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# Base-Accelerated Degradation of Nanosized Platinum Electrocatalysts

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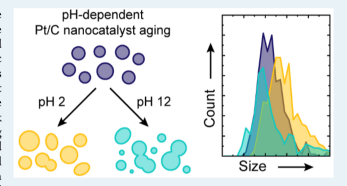
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ABSTRACT: In the pursuit of a hydrogen economy, extensive research has been directed at developing acidic and alkaline hydrogen fuel cells. Such fuel cells often utilize platinum-based catalysts. These materials have been studied extensively in acidic conditions but not in alkaline ones. This focus on acidic systems creates a marked knowledge gap, since recent studies indicate that carbon-supported platinum (Pt/C) electrocatalysts degrade more rapidly in bases than in acids. Addressing this gap, the present work investigates Pt/C degradation at pH 2 and pH 12 using electrochemistry, transmission electron microscopy (TEM), and in situ X-ray absorption spectroscopy (XAS). TEM and XAS reveal accelerated Pt/C degradation at high pH levels, which results in increased Ostwald ripening, Smoluchowski agglomeration, and



nanoparticle detachment. These processes are driven by platinum-catalyzed carbon corrosion and the dissolution and redeposition of platinum nanoparticles. Although these processes take place at both low and high pH levels, basic conditions accelerate the degradation. Base-enhanced Pt dissolution and redeposition was assessed in further detail, revealing an oxidation onset reduction of 100 mV in the base; however, there were no significant differences between undissolved Pt oxidation in acid and in base. The results suggest that soluble Pt oxidation products are stabilized in the base instead. These conclusions are important for translating acid-based literature to alkaline conditions.

KEYWORDS: fuel cells, electrocatalysis, corrosion, catalyst aging, hydrogen economy

#### ■ INTRODUCTION

In an effort to mitigate climate change, governments are stimulating the substitution of fossil fuels with "green" electricity-derived hydrogen.<sup>1</sup> Hydrogen would be used to heat homes, power vehicles, and store renewable energy, thus creating a hydrogen economy.<sup>2</sup> Such a hydrogen economy was already conceived of in the early 1970s,<sup>3</sup> but it is only now nearing economic viability.<sup>4</sup> Facilitating a universally implemented hydrogen economy requires significant cost reductions for green hydrogen production, transportation, storage, and consumption.<sup>1,2</sup>

Affordably producing and utilizing green hydrogen requires efficient electrolyzers and fuel cells. Such devices have lifetimes of tens of thousands of hours, demanding catalysts that are stable for an equally long period. Si,6 Crucially, ensuring catalyst longevity requires a detailed understanding of the degradation mechanisms. This philosophy has spurred extensive research efforts in mapping the aging of platinum-based catalysts because Pt-containing electrodes enable most state-of-the-art fuel cells and electrolyzers. 6-8

Pt aging studies have identified the species that form when platinum is oxidized in acidic electrolytes. Though multiple oxidized Pt species can coexist at a given potential,  $^{9-11}$  it is generally accepted that Pt initially oxidizes to PtOH. This species attains near-monolayer surface coverage between 1.0 and 1.1 V vs the reversible hydrogen electrode (RHE).  $^{9,12-16}$  Pt–OH subsequently oxidizes to PtO $^{9,14}$  and mainly forms  $\alpha$ -PtO $_2$  above  $\sim$ 1.3 V vs RHE.  $^{10,14,15}$  Following  $\alpha$ -PtO $_2$  formation, a variety of Pt oxides can form at higher oxidation potentials and longer polarization times.  $^{14,17-19}$ 

Regardless of the composition of the Pt oxide, modest amounts of Pt dissolve when Pt electrodes are held at voltages above 0.85 V vs RHE through so-called "steady-state" dissolution. More extensive dissolution occurs when Pt oxides are reduced back to their metallic state. Because such dissolution occurs in the short time interval of Pt oxide reduction, it is referred to as "transient dissolution". <sup>21</sup>

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Transient dissolution presumably occurs via the formation of soluble Pt(II) species, <sup>23</sup> which can either be redeposited onto the native Pt electrode or diffuse into the working electrolyte. <sup>24,25</sup>

Although the understanding of transient and steady-state degradation is extensive for Pt in acidic electrolytes, such knowledge is far less comprehensive in alkaline media. This difference is underscored by recent work indicating that acid-based literature does not necessarily translate to Pt in a base: at high pH levels, bulk Pt experiences more transient dissolution than at that at low pH levels. Additionally, carbon-supported platinum (Pt/C) nanoparticles suffer increased Pt-catalyzed corrosion of the carbon support in base. Understanding these degradation processes in base is essential because Pt/C particles are used in large-scale anion-exchange fuel cells that operate under alkaline conditions. Therefore, there is a significant need for studies of Pt catalysts over a wide range of pH values. 7,26,31,32

The present work addresses Pt/C nanoparticle aging in acidic and alkaline electrolytes. The work combines electrochemical characterization, ex situ transmission electron microscopy (TEM), and in situ high-energy resolution fluorescence detection X-ray absorption near-edge structure (HERFD-XANES) spectroscopy. Although the electrochemical results suggest minor differences between the nanoparticle degradation in acid and base, the TEM and HERFD-XANES results point toward strongly accelerated growth and agglomeration in alkaline solutions. This acceleration is attributed to an enhancement of both the carbon corrosion and the diffusion of dissolved Pt species in alkaline media. These results indicate the importance of a combination of ex situ and in situ characterizations of catalysts when quantifying aging. Importantly, our work provides crucial input for the development of technologies to enable the hydrogen economy.

# MATERIALS AND METHODS

**Electrochemical Characterization.** All water used in this study was purified using a Milli-Q Millipore system with a final total organic carbon content below 5 ppb and a resistivity above 18.2 MΩ·cm. Prior to the electrochemical characterization experiments, all glassware was kept in an aqueous solution of 1 g·L<sup>-1</sup> KMnO<sub>4</sub> (Sigma-Aldrich, ACS reagent) and 0.5 M H<sub>2</sub>SO<sub>4</sub> (Sigma-Aldrich, ACS reagent) overnight. This solution was drained on the day of the experiments, and a small amount of H<sub>2</sub>O<sub>2</sub> (Merck, Emprove exp) was used to remove any remaining manganese species from the glassware. The glassware was subsequently boiled in Milli-Q water for 15 min. This 15 min boil was repeated with fresh Milli-Q water until a total of five boiling cycles was reached.

