See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/277892288

In-Situ Monitoring of Electrooxidation Processes at a Gold Single Crystal Surface Using Shell-Isolated Nanoparticle-Enhanced Raman Spectroscopy

ARTICLE in JOURNAL OF THE AMERICAN CHEMICAL SOCIETY - JUNE 2015

Impact Factor: 12.11 · DOI: 10.1021/jacs.5b04670

READS

202

10 AUTHORS, INCLUDING:



Rajapandiyan Panneerselvam Xiamen University

14 PUBLICATIONS 16 CITATIONS

SEE PROFILE



Zhilin Yang

Xiamen University

326 PUBLICATIONS 4,270 CITATIONS

SEE PROFILE



Zhong-Qun Tian

Xiamen University

354 PUBLICATIONS 9,823 CITATIONS

SEE PROFILE



Communication

pubs.acs.org/JACS

In Situ Monitoring of Electrooxidation Processes at a Gold Single Crystal Surface Using Shell-Isolated Nanoparticle-Enhanced Raman Spectroscopy

- ⁴ Chao-Yu Li,^{†,#} Jin-Chao Dong,^{†,#} Xi Jin,[†] Shu Chen,[‡] Rajapandiyan Panneerselvam,[†] ⁵ Alexander V. Rudnev,[§] Zhi-Lin Yang,[‡] Jian-Feng Li,*,^{†,§} Thomas Wandlowski,[§] and Zhong-Qun Tian[†]
- 6 [†]MOE Key Laboratory of Spectrochemical Analysis and Instrumentation, State Key Laboratory of Physical Chemistry of Solid 7 Surfaces, College of Chemistry and Chemical Engineering, and [‡]Department of Physics, Xiamen University, Xiamen 361005, China
- 8 SDepartment of Chemistry and Biochemistry, University of Bern, Freiestrasse 3, Bern CH-3012, Switzerland
- Supporting Information

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27 28

29

ABSTRACT: Identifying the intermediate species in an electrocatalytic reaction can provide a great opportunity to understand the reaction mechanism and fabricate a better catalyst. However, the direct observation of intermediate species at a single crystal surface is a daunting challenge for spectroscopic techniques. In this work, electrochemical shell-isolated nanoparticle-enhanced Raman spectroscopy (EC-SHINERS) is utilized to in situ monitor the electrooxidation processes at atomically flat Au(hkl) single crystal electrode surfaces. We systematically explored the effects of crystallographic orientation, pH value, and anion on electrochemical behavior of intermediate (AuOH/ AuO) species. The experimental results are well correlated with our periodic density functional theory calculations and corroborate the long-standing speculation based on theoretical calculations in previous electrochemical studies. The presented in situ electrochemical SHINERS technique offers a unique way for a real-time investigation of an electrocatalytic reaction pathway at various well-defined noble metal surfaces.

lean energy, along with the booming economy and expanding population, has received great attention among 32 the research community concerned with environmental issues. 33 The fundamental understanding of the electrocatalytic process, 34 the core of fuel cell and electrolysis, will be greatly beneficial to the fabrication of novel catalysts with high efficiency. There is a dearth of knowledge about electrocatalysis; the formation of an 37 oxide film at a metal surface will decisively influence the activity 38 of a catalyst and the reaction mechanism. However, the 39 process of surface electrooxidation is complicated, and the 40 conventional characterization techniques for the identification 41 of key intermediates, adsorbed oxygen species, or hydroxyl ions 42 are still limited by the real-time performance and sensitivity. It 43 is noteworthy to mention that OH- species play a key role in 44 oxide film formation, O₂ reduction, ^{1d,3} and electrooxidation of 45 hydrogen. 4 Because of its importance, the oxidation of gold 46 electrodes by means of electroreflectance spectroscopy was 47 investigated by Nguyen et al.⁵ Though the first direct evidence 48 for the specific adsorption of OH- on gold electrode was 49 obtained, electroreflectance studies were unable to monitor the

real-time oxidation process on an electrode surface. Therefore, 50 it is imperative to investigate the adsorption behavior of 51 hydroxide species on an electrode surface in a precise manner. 52 Although various reaction mechanisms have been proposed 53 based on electrochemical techniques and theoretical calcu- 54 lations, it is still a challenge to prove these results by direct 55 evidence of in situ investigations at an atomically flat 56 surface. 1d,2,6

