 low Pd content the values are constant within the experimental error of the measurements. On the  
408 other hand, for the Pt<sub>88</sub>Pd<sub>12</sub>(111) surface, a significant diminution is observed. In fact, for the  
409 Pd(111) electrode in the same solution, the uncorrected pztc is ca. 0.28 V <sup>45</sup>. Moreover, a  
410 significant positive charge density, ca. 50  $\mu\text{C cm}^{-2}$ , is measured in the apparent double layer  
411 region of the Pd(111) electrode ( $E > 0.35$  V). This fact indicates that OH is already adsorbed on  
412 the Pd(111) electrode at 0.35 V and that the signals corresponding to the final stages of the  
413 hydrogen desorption overlap with those of the OH adsorption. A similar situation is observed for  
414 the Pt<sub>88</sub>Pd<sub>12</sub>(111), for which the total charge at 0.4 V is significantly higher than that observed for  
415 the other electrodes. Owing to the capacity of Pd(111) for adsorbing OH at low potentials, it can  
416 be proposed that the peak centered at ca. 0.32-0.36 V for the alloy electrodes is due to the  
417 competitive adsorption/desorption of H and OH on the Pd atoms.

418 Other important data that can be obtained from the total charge curves is the OH coverage.  
419 As can be seen, for the two alloy electrodes with low Pd content, the charge curves almost overlap  
420 with that of the Pt(111) electrode up to the potential at which the sharp peak for the Pt(111) surface  
421 is observed (ca. 0.8 V). From that point, the charge for the Pt(111) electrode is higher than that of  
422 the two other electrodes, since the sharp peak is absent for the Pd alloy electrodes. The behavior  
423 is similar to that observed for the phase transition in the sulfate layer. Thus, it can be proposed  
424 that the sharp peak at ca. 0.8 V for the Pt(111) electrode is due to a phase transition in the OH  
425 layer. It should be noted that, in those processes, other solution species, such as cations <sup>62,63</sup> or  
426 other anions <sup>64</sup>, are involved, indicating that the interfacial structure is much more complicated



427 than a simple layer of adsorbed anions. For the Pt<sub>88</sub>Pd<sub>12</sub>(111) electrode, the final OH coverage is  
428 similar to that of the Pt(111) electrode, owing to the significant adsorption of OH on the Pd sites,  
429 which takes place on the peak at 0.32 V.

430 For the Pt(111) electrode, as the process of OH and H adsorption are separated by ca.  
431 0.25 V in a region where there are no specifically adsorbed species on the surface, it was possible  
432 to estimate the potential of zero free charge of the electrode (pzfc)<sup>47</sup>. The pzfc is the potential at  
433 which the surface charge of the electrode is zero and can be related to the electronic properties of  
434 the material<sup>65</sup>. On the other hand, the pzfc contains the contribution of the charge exchanged due  
435 to the adsorbed species on the surface and that corresponding to the charge at the metal surface.  
436 To estimate the pzfc, the behavior of the charge curve in the region where no species are adsorbed  
437 on the electrode surface (the so-called double layer region) has to be extrapolated, to the point  
438 where a charge zero is obtained. The corrected curve, the extrapolated line, and the pzfc for the  
439 Pt<sub>88</sub>Pd<sub>12</sub>(111) electrode are shown in Figure 3, and the values of the pzfc for the different  
440 electrodes are given in Table 2. As before, the values obtained for the Pt(111) electrode agree  
441 with those of the literature<sup>47</sup>. On the other hand, for the electrodes with a low Pd content, the pzfc  
442 increases. This increase has been observed for Pt(111) surfaces modified with adatoms, due to the  
443 disruption of the water layer on the Pt(111) surface induced by the presence of a foreign species  
444<sup>66,67</sup>. On the other hand, the estimated value of the pzfc for the Pt<sub>88</sub>Pd<sub>12</sub>(111) electrode is  
445 significantly lower. Two possible reasons can be foreseen. First, the work function of Pd(111) is  
446 lower than that of Pt(111). Due to the linear relationship between the work function and the pzfc  
447<sup>68</sup>, it is expected that an increase in the Pd content results in a diminution of the pzfc. For the very  
448 low Pd coverages, the effect of the disruption of the water adlayer dominates over the change in  
449 the electronic properties, as has been shown for adatoms on Pt(111) with similar electronic effects  
450 to Pd<sup>67</sup>. On the other hand, it is also possible that the determination of the pzfc on the  
451 Pt<sub>88</sub>Pd<sub>12</sub>(111) surface is biased by the presence of some adsorbed OH in the region used for the  
452 extrapolation, resulting in lower values.