Following cleaning, the electrochemical cell was filled with an aqueous working electrolyte, either 0.1 M KClO<sub>4</sub> (Sigma-Aldrich, ACS reagent) with 0.01 M HClO<sub>4</sub> (Sigma-Aldrich, ACS reagent) to achieve pH 2 or 0.09 M KClO<sub>4</sub> with 0.01 M KOH (Fluka, TraceSelect) to achieve pH 12. The cell contained a glassy carbon (GC) counter electrode (Alfa Aesar, type 1,  $\varnothing=5$  mm, length of 100 mm) and a HydroFlex reversible hydrogen reference electrode (Gaskatel). Prior to electrochemical experiments, the electrochemical cell was deaerated by purging Ar (Airgas, UHP grade) through the working electrolyte for at least 15 min. During experiments, an oxygen-free environment was maintained by flowing Ar over the working solution. In the deaerated electrochemical cell, a GC electrode (Alfa Aesar,  $\varnothing=5$  mm) was used as a working

electrode onto which a small amount of Pt/C (PK Catalyst, 5 wt % Pt) was applied. Prior to application, the Pt/C was suspended at a concentration of 7 mg·mL $^{-1}$  (0.35 mg Pt·mL $^{-1}$ ) in a 40/60 (v/v) mixture of isopropanol (Sigma-Aldrich, for HPLC) and ethanol (Merck, Uvasol). This suspension was ultrasonicated for 1 min, then 2.5  $\mu$ L of the solution was applied to the GC electrode and air-dried.

The working electrode was subjected to the following electrochemical protocol. First, the electrode surface was cleaned electrochemically by running 20 cyclic voltammograms (CVs) between 0.06 and 1.4 V vs RHE at a scan rate of 200 mV·s<sup>-1</sup>. Four additional CV cycles were performed between 0.06 and 0.7 V vs RHE at 50 mV·s<sup>-1</sup> to characterize the state of the Pt nanoparticle surface. Following cleaning and characterization, the Pt nanoparticles were held at 0.7 V vs RHE for one minute to oxidize them. The potential was then swept from the anodic holding potential to 0.06 V vs RHE at a 50 mV·s<sup>-1</sup> scan rate to record the reduction charge of any oxidized Pt species. The Pt nanoparticles were then recharacterized by running five voltammetry cycles between 0.06 and 0.7 V vs RHE at 50 mV· s<sup>-1</sup>. This protocol of a 60-s oxidative hold, a subsequent reductive sweep, and voltammetric characterization was repeated at increasingly positive anodic hold potentials. At every iteration, the hold potential was increased by 0.1 V until a final hold potential of 1.4 V vs RHE was reached.

In this work, all currents and charges are normalized as current and charge densities. When normalizing, the hydrogen desorption charge before the first oxidative hold was used. A specific charge of 200  $\mu\text{C}\cdot\text{cm}^{-1}$  was used for the voltammograms collected at pH 2, whereas a specific charge of 145  $\mu\text{C}\cdot\text{cm}^{-1}$  was used for those collected at pH 12.<sup>33</sup>

Electrochemical Characterization during In Situ HERFD-XANES Experiments. A custom-made electrochemical cell was used for in situ X-ray spectroscopy experiments.<sup>34</sup> Specifically, the cell was comprised of a 4.5 mL cuvette (American Scientific Products, polystyrene) in which a 5 mm diameter hole was drilled to accommodate the working electrode. This electrode consisted of a 20 nm thick Au layer that was sputtered on top of a 5 nm thick Cr adhesion layer on a 1  $\mu$ m thick silicon nitride window (Norcada). This window was encased in a 1 cm × 1 cm frame. On top of the window and the frame, 20  $\mu$ L of the Pt/C suspension was drop-casted and dried under a He stream, thus achieving a Pt loading of 7  $\mu$ g·cm<sup>-2</sup>. This catalyst layer did not contain an ionomer. After applying the Pt/C suspension, the working electrode was affixed to the electrochemical cell using Torr Seal (Varian Inc.). The cell was then filled with 3 mL of the pH 2 or pH 12 solution (see the Electrochemical Characterization subsection). This setup used a Pt wire as the counter electrode and a HydroFlex reversible hydrogen reference electrode (Gaskatel). During experiments, the working electrolyte was continuously purged with Ar gas (Airgas, UHP grade) to maintain oxygenfree conditions.

After mounting the electrochemical XAS cell, the working electrode was electrochemically cleaned by cycling between 0 and 1.4 V vs RHE 20 times at a scan rate of 250 mV·s $^{-1}$ . Four subsequent voltammetry cycles were performed between 0.06 and 0.7 V vs RHE at a scan rate of 50 mV·s $^{-1}$  as well as a final cycle between 0.06 and 1.4 V vs RHE. After acquiring these cleaning and surface characterization CVs, the electrode was held at various potentials in order to acquire steady-state HERFD-XANES spectra.

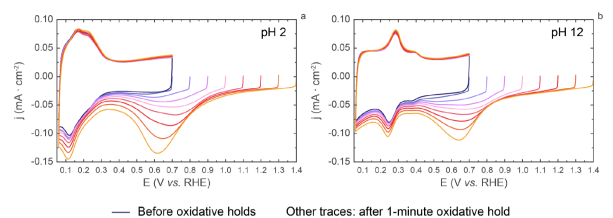


Figure 1. Cyclic voltammograms before and after one-minute holds at various oxidative potentials in (a) pH 2 and (b) pH 12. Electrolytes used were as follows: (a) 100 mM KClO<sub>4</sub> plus 10 mM HClO<sub>4</sub> and (b) 90 mM KClO<sub>4</sub> plus 10 mM KOH. Scan rate was 50 mV·s<sup>-1</sup>.