Surface-enhanced infrared reflection absorption spectroscopy 58 (SEIRAS) is widely used in electrochemical interfaces 59 investigation, 1b,7 but it is difficult to apply an IR technique at 60 low wavenumbers below 800 cm⁻¹, where we can obtain the 61 direct vibrational information on a molecule-metal bond, 62 because of the absorption of infrared light by the thin solution 63 layer and optical window. Surface-enhanced Raman spectros- 64 copy⁸ (SERS) is a unique analytical technique, which has been 65 employed in electrochemical interface investigation, monitor- 66 ing reactions, ^{9c} biological analysis, ¹⁰ and other significant 67 applications. ¹¹ It inherits the advantage of acquiring Raman 68 spectra and can obtain the bonding information at low 69 wavenumbers without any interference from the ensemble. 70 Notably, several efforts have been made using SERS to obtain 71 the spectroscopic information on surface-oxygen species, 72 unfortunately SERS is seriously limited to roughened nano- 73 structured surfaces. 12 Particularly, single crystal surfaces are 74 commonly preferred and used in surface science, because of 75 their well-defined surface state and optic field. In addition, the 76 information obtained from a single crystal surface will be greatly 77 helpful to examine the orientation of adsorbates and to unravel 78 the fundamental surface reactions unambiguously by using 79 surface selection rules. In this contribution, tip-enhanced 80 Raman spectroscopy¹³ (TERS) and attenuated total reflec- 81 tion¹⁴ (ATR) methods have been utilized to accrue spectral 82 details at single crystal surfaces. Unfortunately, TERS was not 83 able to be applied at electrochemical interfaces because of the 84 interference from the tip immersed in solution and the low 85 enhancement by only one tip so that only a few molecules 86 could be used, whereas the latter technique is cumbersome and 87 the enhancement factor (EF) is only 1 to 2 orders of 88 magnitude. Therefore, a detailed description for the adsorption 89

Received: May 5, 2015



90 behavior of an intermediate species such as the OH⁻ species at 91 single crystal electrode surface remains elusive.

In 2010, our group invented a versatile tool called "shell-93 isolated nanoparticle-enhanced Raman spectroscopy" 94 (SHINERS)¹⁵ to open up new avenues in electrochemical 95 interfaces, by resolving the disadvantages associated with TERS 96 and ATR. Remarkably, shell-isolated nanoparticles (SHINs) 97 render discernible Raman signals from single crystal surfaces 98 with significant sensitivity, stability and reproducibility. ^{15,16} As a 99 new technique, EC-SHINERS excels in the investigation of 100 miscellany of adsorbates at single crystal surfaces and provides 101 clear understanding about electrocatalytic processes.

Herein, we employ the in situ EC-SHINERS technique to monitor the surface oxidation processes at low-index Au(*hkl*) 104 single crystal electrodes and systematically evaluate the 105 influence of crystallographic orientation, anion, and pH during 106 electrooxidation. Thus, the formation of hydroxide film is 107 clearly elucidated with in situ EC-SHINERS.

Figure 1 manifests the process of in situ EC-SHINERS at 109 low-index Au(hkl) surfaces. The SHINs utilized in this work

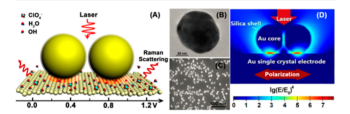


Figure 1. (A) Schematic diagram of in situ EC-SHINERS on low-index $\operatorname{Au}(hkl)$ surfaces. (B) HR-TEM image of $\operatorname{Au@SiO_2}$ nanoparticle. (C) SEM image of $\operatorname{Au}(111)$ single crystal electrode surface modified with SHINs. (D) 3D-FDTD simulations of four SHIN NPs with a model of 2×2 array on an Au substrate.