453 Table 2 Values of the pztc and pzfc for the Pt<sub>100-x</sub>Pd<sub>x</sub>(111) surfaces.View Article Online  
DOI: 10.1039/D4TA01771D

SURFACE	Pztc / V (Uncorrected)	Pztc / V (Corrected)	Pzfc / V (Uncorrected)	Pzfc / V (Corrected)
Pt(111)	0.3365 ± 0.0005	0.385 ± 0.0005	0.160 ± 0.010	0.341 ± 0.010
Pt <sub>97</sub> Pd <sub>03</sub> (111)	0.346 ± 0.001	0.385 ± 0.001	0.164 ± 0.010	0.350 ± 0.010
Pt <sub>96</sub> Pd <sub>04</sub> (111)	0.352 ± 0.001	0.395 ± 0.001	0.204 ± 0.010	0.375 ± 0.010
Pt <sub>88</sub> Pd <sub>12</sub> (111)	0.334 ± 0.003	0.352 ± 0.003	0.073 ± 0.010	0.222 ± 0.010

454

455 CO displacement experiments were also carried out for the (100) electrodes. Transients  
 456 for the different electrodes can be found in Figure S4 and the charge curves for the different (100)  
 457 electrodes are shown in Figure 4 (B). Unlike what is observed for the (111) electrodes, the  
 458 hydrogen coverage at constant potential diminishes as the Pd content increases. The diminution  
 459 is significant at 0.2 V because, at this potential value the hydrogen coverage has almost reached  
 460 the expected value of 1 monolayer for the Pt(100) electrode. On the other hand, the differences at  
 461 0.06 V, the onset of hydrogen evolution, are less important, and the values of the hydrogen charge  
 462 are close to 1 monolayer for the studied electrodes.

463 The curves also allow for calculating the pztc, which can be used to determine whether  
 464 the signals observed at potentials higher than 0.3 V are related to the OH adsorption process.  
 465 Again, the values for the Pt(100) electrode agree with those reported previously<sup>31</sup>. For these  
 466 electrodes, only the pztc can be determined since the estimation of the pzfc requires a region in  
 467 which no adsorbed species are present on the surface. As will be shown, for the Pt<sub>100-x</sub>Pd<sub>x</sub>(100)  
 468 surfaces, the hydrogen, and OH adsorption/desorption processes overlap, preventing any  
 469 reasonable estimation of the pzfc. The values of the pztc obtained from Figure 4 (B) are given in  
 470 Table 3. As can be seen, the pztc is located between 0.4 and 0.3 V, in a region where important  
 471 current signals are observed in the voltammograms. Thus, the final part of the signals observed



472 in the voltammograms, are due to the OH adsorption process, and the final stages of hydrogen  
 473 desorption coincide with the initial stages of OH adsorption. On the other hand, the pztc shifts to  
 474 lower values as the Pd content increases. This means that OH adsorption is favored at lower  
 475 potentials with the addition of Pd. However, the OH coverage at 0.5 or 0.6 V for the different  
 476 electrodes diminishes with the Pd content, as the lower charge density for these electrodes  
 477 indicates. Thus, in the presence of Pd, the initial adsorption of OH is favored, probably on the Pd  
 478 sites, likely due to the Pd effect on the surrounding Pt atoms, the completion of the OH adsorption  
 479 adlayer on Pt is not favored.

480 **Table 3.** Values of the pztc for the  $\text{Pt}_{100-x}\text{Pd}_x(100)$  surfaces.

SURFACE	Pztc / V (Uncorrected)
Pt(100)	$0.407 \pm 0.003$
$\text{Pt}_{99}\text{Pd}_{01}(100)$	$0.400 \pm 0.006$
$\text{Pt}_{93}\text{Pd}_{07}(100)$	$0.369 \pm 0.005$
$\text{Pt}_{85}\text{Pd}_{15}(100)$	$0.355 \pm 0.006$