In Situ HERFD-XANES Experiments. HERFD-XANES experiments were performed at beamline 6-2b of the Stanford Synchrotron Radiation Lightsource. A double-crystal Si(311) monochromator was used to select the energy of the X-ray beam. A Rh-coated parabolic mirror was used to reject higher harmonics and focus the beam on the sample, with a full width at half-maximum of approximately 420  $\mu$ m (horizontally) by 130  $\mu$ m (vertically). Prior to the experiments, the beam energy was calibrated with respect to a metallic Pt foil for which a value of 11563.7 eV was assigned to the first inflection point of the Pt L<sub>III</sub> edge XAS spectrum. During experiments, fluorescent X-rays were detected by a Johann-type X-ray spectrometer. 35 The spectrometer used five Ge(660) spherically bent analyzers (1 m radius of curvature) in order to selectively detect the fluorescent Pt  $L_{\alpha 1}$  X-rays (energy of 9442 eV) with an energy resolution of 1.0 eV. When compared to conventional XAS modes, the energy resolution of the adopted HERFD-XANES provides two critical advantages: a suppression of the intermediate-state lifetime that results in more pronounced spectral fingerprints<sup>36</sup> and a complete rejection of the scattering background, which dramatically enhances the detection sensitivity.

**XANES Data Processing.** The collected HERFD-XANES spectra were aligned in the energy axis using a monochromator glitch feature. These aligned spectra were averaged in the *ATHENA* software<sup>37</sup> and subsequently flattened and normalized by setting the edge jump to one. This normalization was emphasized by plotting all X-ray absorption spectra in *y*-axis units of "edge fraction".<sup>38</sup>

Transmission Electron Microscopy. Transmission electron microscopy was used to analyze the as-received Pt/C and the Pt/C that underwent various amounts of degradation. These aged samples are labeled as "CV treatment", "1.4 V hold", and "extensively aged". The "CV treatment" sample was cycled between 0 and 1.4 V vs RHE 20 times at a scan rate of 250 mV·s<sup>-1</sup>, followed by four voltammetry cycles at 50 mV·s<sup>-1</sup> between 0.06 and 0.7 V vs RHE, two cycles at 50 mV·s<sup>-1</sup> between 0.06 and 1.4 V vs RHE, and two final cycles between 0 and 1.4 V vs RHE. The "1.4 V hold" samples underwent the same procedure as the "CV treatment" samples and were then polarized at 0.4 and 1.4 V vs RHE for 20 min each. Finally, the "extensively aged" sample underwent the same treatment as the "1.4 V hold" sample, followed by successive polarization at 0.0, 0.6, 0.3, 0.0, 0.4, 0.7, 0.8, 0.9, 1.0, 1.1, 1.2, 1.3, and 1.4 V vs

RHE for 20 min each. These potentials match the potential sequence that was used during HERFD-XANES experiments.

Following electrochemistry, the nanoparticle suspension was removed from the glassy carbon working electrode by briefly sonicating the electrode in a mixture of 0.5 mL of water and 0.5 mL of isopropanol. Then, 5  $\mu$ L of this mixture was dropcasted onto a TEM grid (Electron Microscopy Sciences, carbon film on 400 mesh copper) and vacuum-dried. Samples were imaged on an FEI Tecnai transmission electron microscope using an acceleration voltage of 200 kV. The acquired images were used to construct particle-size distributions by analyzing 200 nanoparticles per sample. Particles were sized by tracing their contours in ImageJ and measuring the Feret diameter of the traced particles. If particles had aggregated, the entire aggregate was sized as one particle. Particle sizes are reported as median sizes with single standard deviations. Representative TEM images are displayed in Figure S1 for as-received particles, Figure S2 for particles aged at pH 2, and in Figure S3 for particles aged at pH 12.

#### RESULTS

In studying pH-dependent Pt/C degradation, this work focuses on pH 2 and pH 12. These pH values were obtained by adding either HClO<sub>4</sub> or KOH to a KClO<sub>4</sub> supporting electrolyte. This KClO<sub>4</sub>-based supporting electrolyte was chosen to match the general electrolyte conditions between pH values while minimizing the surface adsorption of electrolyte ions. However, note that completely eliminating electrolyte—surface interactions is not achievable due to ClO<sub>4</sub><sup>-</sup> interacting with adsorbed \*OH in acidic conditions <sup>15,39</sup> and K<sup>+</sup> interacting with adsorbed \*H<sub>2</sub>O or \*OH in alkaline conditions. <sup>40–42</sup> In both acidic and alkaline conditions, Pt/C degradation was studied through electrochemistry, *ex situ* TEM, and *in situ* HERFD-XANES. The results of these studies will be presented in the following subsections.

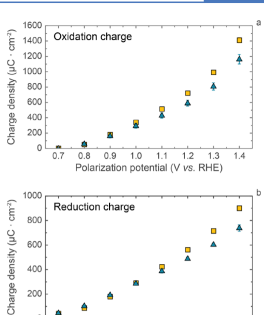
Electrochemical Oxidation and Reduction. Figure 1 displays surface-normalized voltammograms that illustrate Pt oxidation and reduction in acidic and alkaline conditions. These processes were probed electrochemically by holding the working electrode at a given oxidative potential for one minute. Following this positive hold, the working electrode was swept to 0.06 V vs RHE in order to reduce the Pt surface, which produced a reduction peak between 0.4 and 1.0 V vs RHE. Five more cyclic voltammograms were performed between 0.06 and 0.7 V vs RHE to recharacterize the Pt/C surface

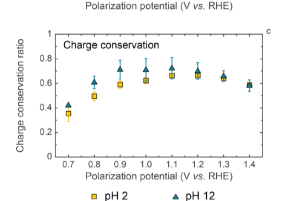
through the hydrogen adsorption and desorption features that are visible below 0.4 V vs RHE.<sup>33</sup> After recharacterizing the surface, the next oxidative hold was performed. This procedure was repeated until the Pt surface was held at 1.4 V vs RHE and then subsequently reduced through a final negative-going voltammetric sweep. In Figure 1, the absolute hydrogen desorption charge between 0.06 and 0.3 V vs RHE varies less than several percent between potential holds, suggesting that no significant amount of the surface area was created or lost during the electrochemical oxidation and reduction protocol.