110 consist of uniform Au nanoparticles (~55 nm) with a 111 chemically grown SiO₂ shell (~2 nm). The size distribution 112 of more than 100 SHINs is shown in Supporting Information, 113 Figure S1. The as-prepared SHINs were carefully drop-casted 114 onto a freshly prepared Au(hkl) single crystal half-bead 115 electrode, and the electromagnetic simulations were performed 116 using the finite-difference time-domain (FDTD) method. Four 117 shell-isolated nanoparticles are placed on a gold single crystal 118 substrate and illuminated using a linearly polarized plane wave 119 with electric field amplitude of 1 V/m. As the electric field 120 distribution shows (Figure 1D), the hot spots are located in the 121 particle-film junctions under the 633 nm laser line. The modified electrode was subsequently mounted in a custom-made spectroelectrochemical cell. Pt and Ag/AgCl electrodes were used as counter and reference electrodes, respectively. 125 After the modification of SHINs on the electrode surface, a 126 hydrogen evolution reaction (HER) cleaning procedure is 127 applied to remove the possible contamination. The electrochemical response of the modified single crystal surface is 129 unaffected, which is evidenced by the cyclic voltammograms (CV). 16a All potentials are reported with respect to Ag/AgCl 131 electrode in this paper. The average enhancement factor of this 132 configuration is about 1.1×10^6 on the Au(110) surface. ^{16b} 133 Further details of the experiments can be found in the 134 Supporting Information.

Figure 2 shows the CV of the Au(111) electrode in deaerated 136 0.1 M NaClO₄ solution (pH was tuned to ~9 with NaOH) at 137 room temperature and corresponding in situ SHINERS spectra

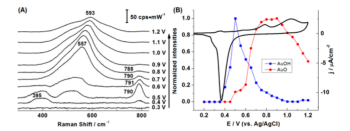


Figure 2. (A) In situ EC-SHINERS spectra of electrooxidation of the Au(111) surface in 0.1 M $NaClO_4$ (pH is \sim 9). (B) Normalized EC-SHINERS intensities of the stretching mode of AuO and the bending mode of AuOH at different potentials. CV of Au(111) electrode in 0.1 M $NaClO_4$ is presented (pH is \sim 9, scan rate is 2 mV/s).

during positive potential scan. During the positive potential 138 excursion, there is no observable Raman signature in the range 139 of 300 to 800 cm⁻¹ until 0.4 V. However, at 0.4 V, a peak 140 begins to appear around 790 cm⁻¹ and the intensity reaches a 141 maximum at ~0.5 V. On the basis of the periodic density 142 functional theory (DFT) calculations on these three low index 143 crystal surfaces, (as shown in the Supporting Information, 144 Table S1 and Figure S6), and the literature data, 12a,d we 145 attribute this band to the gold-hydroxide bending mode $\delta_{ ext{AuOH}}$ 146 of the adsorption on top sites. Generally, hydroxide ions could 147 specifically adsorb on the Au(111) surface and form $AuOH_{ad}$ 148 through one-electron oxidation. Importantly, at 0.5 V, the 149 bending mode of AuOH is strongest which vividly emphasizes 150 the maximum adsorption of the hydroxide ions on the Au(111) 151 surface before their further oxidation to Au oxide. When the 152 potential is higher, the oxidation and deprotonation of AuOH 153 weakens the bending mode of AuOH (~790 cm⁻¹) which 154 diminishes after complete oxidation of the Au (111) surface 155 (>1.0 V). The peak slightly blue shifts from 790 to 791 cm⁻¹ as 156 potential increased from 0.5 to 0.6 V, and red shifts to 788 157 cm⁻¹ as potential increased to 0.9 V. To ascertain our 158 conclusion, a deuterium isotopic substitution measurement 159 (as illustrated in Figure S2) was also carried out, and the 160 bending mode of AuOH at 790 cm⁻¹ shift toward the lower 161 wavenumber 694 cm⁻¹ in deuterated water was observed. This 162 observation clearly implies that the band is attributed to the 163 gold-hydroxide bending mode δ_{AuOH} , and accords well with the l64 earlier reports on the roughened Au surface. In the low- l65 frequency region, a broad band is present at ~360-420 cm⁻¹ 166 when the potential is set at 0.3 V. This band is ~20 cm⁻¹ blue- 167 shifted in deuterated water, which is similar to the experiments 168 in previous publications, and it was assigned to Au-OH 169 stretching. 12b,c