481

### 482 3.4. CO oxidation on the electrodes.

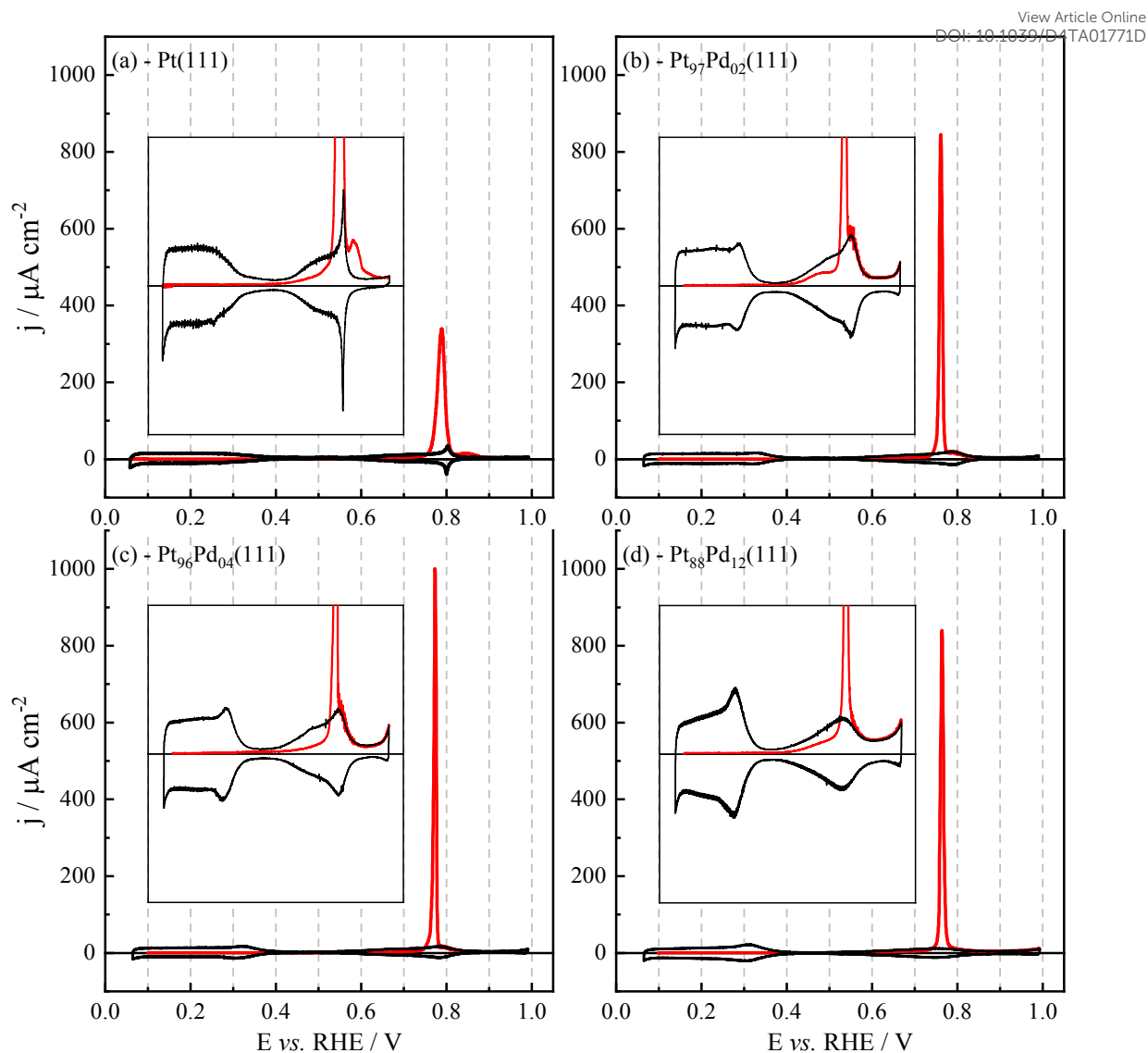
483 The electrooxidation of CO on platinum is a well-explored reaction due to its frequent  
 484 occurrence as a poisoning species in various electrochemical reactions, particularly during the  
 485 oxidation of organic compounds or fuels. This reaction is highly sensitive to surface structure  
 486 variations, and in this study, it has been used to further characterize the properties of the single-  
 487 crystal electrodes. Figure 5 shows the voltammetric profiles for CO oxidation on Pt(111) and  
 488  $\text{Pt}_{100-x}\text{Pd}_x(111)$  surfaces. Studies on Pd(111) showed that CO oxidation occurs at potentials higher  
 489 than 0.9 V<sup>45</sup>, unlike what is observed here for  $\text{Pt}_{100-x}\text{Pd}_x(111)$  surfaces, where  $\text{CO}_{\text{ad}}$  is fully  
 490 oxidized at potentials lower than 0.83 V. From these curves, two different parameters can be  
 491 extracted: the CO oxidation charge and the peak potential. For the CO oxidation charge, it should  
 492 be noted that the oxidation peak contains charge contributions not only from the CO oxidation





493 process but also those related to the changes in the double-layer charge from a CO-covered  
494 electrode to a clean surface. Thus, during the oxidation process, the total charge of the electrode  
495 changes from that obtained for a CO-covered electrode to that registered on the clean surface.  
496 This correction is equal to the uncorrected total charge measured at 0.9 V. Thus, the integrated  
497 charge for the CO oxidation peak between the initial potential and 0.9 V should be corrected by  
498 the uncorrected total charge at 0.9 V. Peak potential values and CO oxidation charges are given  
499 in table S3. As can be seen, the charge values for the (111) surfaces are around  $330 \mu\text{C cm}^{-2}$ ,  
500 which results in a coverage of ca. 0.69. For the Pt(111) electrode, in the absence of CO in the  
501 solution, STM results show that the CO adlayer forms an ordered  $\sqrt{19} \times \sqrt{19}$ -13 CO with a CO  
502 coverage of 0.68<sup>69</sup>. Thus, the results indicate that the CO coverage is nearly independent of the  
503 Pd content, and probably, the same CO adlayer structure is obtained. However, a significant  
504 diminution of the peak potential is observed when Pd is introduced in the adlayer. Model  
505 calculations for the CO oxidation process in voltammetry on the Pt(111) electrode show that the  
506 peak potential is very dependent on the number of initial defects on the CO adlayer<sup>70</sup>. Since CO  
507 oxidation on Pt takes place according to a Langmuir-Hinshelwood mechanism between adsorbed  
508 CO and adsorbed OH, the initial stages require the presence of some defects in the adlayer where  
509 OH adsorption can occur. As mentioned before, the presence of Pd creates sites with different  
510 adsorption energy and thus, the mobility of the adsorbed species to form ordered structures is  
511 hindered. Thus, it is expected that the CO adlayer in the Pd-containing electrodes has more defects  
512 than that on a perfect Pt(111) surface. The consequence of the higher number of defects in the CO  
513 adlayer is the lower onset and peak potential for the CO oxidation process.