Figure 1 also highlights other interesting characteristics of the Pt surface reduction. Interestingly, a reduction peak is already visible after mildly anodic polarization at 0.7 V vs RHE. Although this feature occurs in the potential window where the reduction of the carbon support occurs, 43,44 the additional reductive current after a hold at 0.7 V vs RHE likely corresponds to \*OH, as a small amount of \*OH adsorbs on Pt at 0.7 V vs RHE. 15,45 This assertion will further be explored in the In Situ HERFD-XANES section. Notably, this Pt reduction peak between 0.4 and 1.0 V vs RHE grows in size with increasing anodic hold potentials and is actually composed of two peaks in both the acid and the base. This peak duality has been noted before in acid, 11,12 with the earlier (more positive) reduction peak being assigned to \*OH reduction and the later (more negative) peak corresponding to \*O reduction. 11 Precise peak positions for \*OH and \*O reduction cannot be assigned due to the peaks shifting with increasing oxidation potentials. Both the early \*OH reduction feature and the peak duality are seen at pH 2 and pH 12. However, these reduction features appear both larger and sharper in pH 2. It is therefore instructive to quantitatively assess this Pt reduction and the oxidation that precedes it.

A quantitative assessment is presented in Figure 2, where the oxidative charge density from the one minute oxidative holds is presented in panel a. The oxidative charge density increased with the increasing potential and is similar in both the acid and the base until 1.0 V vs RHE. At higher potentials, more oxidative charge is passed in the acid. This trend is matched by the reductive charge density (Figure 2b), which is similar between electrolytes until 1.0 V vs RHE and diverges at higher voltages. This divergence might suggest a change in the dissolution mechanism above 1.0 V vs RHE; in acidic conditions, steady-state Pt dissolution outweighs transient dissolution below 1.0 V vs RHE and is less relevant above this potential.<sup>20</sup>

Pt reduction and oxidation in pH 2 and ph 12 is explored further in Figure 2c, which displays the charge conservation ratio. This ratio was obtained by dividing the reductive charge density in Figure 2b by the oxidative charge density in Figure 2a. The thus-achieved ratio is less than one, indicating that the oxidative charge density is larger than the reductive charge density. This observation is consistent with previous literature in acidic conditions. 12,46,47 The charge conservation ratio increases between 0.7 and 1.0 V vs RHE and then decreases up to 1.4 V vs RHE in both the acid and the base. Within experimental error, the charge conservation ratios in the acid and the base are similar in magnitude at more positive potentials (Figure 2c). The matched ratio between pH 2 and pH 12 demonstrates that the increased oxidative charge density in pH 2 in Figure 2a is proportional to the increase in reduction in Figure 2b. Overall, a strong pH-dependent difference in Pt oxidation and reduction is therefore not apparent from the charge density quantification in Figure 2.





0.9

10 11

0.8

12

Λ

Figure 2. (a) Oxidative charge density during one minute oxidative holds, (b) reductive charge density after one minute oxidative holds, and (c) charge conservation. (a) Oxidative charge densities were determined by integrating the chronoamperometry current density during oxidation. (b) Reductive charge densities were determined by integrating the reductive voltammograms after one-minute oxidative holds. The negative integration limit was 0.4 V, whereas the positive integration limit was the upper voltammetric potential bound. A horizontal baseline was used to compensate for the double layer current. (c) Charge conservation was determined as the ratio between the oxidative and reductive current densities. Each data point is the average of three experiments, and error bars represent one standard deviation.

Though the data in Figure 2 indicate only minor electrochemical pH-dependent Pt oxidation and reduction differences, these data alone do not rule out differences in Pt/C catalyst degradation. Perhaps the most critical note regarding Figure 2 is that the charge conservation ratio does not correlate with the amount of Pt that is dissolved through oxidation and subsequently redeposited. This lack of a one-to-one correlation between Pt charge conservation and Pt atom conservation is due to three reasons. First, part of the recorded charge could partially correspond to the oxidation of the carbon support instead of that singly from Pt. For example, it is known that Pt can electrochemically oxidize its carbon support to  $\mathrm{CO}_2$ . Such an oxidation current will not correspond directly to Pt dissolution. Second, the presented Pt reduction

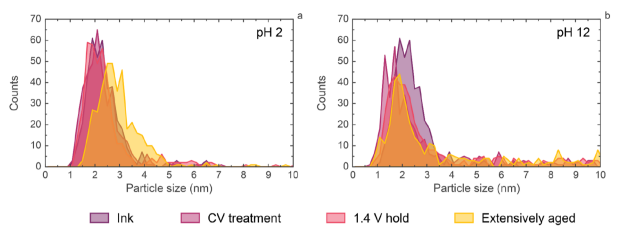


Figure 3. Particle size distributions of the as-received Pt/C nanoparticle ink and those of the same nanoparticles after various electrochemical treatments in (a) pH 2 and (b) pH 12. Each size distribution combines the results of two independent samples, accounting for 200 nanoparticles per sample (400 per distribution). Representative images for each sample are presented in the Supporting Information, including a version of panel b that extends to 30 nm.

data (Figure 2b) were obtained with a lower charge integration limit of 0.4 V vs RHE. This integral therefore ignores the small amount of Pt oxide reduction that overlaps with hydrogen adsorption features (Figure 1) and is challenging to unambiguously background correct. Hence, ignoring this amount of  $PtO_x$  reduction will artificially diminish the calculated reduction charge in Figure 2b and the charge conservation ratio in Figure 2c. Third, and perhaps most importantly, Pt reduction (Figure 2b) corresponds even more strongly to Pt dissolution than Pt oxidation does (Figure 2a), as significant amounts of Pt are dissolved during  $PtO_x$  reduction.  $^{21}$ 

Notably, these three considerations underscore that the data in Figure 2 should not be taken as an indication that Pt/C aging is near-identical in acidic and basic conditions. In fact, Figure 2 indicates that the similarity in acidic and alkaline electrochemistry can mask the well-documented pH-dependent differences in Pt/C degradation. A critical evaluation of Figure 2 therefore indicates a crucial need for complementary analysis techniques when studying Pt/C aging. Such a complementary analysis is provided in the following sections through microscopy and spectroscopy.