Concurrently, at 0.5 V, a broad hump develops around 489—171 524 cm⁻¹ which confirms the further oxidation of AuOH_{ad} to 172 AuO_{ad}. This is also confirmed by our DFT calculation of the 173 coadsorption model, as shown in Table S1 and Figure S7. 174 Interestingly, the hump at higher wavenumber intensifies with 175 increasing potential. At 0.7–1.2 V, the peak position shifts 176 linearly from 557 to 593 cm⁻¹, and the intensity reaches a 177 maximum at 0.8–0.9 V. It is evident that this potential 178 dependent behavior is derived from gold oxo-species at the 179 electrode surface rather than that in the bulk. As demonstrated 180 by Weaver's group, on the poly-gold electrode surface with ill-181 defined morphology, ^{12b},c the band around 520–580 cm⁻¹ is 182 attributed to the AuO stretching mode. The frequency shift 183 with electrochemical potential of the Au–O stretching mode is 184 more significant than that of the bending mode. This may 185

186 because of the dipole variation of the adsorbates during the 187 Au—O stretching vibration which is parallel to the surface 188 electric field. Apparently, a decrement in Raman signal was 189 observed with the growth of oxide film, because it weakens the 190 electromagnetic coupling between the electrode surface and 191 SHINs. Furthermore, the corresponding electro-reduction 192 processes are followed as shown in Figure S3, and the 193 reappearance of δ_{AuOH} peak is observed. During a negative scan, 194 abundant surface oxide species at high potential were reduced 195 to gold hydroxide gradually, which results in greater content of 196 AuOH_{ad} and thus stronger Raman intensity of δ_{AuOH} was 197 observed.

Herein, the whole surface electrooxidation process at the 199 atomically flat Au(111) surface as well as the potential 200 dependent evolution of AuOH species have been in situ 201 monitored using EC-SHINERS technique. These distinguishing 202 spectral features corroborate the long-term speculation based 203 on theoretical calculations in the electrochemical studies of 204 oxidation intermediates. 2,6a,f,17,19

To examine the effect of pH value on the formation of intermediate AuOH, we carried out controlled experiments with different pH values as well. As depicted in Figure 3, the

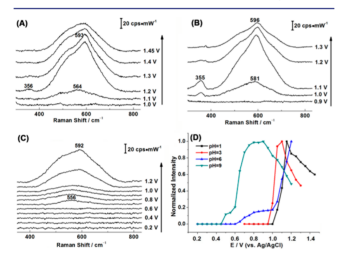


Figure 3. In situ EC-SHINERS spectra of electrooxidation of Au(111) surface during a positive scan in 0.1 M ClO₄⁻ solutions with different pH values. The pH values of the electrolytes in (A) 0.1 M HClO₄, (B) 0.1 M NaClO₄, and (C) 0.1 M NaClO₄ are 1, 3, and 6, respectively. (D) The related potential dependences of normalized Raman intensities for stretching mode of AuO in electrolytes with different pH values.

208 bending mode of AuOH is absent during the entire 209 electrooxidation process when pH values are tuned to about 210 1, 3, and 6 (δ_{AuOH} mode at 796 cm⁻¹ is observable in electrolyte 211 with pH of 11, as shown in Figure S4). But the broad hump 212 around 590 cm⁻¹ remained in acid solution which confirms the 213 contribution from the stretching mode of AuO. Apparently, the 214 potential-dependent appearance of ν_{AuO} becomes lower with 215 the increasing pH values, during the positive sweep (the 216 potential-dependent appearance of ν_{AuO} in electrolytes with pH 217 values of 1, 3, 6, 9, and 11 are 1.05, 1.0, 0.65, 0.5, and 0.3 V, 218 respectively). This behavior clearly indicates that the onset of 219 oxidation of the Au surface in basic solution is shifted to lower 220 potential. Specifically, with the emergence of ν_{AuO} (potential is 221 higher than 1.0 V), a band at ~355 cm⁻¹ is present and the 222 intensity decreases during extended surface oxidation. The Au-223 OH stretching mode located around 400 cm⁻¹ is different from

the broad band that appeared before the complete surface 224 oxidation in basic solution. This narrow band (at $\sim 355~\text{cm}^{-1}$) 225 could be assigned to bridging oxygen adsorption (Figure 226 S7c). It is evidenced that the surface electro-oxidation 227 mechanisms are rather different under different pH values.