514

515 Figure 5 Cyclic voltammetric profile for CO oxidation, at  $0.02 \text{ V s}^{-1}$ , on  $\text{Pt}_{100-x}\text{Pd}_x(111)$  surfaces.

516 The insets show a magnification of the voltammetric profiles.

517

518 The CO oxidation study was also carried out on  $\text{Pt}(100)$  and  $\text{Pt}_{100-x}\text{Pd}_x(100)$  surfaces under  
 519 the same experimental conditions. The cyclic voltammetric profiles for CO oxidation on (a)  
 520  $\text{Pt}(100)$  or (b-d)  $\text{Pt}_{100-x}\text{Pd}_x(100)$  surfaces are shown in Figure 6. For this family of electrodes, CO  
 521 has been completely oxidized at  $0.80 \text{ V}$ <sup>45</sup> (maximum potential used in the studies of the previous  
 522 topics for these surfaces), and the charges were integrated up to  $0.85 \text{ V}$ . Values of the integrated  
 523 charges and peak potentials are given in table S4. The value for the  $\text{Pt}(100)$  electrode agrees with  
 524 previous results<sup>71,72</sup>. The increase of the Pd content leads to changes in the CO oxidation behavior.

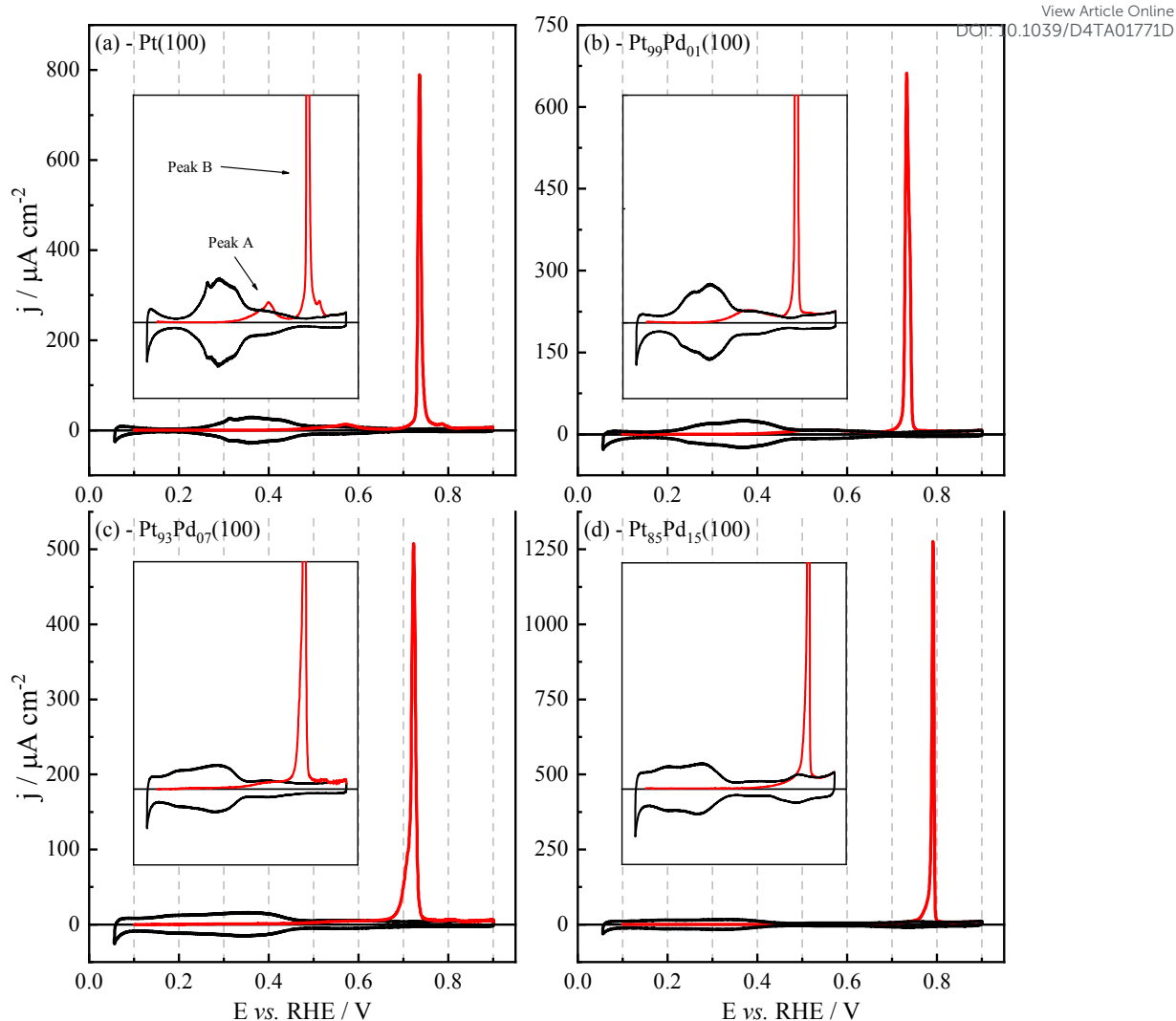
25



525 First, the CO oxidation charge increases from 330 to ca. 350  $\mu\text{C cm}^{-2}$ . Thus, the CO coverage  
526 increases from 0.75 to 0.83. Another observed change in the voltammetric profiles is the presence  
527 of a pre-wave, as shown in the insets of Figure 6. As can be seen, the currents in the pre-wave  
528 region diminish with increasing Pd content. These pre-oxidation waves have been assigned to the  
529 presence of surface defects<sup>73,74</sup>. According to this, the presence of defects on the CO adlayer  
530 diminishes with the content of Pd. As mentioned before, the lifting of the reconstruction when the  
531 (100) electrodes are immersed in the electrolyte solution leads to the formation of mesas, whose  
532 borders act as surface defects. The absence of reconstruction for the Pt<sub>85</sub>Pd<sub>15</sub>(100) electrode  
533 should give rise to a much-ordered surface structure, with a much lower number of defects,  
534 justifying the absence of the pre-wave for this electrode.

535 The Pd content also affects the peak potential. As shown in Table S4, the peak potential  