Transmission Electron Microscopy. TEM is commonly utilized to visualize nanoparticle size distributions. Such size distributions are shown in Figure 3 for as-received Pt/C nanoparticles as well as nanoparticles that underwent progressive amounts of degradation. The degraded samples were treated as described in the Transmission Electron Microscopy subsection of the Experimental section and are labeled "CV treatment", "1.4 V hold", and "extensively aged". The "CV treatment" protocol mimics the conditions that are typically used to electrochemically clean Pt/C catalysts before experiments, 23,49,50 or to stress-test such nanoparticles. 48,49,51,52 The next sample set in Figure 3 is labeled "1.4 V hold" and identifies nanoparticle degradation after electrochemical cleaning and a single subsequent oxidative potential hold for 20 min. The final degraded sample is labeled "extensively aged" and mimics degradation after a sequence of potential holds that might simulate start/stop catalytic conditions or varying fuel cell loads.<sup>53</sup> Additionally, the "extensively aged" potentials match the potentials that were applied during HERFD-XANES data collection in this work.

Combined, the "extensively aged", "1.4 V hold", and "CV treatment" samples in Figure 3 represent a variety of protocols that are encountered in nanoparticle degradation studies.

Figure 3a shows a narrow size distribution for the asreceived Pt/C nanoparticle ink, which has a mean particle size of 2.2  $\pm$  0.5 nm. After the CV treatment in acid, the mean particle size remains similar at  $2.1 \pm 0.6$  nm. This lack of severe nanoparticle growth after cycling in acid matches previous work.<sup>54</sup> Further aging at 1.4 V vs RHE broadens the nanoparticle size distribution, yielding particles with a mean particle size of  $2.3 \pm 1.0$  nm. Notably, both the 1.4 V hold and the CV treatment leave the main particle size feature relatively intact, as is indicated by a shift of the median particle size from 2.1 (ink) to 2.1 (CV treatment) and 2.0 nm (1.4 V hold). More pronounced aggregation is visible after extensive aging, which pushes the mean particle size to  $2.8 \pm 0.8$  nm (2.6 nm median). Overall, the 0.6-nm mean nanoparticle size shift between the ink and extensive aging remains relatively mild in acid conditions.

In contrast, pronounced degradation is visible at pH 12, as the CV treatment yields a mean particle size of  $3.4 \pm 3.0$  nm (2.3 nm median size). Notably, the size distribution after the CV treatment contains  $\sim$ 7% of nanoparticles in the 0.9–1.1 nm range, which are not present in fractions above 1% in either the as-received nanoparticle ink or that after degradation in acid. Additionally, particle agglomerates as large as 20 nm are present. These agglomerates appear to be responsible for skewing the mean particle size (3.4 nm) away from the median size (2.3 nm). The large particles are visible in the micrographs in Figure S3, and the full-range particle size distribution is shown in Figure S4. Both these large agglomerates and the smaller 0.9-1.1 nm particles are also present after the hold at 1.4 V vs RHE for which a mean particle size of 3.8  $\pm$  4.0 nm and a median size of 2.1 nm were obtained. Finally, extensive aging generates a broad size distribution with a mean size of  $5.5 \pm 4.9 \text{ nm}$  (3.4 nm median).

The above results indicate that Pt/C degradation in pH 12 is more severe than that in pH 2. This is explicitly apparent when overlaying the particle size distributions after an identical treatment at low and high pH levels (Figure S5). The presently observed base-accelerated degradation is consistent with the literature; acidic electrolytes promote relatively mild nano-

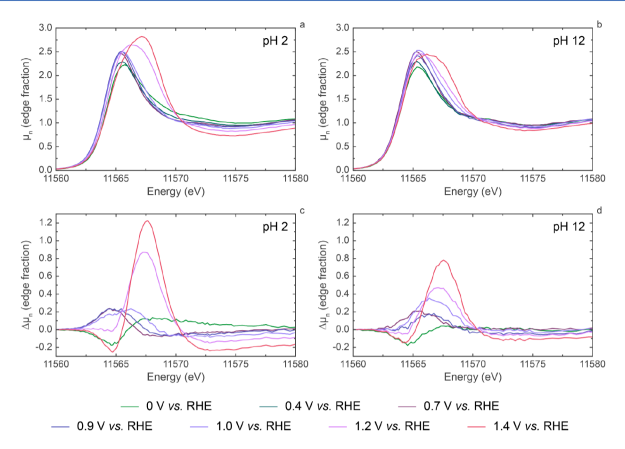


Figure 4. (a and b) Pt  $L_{\rm III}$ -edge HERFD-XANES spectra and (c and d) difference spectra of Pt/C nanoparticles at various potential holds in (a and c) pH 2 and(b and d) pH 12 electrolytes. In order to construct difference spectra for a given sample, the *in situ* spectrum of the same sample at 0.4 V vs RHE was used as a reference spectrum.

particle growth,<sup>29,54</sup> while alkaline electrolytes induce severe growth that is dominated by particle aggregation.<sup>29</sup> Overall, the significant pH-dependent differences warrant the further spectroscopic investigation of the studied Pt/C catalysts.

In Situ HERFD-XANES. In-situ HERFD-XAS spectra of Pt/C in pH 2 and pH 12 are presented in panels a and b, respectively, of Figure 4. The same figure also contains difference spectra for pH 2 and pH 12 in panels c and d, respectively. These difference spectra were obtained by subtracting the spectrum at 0.4 V vs RHE in panels a and b from the other spectra in the respective panels. This difference-based procedure highlights small spectral changes and can be used to identify adsorbates on Pt (electro)catalysts. A5,56-59 More specifically, the shape of a difference spectrum indicates the type of adsorbate, whereas the magnitude of a difference spectra indicates the relative abundances of these adsorbates. Including difference spectra in Figure 4 therefore helps highlight subtle chemical changes in the Pt/C catalysts.