Furthermore, to ascertain the role of anions, we performed 229 the experiments in 0.1 M Na₂SO₄ solution in which the anion 230 ion SO₄ $^{2-}$ is stronger than ClO₄ $^{-}$ to chemically adsorb on the 231 Au surface. As shown in Figure S5, the occurrence potential of 232 δ_{AuOH} in Na₂SO₄ is delayed to ~0.45 V and the Raman 233 intensity decreased as well. Explicitly, we infer that the 234 competitive adsorption between hydroxyl and sulfate ions 235 inhibits the onset of hydroxide formation.

To further investigate the effect of crystallographic 237 orientation, comparative experiments were conducted at three 238 low-index $\mathrm{Au}(hkl)$ single crystal surfaces— $\mathrm{Au}(111)$, $\mathrm{Au}(100)$, 239 and $\mathrm{Au}(110)$ —under identical condition (all electrolytes were 240 deaerated 0.1 M $\mathrm{NaClO_4}$ with a pH value of \sim 9). Evidently, the 241 peak at 790 cm⁻¹ features the existence of the intermediate 242 AuOH species, and that peak is chosen to analyze the 243 electrooxidation process on different single crystal surfaces.

Figure 4 displays the intensities of bending mode δ_{AuOH} on 245 f4 three single crystal surfaces increasing in the order of Au(111) 246

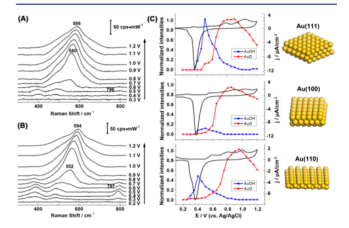


Figure 4. (A,B) EC-SHINERS spectra of electrooxidation of Au(100) and Au(110) surfaces in 0.1 M NaClO₄ (pH is \sim 9). (C) EC-SHINERS intensities of the stretching mode of AuO and the bending mode of AuOH at three low-index Au(hkl) surfaces are all scaled to the same maximum value. The corresponding CVs (scan rate is 2 mV/s) are presented as well.

> Au(110) \gg Au(100), which is consistent with the case of ²⁴⁷ facet-dependent reduction processes presented in Figure S3. ²⁴⁸ The distinct data from our experiments suggest that the ²⁴⁹ energetic favorable formation of hydroxide film on the Au(111) ²⁵⁰ surface occurs through one-electron oxidation of OH⁻ ions. On ²⁵¹ the other hand, the intensity of bending mode δ_{AuOH} increases ²⁵² in the order which is in contrast to the activities of the above ²⁵³ three gold surfaces in the oxygen reduction reaction (the ²⁵⁴ reaction activity sequence is Au(100) \gg Au(110) > ²⁵⁵ Au(111)). ²⁰ In essence, we suppose the formation of hydroxide ²⁵⁶ ions during the reduction of the gold surface may retard the ²⁵⁷ oxygen reduction reaction. A detailed mechanism study of this ²⁵⁸ process is currently underway in our group.

In conclusion, in situ EC-SHINERS, along with the 260 crystallographic orientation, pH, and anion effects, was used 261 to systematically characterize and monitor the electrooxidation 262 process on gold single crystal electrodes. The direct observation 263

264 of the chemical nature of the intermediate, AuOH/AuO 265 species, is achieved with the combination of EC-SHINERS and 266 theoretical modeling. This technique will be further applied to 267 examine electrocatalytic processes at noble metal single crystal 268 surfaces to pave a way for technological innovations in energy 269 materials.

270 **ASSOCIATED CONTENT**

Supporting Information

272 Experimental section, computational details, deuterium isotopic 273 substitution measurement, EC-SHINERS spectra of Au(hkl) 274 electrooxidation and -reduction in 0.1 M NaClO₄ and Na₂SO₄. 275 The Supporting Information is available free of charge on the 276 ACS Publications website at DOI: 10.1021/jacs.5b04670.