536 for the CO oxidation peak first diminishes to increase for the Pt<sub>85</sub>Pd<sub>15</sub>(100). According to the  
537 Langmuir-Hinshelwood model for CO oxidation, the peak potential for CO oxidation is governed  
538 by three factors<sup>70</sup>: CO coverage, number of defects on the adlayer, and energetics of OH  
539 adsorption. Normally, an increase in the coverage and a diminution of defects in the adlayer, as  
540 observed for the Pt<sub>99</sub>Pd<sub>01</sub>(100) and Pt<sub>93</sub>Pd<sub>07</sub>(100), are associated with a higher peak potential.  
541 However, for these electrodes, a diminution of the peak potential is observed. Then, the reason  
542 for the diminution should be a more favorable OH adsorption, as observed from the CO  
543 displacement experiments, which show that the onset for OH adsorption is displaced to lower  
544 potential values. On the other hand, a significant increase in the peak potential is observed from  
545 the Pt<sub>85</sub>Pd<sub>15</sub>(100). In this case, the potential increase should be assigned to the significant  
546 diminution of the number of defects in the adlayer, which overcompensates the more favorable  
547 OH adsorption.





548  
549 Figure 6 Cyclic voltammetric profile for CO oxidation, at  $0.02 \text{ V s}^{-1}$ , on  $\text{Pt}_{100-x}\text{Pd}_x(100)$  surfaces.

550 The insets show a magnification of the voltammetric profiles.

551

#### 552 4. Conclusions

553 The validation of Clavilier's method for manufacturing well-defined Pt-Pd alloy single-  
554 crystal surfaces has been achieved through a combination of electrochemical and UHV surface  
555 science characterization methods. Insights into the nature of the adsorbed species in different  
556 acidic electrolytes, namely, perchloric and sulfuric acids, have been attained via the analysis of  
557 the voltammetric profiles and the determination of the pztc. Our results reveal that anion  
558 interaction with the alloyed surfaces is stronger in comparison to pure platinum. Although the  
559 pztc of  $\text{Pt}_{100-x}\text{Pd}_x(111)$  surfaces remains almost constant with increasing Pd content, it has been