In pH 2, the Pt/C catalyst is in its metallic state at 0.4 V vs RHE; it is not covered by hydrogen, but a finite amount of adsorbed \*OH is likely present at the top of Pt step edges and at corner sites. <sup>40,41,53,60</sup> This \*OH is substituted by \*H when the electrode is stepped down to 0 V vs RHE. Such \*H adsorption broadens the HERFD-XANES spectrum in Figure 4a, thus producing corresponding features in the difference spectrum (Figure 4c). *i.e.*, a negative feature around 11564.7 eV and a positive shoulder that gradually decreases in magnitude above 11567.1 eV. These changes in the normal and difference spectra match those in previous work <sup>53,61</sup> and

correspond to \*H that adsorbs in the "hollow" position, <sup>56</sup> as \*H adsorption creates antibonding electronic states that broaden the XANES spectrum. <sup>62–64</sup> Importantly, these hydrogen-induced changes are also visible at pH 12 (Figure 4b and d); however, they are approximately fourfold smaller in magnitude than those in pH 2. The decreased magnitude in the base is consistent with the TEM-observed particle growth at pH 12 after the CV cleaning treatment (Figure 3b), as these larger particles have a lower surface-to-bulk ratio than the smaller particles. Because of the reduced amount of surface atoms at pH 12, less Pt atoms are available for hydrogen adsorption and, consequently, smaller hydrogen features were observed for pH 12 in Figure 4 via HERFD-XANES.

Analogously to the HERFD-XANES features caused by the cathodically adsorbed \*H, features corresponding to the anodically adsorbed oxygen species are similar in shape but different in magnitude in the acid and the base. In both electrolytes, the main absorption peak (the whiteline) increases in intensity when the electrode potential is stepped from 0.4 to 0.7 V vs RHE. This increase yields a peak in the difference spectrum at 11564.5 eV, which signifies \*OH adsorption. 45,53,55,57 This spectroscopic fingerprint of \*OH confirms that the reductive current after oxidation at 0.7 V vs RHE in Figure 1 corresponds to Pt-OH reduction. Signs of additional adsorbates appear when the working electrode is polarized at 0.9 V vs RHE and above, as the intensity of the whiteline increases and its position gradually shifts. These changes in the spectra match those of the difference spectra, where a negative peak at 11564.5 eV and a positive peak at 11567.6 eV gradually

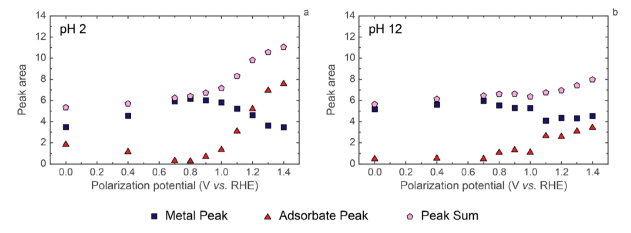


Figure 5. Areas of the fitted metal peak, the adsorbate peak, and the sum of both peaks. Full fit results are given in the Supporting Information.

develop. The positive difference peak corresponds to multifoldbound oxygen<sup>57</sup> and therefore suggests platinum oxide formation. The platinum oxide feature in the difference spectrum is more pronounced for pH 12 than for pH 2 when the difference spectra at 0.9 V vs RHE are compared. This suggests an earlier onset for Pt oxide formation in the base. The latter observation is supported by earlier reports of dissolved Pt with online ICP-MS<sup>27</sup> and with early onset oxidation currents during voltammetry in a base. 33 Oxidized Pt that does not dissolve remains on the surface as Pt oxides (PtO<sub>x</sub>). The identities of these oxides appear to be similar between the acid and the base, which is indicated by the difference spectra in these electrolytes having similar shapes.<sup>55</sup> At potentials above 1.4 V vs RHE, the most predominant oxide phase is likely PtO<sub>2</sub> in both the acid and the base; the shape of the difference spectra in Figure 4 matches that of a PtO<sub>2</sub>·H<sub>2</sub>O standard compound with a negative peak at 11564.4 eV and a positive peak at 11567.6 eV (Figure S6). Interestingly, in absolute terms the amount of PtO2 formed is largest in the acid because the difference spectrum magnitude at 1.4 V vs RHE is larger in the acid than in the base.

The degree of Pt oxidation is explicitly explored in Figure 5, which utilizes a HERFD-XANES fitting procedure to approximate the relative amount of metallic and oxidized Pt. S3,61 Full fitting details are provided in Supporting Information Figure S7. Briefly, a simplified model assumes a pseudo-Voigt peak at 11564.50 eV that is used to quantify the amount of metallic Pt, whereas a pseudo-Voigt peak at 11567.55 eV accounts for the amount of adsorbate-covered Pt. The adsorbate peak corresponds to \*H-covered Pt at potentials below 0.4 V vs RHE, \*O(H)-covered Pt, and Pt oxides at higher potentials. The sum of the adsorbate peak and the metallic peak is also displayed in Figure 5. The peak sum correlates with the amount of empty Pt d-states and is therefore a measure of Pt oxidation, where a higher peak sum indicates more oxidation.

As Figure 5 demonstrates, the metal peak, the adsorbate peak, and the peak sum vary as a function of the electrode potential. When lowering the electrode potential from 0.4 to 0 V vs RHE, the adsorbate peak decreases, and the metal peak increases. As mentioned in the previous paragraph, these changes correspond to \*H adsorption. <sup>53</sup> Additionally, the peak sum decreases slightly when moving toward 0 V vs RHE, which suggests a more reduced Pt state upon hydrogen adsorption. These changes in the adsorbate and metal peaks

are 3–4 times more pronounced at pH 2 than at pH 12. Analogously, the peak integrals at higher potentials vary more at low pH values than at high pH values; the adsorbate and sum peaks are higher in the acid than in the base, whereas the metal peak is lower. These changes indicate that a larger relative amount of Pt is oxidized at low pH values. However, as is also suggested by Figure 4, oxidation commences at lower potentials in the base, as the adsorbate peak starts growing at 0.8 V vs RHE in the base and at 0.9 V vs RHE in the acid. In short, the onset of Pt oxidation appears at ~100 mV less-positive potentials in the base, but a larger fraction of Pt atoms ultimately oxidizes in the acid at the highest potential.