277 **AUTHOR INFORMATION**

278 Corresponding Author

279 Li@xmu.edu.cn

280 Author Contributions

²⁸¹ *C.-Y.L. and J.-C.D. contributed equally.

282 Notes

283 The authors declare no competing financial interest.

284 ACKNOWLEDGMENTS

285 We thank Z.Y. Zhou for helpful discussion. This research was 286 supported by Thousand Youth Talents Plan of China, the 287 MOST of China (2010IM040100), the Swiss National Science 288 Foundation (200020-144471, and 200021-124643).

289 REFERENCES

290 (1) (a) Burke, L. D.; Nugent, P. F. Gold Bull. 1998, 31, 39. 291 (b) Osawa, M. Bull. Chem. Soc. Jpn. 1997, 70, 2861. (c) Avramov-Ivić, 292 M.; Jovanović, V.; Vlajnić, G.; Popić, J. J. Electroanal. Chem. 1997, 423, 293 119. (d) Štrbac, S.; Adžić, R. R. J. Electroanal. Chem. 1996, 403, 169. 294 (e) Hughes, M. D.; Xu, Y. J.; Jenkins, P.; McMorn, P.; Landon, P.; 295 Enache, D. I.; Carley, A. F.; Attard, G. A.; Hutchings, G. J.; King, F.; 296 Stitt, E. H.; Johnston, P.; Griffin, K.; Kiely, C. J. Nature 2005, 437, 297 1132. (f) Marković, N. M.; Ross, P. N., Jr. Surf. Sci. Rep. 2002, 45, 117. 298 (g) Björling, A.; Herrero, E.; Feliu, J. M. J. Phys. Chem. C 2011, 115, 299 15509. (h) Wang, D.; Wan, L. J. J. Phys. Chem. C 2007, 111, 16109. 300 (2) Conway, B. E. Prog. Surf. Sci. 1995, 49, 331.

- 301 (3) Attard, G. A.; Brew, A.; Ye, J. Y.; Morgan, D.; Sun, S. G. 302 ChemPhysChem **2014**, 15, 2044.
- 303 (4) Angerstein-Kozlowska, H.; Conway, B. E.; Hamelin, A. J. 304 Electroanal. Chem. Interfac. 1990, 277, 233.
- 305 (5) Nguyen Van Huong, G.; Hinnen, C.; Lecoeur, J. J. Electroanal. 306 Chem. Interfac. 1980, 106, 185.
- 307 (6) (a) Shubina, T. E.; Hartnig, C.; Koper, M. T. M. Phys. Chem.
 308 Chem. Phys. 2004, 6, 4215. (b) Dickinson, T.; Povey, A. F.; Sherwood,
 309 P. M. A. J. Chem. Soc. Faraday Trans. 1975, 71, 298. (c) Vassilev, P.;
 310 Koper, M. T. M. J. Phys. Chem. C 2007, 111, 2607. (d) Pessoa, A. M.;
 311 Fajín, J. L. C.; Gomes, J. R. B.; Cordeiro, M. N. D. S. J. Mol.
 312 Struct.:THEOCHEM 2010, 946, 43. (e) Adžić, R. R.; Marković, N. M.
 313 J. Electroanal. Chem. Interfac 1982, 138, 443. (f) Koper, M. T. M.; van
 314 Santen, R. A. J. Electroanal. Chem. 1999, 472, 126.
- 315 (7) Chen, A.; Lipkowski, J. J. Phys. Chem. B 1999, 103, 682.
- (8) (a) Fleischmann, M.; Hendra, P. J.; McQuillan, A. J. Chem. Phys. 117 Lett. 1974, 26, 163. (b) Jeanmaire, D. L.; Van Duyne, R. P. J. 118 Electroanal. Chem. Interfac. 1977, 84, 1. (c) Tian, Z. Q.; Ren, B.; Wu, 119 D. Y. J. Phys. Chem. B 2002, 106, 9463. (d) McCreery, R. L. Raman 120 Spectroscopy for Chemical Analysis; John Wiley & Sons: 2005.
- 321 (9) (a) Zou, S.; Weaver, M. J. J. Phys. Chem. 1996, 100, 4237. 322 (b) Huang, Y. F.; Wu, D. Y.; Wang, A.; Ren, B.; Rondinini, S.; Tian, Z. 323 Q.; Amatore, C. J. Am. Chem. Soc. 2010, 132, 17199. (c) Li, L.; Steiner, 324 U.; Mahajan, S. Nano Lett. 2014, 14, 495.