560 found that hydrogen desorption overlaps with the adsorption of OH, which takes place at already  
561 0.35 V. On Pt<sub>100-x</sub>Pd<sub>x</sub>(100) surfaces, the OH adsorption is favored at lower potentials, as pointed  
562 out by the shift of the pztc to lower values as the Pd content increases. CO oxidation experiments  
563 indicate that the CO coverage and adlayer structure on the electrodes with (111) orientation are  
564 similar to those on Pt(111). However, a lower onset for the CO oxidation process is observed due  
565 to the earlier adsorption of OH. Similar behavior for the onset of CO oxidation was found for  
566 Pt<sub>100-x</sub>Pd<sub>x</sub>(100) surfaces, although higher CO coverages are achieved on these electrodes due to  
567 the well-ordered surface structure induced by the inclusion of Pd on the Pt(100) structure. Various  
568 reaction mechanisms are significantly influenced by the presence of OH<sub>ads</sub> species, including the  
569 oxidation reactions of organic molecules and oxygen reduction reaction, with OH<sub>ads</sub> being  
570 identified as the principal intermediate. Therefore, the fundamental insights here reported on  
571 platinum-palladium electrodes are of particular importance for the proper interpretation of the  
572 catalytic properties of these materials.

573

#### 574 **Author Contributions**

575 **Gabriel Melle:** Investigation, Methodology, Writing – review & editing. **Fabian Scholten:**  
576 Investigation, Methodology, Writing – review & editing. **Juan M. Feliu:** Writing – review  
577 & editing., **Enrique Herrero:** Writing – review & editing, Supervision. **Beatriz**  
578 **RoldanCuenya:** Writing – review & editing. **Rosa M. Arán-Ais** Conceptualization,  
579 Methodology, Investigation, Writing – review & editing, Supervision.

580

#### 581 **Conflicts of interest**

582 The authors declare no competing financial interest.

#### 583 **Acknowledgment**

584 The authors thank the funding by the Max Planck Society through the funding of the Max  
585 Planck Partner Group on “Mechanistic insights into electrocatalytic reactions on well-defined



586 bimetallic surfaces". Financial support from the Ministerio de Ciencia e Innovación (project  
587 PID2022-137350NB-I00) is also acknowledged.

588

## 589 Notes and references

590 1 C. Roth, A. J. Papworth, I. Hussain, R. J. Nichols and D. J. Schiffrin, *J. Electroanal.*  
591 *Chem.*, 2005, **581**, 79–85.

592 2 E. Herrero, A. Fernández-Vega, J. M. Feliu and A. Aldaz, *J. Electroanal. Chem.*, 1993,  
593 **350**, 73–88.

594 3 M. Mavrikakis, B. Hammer and J. K. Nørskov, *Phys. Rev. Lett.*, 1998, **81**, 2819–2822.

595 4 G. B. Melle, T. Altair, R. L. Romano and H. Varela, *Energy & Fuels*, 2021, **35**, 6202–  
596 6209.

597 5 A. Capon and R. Parson, *J. Electroanal. Chem. Interfacial Electrochem.*, 1973, **44**, 1–7.

598 6 R. Parsons and T. VanderNoot, *J. Electroanal. Chem. Interfacial Electrochem.*, 1988, **257**,  
599 9–45.

600 7 C. A. Rice, A. Bauskar and P. G. Pickup, in *Electrocatalysis in Fuel Cells*, 2013, vol. 9,  
601 pp. 69–87.

602 8 C. A. Rice and A. Wieckowski, 2013, pp. 43–67.

603 9 X. Chen, L. P. Granda-Marulanda, I. T. McCrum and M. T. M. Koper, *Nat. Commun.*,  
604 2022, **13**, 38.

605 10 N. de Z. M. J. N. Pourbaix, J. Van Muylder, *Platin. Met. Rev.*, 1959, **3**, 47.

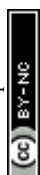
606 11 M. Grdeń, M. Łukaszewski, G. Jerkiewicz and A. Czerwiński, *Electrochim. Acta*, 2008,  
607 **53**, 7583–7598.



- 608 12 B. A. F. Previdello, E. Sibert, M. Maret and Y. Soldo-Olivier, *Langmuir*, 2017, **33**, 2087–  
609 2095. View Article Online  
DOI: 10.1039/D4TA01771D
- 610 13 M. J. Llorca, J. M. Feliu, A. Aldaz and J. Clavilier, *J. Electroanal. Chem.*, 1993, **351**, 299–  
611 319.
- 612 14 R. Gómez, A. Rodes, J. M. Pérez, J. M. Feliu and A. Aldaz, *Surf. Sci.*, 1995, **344**, 85–97.
- 613 15 B. Álvarez, A. Berná, A. Rodes and J. M. Feliu, *Surf. Sci.*, 2004, **573**, 32–46.
- 614 16 X. Chen, K. Ojha and M. T. M. Koper, *JACS Au*, 2023, **3**, 2780–2789.
- 615 17 C. Ye, F. Dattila, X. Chen, N. López and M. T. M. Koper, *J. Am. Chem. Soc.*, 2023, **145**,  
616 19601–19610.
- 617 18 X. Chen, L. P. Granda-Marulanda, I. T. McCrum and M. T. M. Koper, *Chem. Sci.*, 2020,  
618 **11**, 1703–1713.
- 619 19 D. J. Watson and G. A. Attard, *Electrochim. Acta*, 2001, **46**, 3157–3161.
- 620 20 D. J. Watson and G. A. Attard, *Surf. Sci.*, 2002, **515**, 87–93.
- 621 21 T. J. Schmidt, N. M. Markovic, V. Stamenkovic, P. N. Ross, G. A. Attard and D. J.  
622 Watson, *Langmuir*, 2002, **18**, 6969–6975.
- 623 22 B. Álvarez, V. Climent, A. Rodes and J. M. Feliu, *Phys. Chem. Chem. Phys.*, 2001, **3**,  
624 3269–3276.
- 625 23 R. Rizo, E. Herrero, V. Climent and J. M. Feliu, *Curr. Opin. Electrochem.*, 2023, **38**,  
626 101240.
- 627 24 D. S. Mekazni, R. M. Arán-Ais, A. Ferre-Vilaplana and E. Herrero, *ACS Catal.*, 2022, **12**,  
628 1965–1970.