# DISCUSSION

**Degradation Mechanisms.** The HERFD-XANES results (Figures 4 and 5) indicate that less Pt/C oxidation occurs in the base than in the acid. The reduced amount of oxidation in the base is caused by a larger nanoparticle size at a high pH. The HERFD-XANES results are therefore consistent with the complementary TEM results (Figure 3), which reveal that markedly more nanoparticle degradation occurs in the base. Notably, *in situ* spectroscopy captures minute chemical changes on the surface of Pt nanoparticles, showing the early formation of oxides in the base. Ultimately, both XAS and TEM demonstrate base-enhanced Pt/C degradation.

In discussing base-enhanced electrode degradation within the context of nanoparticle catalysts, primary and secondary degradation are distinguished. Briefly, primary degradation refers to the chemical mechanisms that cause aging, such as platinum dissolution and platinum-catalyzed carbon corrosion. In turn, these primary mechanisms drive secondary processes that change the nanoparticle morphology, such as Ostwald ripening, particle migration or agglomeration (Smoluchowski agglomeration), and nanoparticle detachment. Typically, the detection of such secondary changes can indicate which primary mechanism is predominant in the system.

Between the primary mechanisms, carbon corrosion has been suggested to cause enhanced Pt/C degradation in base.<sup>26</sup> Such corrosion is Pt-catalyzed<sup>48</sup> and is hypothesized to be accelerated by hydroxide ions in base,<sup>26</sup> as carbon corrosion could degrade the metal—support interactions and form carbon dioxide.<sup>48,67</sup> In alkaline solutions, this carbon dioxide would precipitate as carbonate salts. Within this suggested mechanism, the physical stress caused by the formation of solid carbonates and the weakened Pt—support bonds would cause

the detachment of Pt nanoparticles.<sup>26</sup> Once detached, Pt particles can diffuse into the working electrolyte, thus decreasing the nanoparticle density and the overall Pt loading on the support.<sup>29,30</sup> Alternatively, they could undergo Smoluchowski agglomeration by migrating within the support structure and colliding with other Pt particles.<sup>68</sup> Smoluchowski agglomeration generally causes the nanoparticle size distribution to tail toward large agglomerated particles.<sup>69,70</sup>

Smoluchowski agglomeration could also be caused by Pt dissolution and redeposition because Pt atoms dissolve from one side of a nanoparticle and are redeposited on a different side, thus causing a net migration of the particle. <sup>71</sup> In addition, Pt dissolution and redeposition drives Ostwald ripening. <sup>68</sup> Although the effects of Ostwald ripening on particle size distributions vary, <sup>72,73</sup> there is one prominent Ostwald ripening feature: the presence of nanoparticles that are smaller than those in the initial nanoparticle ink. <sup>74</sup> These particles form because Ostwald ripening preferentially consumes small particles, which will shrink to sizes that are not present in the initial nanoparticle size distribution. The presence of these minuscule particles is therefore an indicator of Ostwald ripening and, by extension, Pt dissolution. <sup>74</sup>

Based on the outlined degradation processes, one can set out to distinguish Pt dissolution and carbon corrosion. Of the secondary processes, Smoluchowski agglomeration is the most prominent secondary change in the present work. A pronounced agglomeration-induced tail is visible in the particle size distributions for both pH 2 and pH 12 in Figure 3, with exacerbated tailing visible in pH 12. Similarly, large particle agglomerates are visible for the base-aged samples in Figures S3 and S4. Notably, Smoluchowski agglomeration can be caused by both Pt dissolution and carbon corrosion. When distinguishing these primary processes, one should therefore rely on Ostwald ripening and nanoparticle detachment, which respectively indicate Pt dissolution/redeposition and carbon corrosion.

The present work reveals a reduced nanoparticle density for the alkaline samples in the micrographs, as seen in Figure S3. The loss of particles points toward carbon corrosion, which is consistent with previous degradation reports. <sup>29,30</sup> Simultaneously, TEM reveals the presence of particles with sub-1.2 nm diameters after moderate aging at pH 12 (Figure 3b). Such small nanoparticles are not present in the initial nanoparticle ink and therefore indicate Ostwald ripening. In turn, Ostwald ripening points toward degradation through Pt dissolution/redeposition. As such, the present TEM results support the simultaneous occurrence of Pt dissolution and carbon corrosion, which are enhanced in the base.

**Base-Enhanced Pt-Based Degradation.** The reasons for this alkaline enhancement are actively being researched for both carbon corrosion and Pt dissolution. Although the carbon corrosion mechanism is beyond the scope of this work, the present results do allow for conclusions on Pt-based degradation. Because such degradation inherently involves both the dissolution and redeposition of Pt, each of these components will be discussed separately.

Pt dissolution will be pH-dependent if the stability of the oxidized Pt species is also pH-dependent. Consequentially, different types or varying amounts of oxidized Pt species would form at different pH values. From an electrochemical perspective, there is only a  $\sim\!20\%$  difference between oxidation and reduction charge densities between electrolytes. Importantly, this difference favors oxidation in acid and thus does not

explain base-enhanced aging, which is also supported by the higher degree of oxidation in the acid above 1.0 V vs RHE. Analogously, HERFD-XANES reveals an earlier oxidation onset in the base than in the acid (Figures 4 and 5, respectively), but does not support a pronounced speciation difference in Pt oxidation at higher potentials, *i.e.*, a PtO<sub>2</sub>-like oxide forms at both pH 2 and pH 12 at potentials higher than 1.2–1.4 V vs RHE. This similarity between the acid and the base is supported by the similar shapes of the difference spectra in Figure 4c and d, which match the shape of hydrated PtO<sub>2</sub> (Figure S6).

With Pt dissolution seeming similar in both the acid and the base, it appears that Pt redeposition differs between acidic and alkaline Pt/C degradation. 24,25,28 This hypothesis relies on alkaline electrolytes containing hydroxide ions, which form complexes with dissolved Pt. As a result, the dissolved Pt is stabilized in base and is more likely to diffuse prior to redeposition.<sup>24,25</sup> Such solvated Pt is generally lost when bulk Pt electrodes are involved, 24 but carbon-supported Pt electrodes allow for diffusion within the 3D carbon support. 52,68,75 In these 3D systems, diffusion and subsequent redeposition will cause enhanced Ostwald ripening and Smoluchowski agglomeration, respectively. These effects are consistent with those observed in the present work. Similar results are obtained if Pt/ C is cycled in acidic solutions, but only if complexing ions such as chloride are present.<sup>76</sup> In the present work, only the highpH systems contain complexing ions in the form of OH-. These complexing ions stabilize the dissolved Pt species at pH 12, which causes base-enhanced diffusion of Pt species and subsequent catalyst degradation. This diffusion should be enhanced if more OH is present, which has indeed been shown when Pt dissolution is compared in 0.02 and 0.2 M KOH.