(10) (a) Zheng, J.; Jiao, A.; Yang, R.; Li, H.; Li, J.; Shi, M.; Ma, C.; 325 Jiang, Y.; Deng, L.; Tan, W. J. Am. Chem. Soc. 2012, 134, 19957. 326 (b) Guerrini, L.; Krpetić, Ž.; van Lierop, D.; Alvarez-Puebla, R. A.; 327 Graham, D. Angew. Chem., Int. Ed. 2015, 54, 1144. (11) (a) Abdelsalam, M. E.; Mahajan, S.; Bartlett, P. N.; Baumberg, J. 329 J.; Russell, A. E. J. Am. Chem. Soc. 2007, 129, 7399. (b) Hu, J.; Tanabe, 330 M.; Sato, J.; Uosaki, K.; Ikeda, K. J. Am. Chem. Soc. 2014, 136, 10299. 331 (12) (a) Kim, J.; Gewirth, A. A. J. Phys. Chem. B 2006, 110, 2565. 332 (b) Zhang, Y.; Gao, X.; Weaver, M. J. J. Phys. Chem. 1993, 97, 8656. 333 (c) Desilvestro, J.; Weaver, M. J. J. Electroanal. Chem. Interface 1986, 334 209, 377. (d) Li, X.; Gewirth, A. A. J. Am. Chem. Soc. 2003, 125, 7086. 335 (13) (a) Ren, B.; Picardi, G.; Pettinger, B.; Schuster, R.; Ertl, G. 336 Angew. Chem., Int. Ed. 2005, 44, 139. (b) Schmid, T.; Opilik, L.; Blum, 337 C.; Zenobi, R. Angew. Chem., Int. Ed. 2013, 52, 5940. 338 (14) Bruckbauer, A.; Otto, A. J. Raman Spectrosc. 1998, 29, 665. 339 (15) Li, J. F.; Huang, Y. F.; Ding, Y.; Yang, Z. L.; Li, S. B.; Fan, F. R.; 340 Zhang, W.; Zhou, Z. Y.; Wang, Z.; Tian, Z. Q. Nature 2010, 464, 392. 341 (16) (a) Li, J. F.; Rudnev, A.; Fu, Y.; Bodappa, N.; Wandlowski, T. 342 ACS Nano 2013, 7, 8940. (b) Li, J. F.; Ding, S. Y.; Yang, Z. L.; Bai, M. 343 L.; Anema, J. R.; Wang, X.; Wang, A.; Wu, D. Y.; Ren, B.; Hou, S. M.; 344 Wandlowski, T.; Tian, Z. Q. J. Am. Chem. Soc. 2011, 133, 15922. 345 (c) Honesty, N. R.; Gewirth, A. A. J. Raman Spectrosc. 2012, 43, 46. 346 (d) Butcher, D. P.; Boulos, S. P.; Murphy, C. J.; Ambrosio, R. C.; 347 Gewirth, A. A. J. Phys. Chem. C 2012, 116, 5128. (17) Strbac, S.; Hamelin, A.; Adzić, R. R. J. Electroanal. Chem. 1993, 349 362. 47 350 (18) Bublitz, G. U.; Boxer, S. G. Annu. Rev. Phys. Chem. 1997, 48, 351 (19) Angerstein-Kozlowska, H.; Conway, B. E.; Barnett, B.; Mozota, 353 J. J. Electroanal. Chem. Interface 1979, 100, 417. 354 (20) Schmidt, T. J.; Stamenkovic, V.; Arenz, M.; Markovic, N. M.; 355 Ross, P. N., Jr. Electrochim. Acta 2002, 47, 3765.