- 629 25 Y. Yang, R. G. Agarwal, P. Hutchison, R. Rizo, A. V. Soudackov, X. Lu, E. Herrero, J. M. Feliu, S. Hammes-Schiffer, J. M. Mayer and H. D. Abruña, *Nat. Chem.*, 2023, **15**, 271–  
630 277. View Article Online  
DOI: 10.1039/D4TA01771D
- 631
- 632 26 V. Briega-Martos, E. Herrero and J. M. Feliu, *Electrochim. Acta*, 2017, **241**, 497–509.
- 633 27 J. Clavilier, D. Armand, S. G. Sun and M. Petit, *J. Electroanal. Chem. Interfacial*  
634 *Electrochem.*, 1986, **205**, 267–277.
- 635 28 D. J. Watson and G. A. Attard, *Surf. Sci.*, 2002, **515**, 87–93.
- 636 29 F. J. Vidal-Iglesias, A. Al-Akl, D. Watson and G. A. Attard, *J. Electroanal. Chem.*, 2007,  
637 **611**, 117–125.
- 638 30 D. J. Watson and G. A. Attard, *Surf. Sci.*, 2002, **515**, 87–93.
- 639 31 R. M. Arán-Ais, M. C. Figueiredo, F. J. Vidal-Iglesias, V. Climent, E. Herrero and J. M.  
640 Feliu, *Electrochim. Acta*, 2011, **58**, 184–192.
- 641 32 J. M. Orts, R. Gómez, J. M. Feliu, A. Aldaz and J. Clavilier, *Electrochim. Acta*, 1994, **39**,  
642 1519–1524.
- 643 33 T. B. Massalski, H. Okamoto and A. S. M. International, *Binary alloy phase diagrams*,  
644 ASM International, Materials Park, Ohio SE -, 2nd ed., 1990.
- 645 34 A. V Ruban, H. L. Skriver and J. K. Nørskov, *Phys. Rev. B*, 1999, **59**, 15990–16000.
- 646 35 Z. Liu, Y. Li, X. Zhang, S. Rao, J. Li, W. Wang, Z. Sun and J. Yang, *ACS Appl. Mater.*  
647 *Interfaces*, 2022, **14**, 28816–28825.
- 648 36 S. A. Krasnikov, S. Murphy, N. Berdunov, A. P. McCoy, K. Radican and I. V Shvets,  
649 *Nanotechnology*, 2010, **21**, 335301.
- 650 37 M. Wakisaka, M. Sugimasa, J. Inukai and K. Itaya, *J. Electrochem. Soc.*, 2003, **150**, E81.





- 651 38 T. E. Felter and A. T. Hubbard, *J. Electroanal. Chem. Interfacial Electrochem.*, 1979, **100**,  
652 473–491. Open Access Article Online  
DOI: 10.1039/D4TA01771D
- 653 39 G. W. Simmons, Y. N. Wang, J. Marcos and K. Klier, *J. Phys. Chem.*, 1991, **95**, 4522–  
654 4528.
- 655 40 R. Rizo, S. Pérez-Rodríguez and G. García, *ChemElectroChem*, 2019, **6**, 4725–4738.
- 656 41 A. Ferre-Vilaplana, J. V. Perales-Rondón, C. Buso-Rogero, J. M. Feliu and E. Herrero, *J.*  
657 *Mater. Chem. A*, 2017, **5**, 21773–21784.
- 658 42 T. Solomun, *J. Electroanal. Chem. Interfacial Electrochem.*, 1987, **217**, 435–441.
- 659 43 T. Solomun, *J. Electroanal. Chem. Interfacial Electrochem.*, 1988, **255**, 163–177.
- 660 44 T. Solomun, *J. Electroanal. Chem. Interfacial Electrochem.*, 1991, **302**, 31–46.
- 661 45 M. Hara, U. Linke and T. Wandlowski, *Electrochim. Acta*, 2007, **52**, 5733–5748.
- 662 46 R. Gisbert, G. García and M. T. M. Koper, *Electrochim. Acta*, 2010, **55**, 7961–7968.
- 663 47 R. Rizo, E. Sitta, E. Herrero, V. Climent and J. M. Feliu, *Electrochim. Acta*, 2015, **162**,  
664 138–145.
- 665 48 K. N. da Silva and E. Sitta, *J. Solid State Electrochem.*, 2020, **24**, 1921–1926.
- 666 49 U. Stimming and R. Vogel, , DOI:[https://doi.org/10.1016/S0022-0728\(96\)05051-6](https://doi.org/10.1016/S0022-0728(96)05051-6).
- 667 50 A. Cuesta, M. Kleinert and D. M. Kolb, 2000, 5684–5690.
- 668 51 K. Sato, S. Yoshimoto, J. Inukai and K. Itaya, 2006, **8**, 725–730.
- 669 52 L. Wan, T. Suzuki, K. Sashikata, J. Okada, J. Inukai and K. Itaya, 2000, **484**, 189–193.
- 670 53 Y. G. Kim, J. B. Soriaga, G. Vigh and M. P. Soriaga, *J. Colloid Interface Sci.*, 2000, **227**,