The aforementioned diffusion-promoting role of OH<sup>-</sup> is consistent with recent work that compares Pt/C degradation in a liquid 0.1 M NaOH electrolyte and in a dry solid-polymer electrolyte.<sup>30</sup> This comparison demonstrated reduced aging in the solid electrolyte, which was ascribed to the reduced mobility of dissolved Pt species prior to redeposition. Likewise, Pt loss from fuel cell catalysts can be reduced by impeding the mass transport of dissolved Pt species.<sup>52</sup> Such mass transport effects can generate different nanoparticle growth regimes throughout thick catalyst layers.<sup>75</sup> Although such spatial inhomogeneities cannot be assessed in the present study, our current results explicitly underscore the importance of the diffusion of Ptn+ that occurs between dissolution and redeposition. It is therefore crucial to look beyond the oxidation and reduction behavior of Pt when assessing fuel cell catalyst stability, as the diffusion of dissolved species appears to be equally relevant. Furthermore, this relevance could even vary with operational parameters such as the temperature, which is known to affect Pt catalyst degradation. Additionally, the type of complexing anions will likely determine the type of Pt complex that forms in solution. Consequently, the stability of the dissolved Pt will depend on the type of complexing anion. Determining the exact type of the dissolved Pt species would be a notable future research goal if solution-based spectroscopy or online chromatography and mass spectrometry techniques are optimized to distinguish the minuscule amounts of dissolved Pt.

Analysis of Aging Effects. The previous sections underscore the simultaneous occurrence of carbon corrosion and platinum dissolution while subsequently discussing base-

enhanced Pt dissolution in light of the results presented here and in recent literature. Having done this, a final discussion will be devoted to generally summarizing Pt/C aging in the present work

As discussed in the Degradation Mechanisms subsection, Pt/C aging in this work occurs through a combination of Ostwald ripening, Smoluchowski agglomeration, and Pt detachment. Although signs for all of these mechanisms are present, it appears that Smoluchowski agglomeration eventually dominates the aging process. This dominance is relatively apparent at pH 12, where nanoparticle agglomerates are already visible after a simple cyclic voltammetry procedure (Figure 3b). Agglomerates larger than 5 nm are relatively rare in pH 2, but a positively skewed particle size distribution in Figure 3a also indicates the prevailing Smoluchowski agglomeration after extensive aging.<sup>70</sup>

A final scrutiny of the size distributions in Figure 3 sheds light on the relative importance of steady-state and transient dissolution. As mentioned in the Introduction, steady-state dissolution occurs during constant anodic polarization, whereas transient dissolution takes place during oxidation—reduction cycles.<sup>21</sup> In the case of platinum, transient dissolution generally causes more Pt loss than steady-state dissolution. 21,27 This behavior is parallel to alkaline aging in this work since the most dramatic Pt/C growth occurs after the CV treatment (Figure 3b). However, this behavior is unmatched by Pt/C at pH 2. As reported previously, 54 only marginal Pt/C growth is detectable after both the CV treatment and a subsequent 1.4 V hold, whereas clear particle growth is only visible in the acid after extensive aging (Figure 3a). The lack of severe Pt/C aging after cycling might be caused by passivation effects for the small nanoparticle sizes in this work<sup>7,51</sup> and deserves further attention in future Pt/C aging studies.

# CONCLUSIONS

Summarizing, this work has demonstrated pH-dependent Pt/C aging during both cyclic voltammetry and constant anodic polarization. The results show that more Pt/C degradation occurs in base than in acid, which is consistent with previous reports. 24,26,28-30 The details of base-accelerated degradation were rigorously probed through a combination of electrochemistry, ex situ TEM, and in situ HERFD-XANES. Our analysis indicates that catalyst aging is a result of a combination of Ostwald ripening, Smoluchowski agglomeration, and Pt nanoparticle detachment. In turn, these secondary processes suggest an interplay between both carbon corrosion and Pt dissolution/redeposition as primary degradation mechanisms. Electrochemistry revealed no signs of different oxidation mechanisms between the acid and the base, as both electrolytes produced similar Pt oxidation and reduction charges. Furthermore, HERFD-XANES indicated the generation of a PtO2 phase at both low and high pH levels at potentials larger than 1.2 V vs RHE. Interestingly, in situ XAS suggests a ~100 mV earlier onset for the formation of oxides of Pt in base. Finally, it seems that base-induced Pt complexation facilitates the diffusion of dissolved Pt in base. This increased stability and mobility of solution-phase Pt would contribute to the base-enhanced Ostwald ripening and Smoluchowski agglomeration upon redeposition. Importantly, this conclusion indicates that similar fundamental Pt oxidation and reduction processes occur in the acid and the base, but the pHdependent chemistry of dissolved Pt is the cause of the baseenhanced Pt degradation through dissolution and redeposition.

Notably, these insights on the coexistence of Pt-based degradation and carbon corrosion would not have been reached without combining *ex situ* and *in situ* catalyst characterizations and shed valuable additional light on the relatively understudied issue of base-accelerated Pt catalyst degradation. The results additionally show that electrochemistry can conceal pH-dependent differences in Pt/C degradation, and *in situ* characterizations are critical for further advances. Such an improved understanding is required to implement Pt-catalyzed alkaline electrochemical reactions in general and alkaline fuel cells specifically. The present insights therefore mark relevant progress toward achieving a functional hydrogen economy.

## ASSOCIATED CONTENT

# Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscatal.1c02468.

Representative transmission electron micrographs, additional particle size distributions, comparison of difference spectra with those of standard compounds, and HERFD-XANES fit details (PDF)

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# **Author Contributions**

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#### Notes

The authors declare no competing financial interest.

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