- 671 505–509.
- 672 54 L. Wan, 1995, 9507–9513.
- 673 55 J. Clavilier, 2002, **4**, 379–383.
- 674 56 A. Rodes, M. A. Zamakhchari, K. El Achi and J. Clavilier, *J. Electroanal. Chem.*  
675 *Interfacial Electrochem.*, 1991, **305**, 115–129.
- 676 57 N. Garcia-Araez, J. J. Lukkien, M. T. M. Koper and J. M. Feliu, *J. Electroanal. Chem.*,  
677 2006, **588**, 1–14.
- 678 58 V. Climent, R. Gomez, J. M. Orts, A. Rodes, A. Aldaz and J. M. Feliu, 1999.
- 679 59 R. M. Arán-Ais, M. C. Figueiredo, F. J. Vidal-Iglesias, V. Climent, E. Herrero and J. M.  
680 Feliu, *Electrochim. Acta*, 2011, **58**, 184–192.
- 681 60 R. Rizo, J. Fernández-Vidal, L. J. Hardwick, G. A. Attard, F. J. Vidal-Iglesias, V. Climent,  
682 E. Herrero and J. M. Feliu, *Nat. Commun.*, 2022, **13**, 2550.
- 683 61 M. J. Weaver, S.-C. Chang, L.-W. H. Leung, X. Jiang, M. Rubel, M. Szklarczyk, D.  
684 Zurawski and A. Wieckowski, *J. Electroanal. Chem.*, 1992, **327**, 247–260.
- 685 62 N. Garcia-Araez, V. Climent, P. Rodriguez and J. M. Feliu, *Phys. Chem. Chem. Phys.*,  
686 2010, **12**, 12146.
- 687 63 V. Climent, N. García-Araez and J. M. Feliu, *Electrochem. commun.*, 2006, **8**, 1577–1582.
- 688 64 A. BERNA, V. CLIMENT and J. FELIU, *Electrochem. commun.*, 2007, **9**, 2789–2794.
- 689 65 S. Trasatti and E. Lust, in *Modern Aspects of Electrochemistry*, 2002, pp. 1–215.
- 690 66 V. Climent, *Electrochem. commun.*, 2001, **3**, 590–594.
- 691 67 N. García-Araez, V. Climent and J. M. Feliu, *J. Am. Chem. Soc.*, 2008, **130**, 3824–3833.



- 692 68 William M. Haynes, *HANDBOOK OF CHEMISTRY and PHYSICS*, CRC Press: 97<sup>th</sup> Edition  
693 edição (24 junho 2016), 97th edn., 2554. DOI: 10.1039/D4TA01771D
- 694 69 I. Villegas and M. J. Weaver, *J. Chem. Phys.*, 1994, **101**, 1648–1660.
- 695 70 C. A. Angelucci, E. Herrero and J. M. Feliu, *J. Phys. Chem. C*, 2010, **114**, 14154–14163.
- 696 71 K. Domke, E. Herrero, A. Rodes and J. M. Feliu, *J. Electroanal. Chem.*, 2003, **552**, 115–  
697 128.
- 698 72 C. K. Rhee, J. M. Feliu, E. Herrero, P. Mrozek and A. Wieckowski, *J. Phys. Chem.*, 1993,  
699 **97**, 9730–9735.
- 700 73 A. López-Cudero, A. Cuesta and C. Gutiérrez, *J. Electroanal. Chem.*, 2005, **579**, 1–12.
- 701 74 M. J. S. Farias, B. A. V. Lima, G. Tremiliosi-Filho and E. Herrero, *J. Electroanal. Chem.*,  
702 2021, **896**, 115382.
- 703

