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High-resolution photoelectron spectroscopy of TiO$_3$H$_2^-$: Probing the TiO$_2^-$ + H$_2$O dissociative adduct

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Slow electron velocity-map imaging spectroscopy of cryogenically cooled TiO$_3$H$_2^-$ anions is used to probe the simplest titania/water reaction, TiO$_2^{3/-}$ + H$_2$O. The resultant spectra show vibrationally resolved structure assigned to detachment from the cis-dihydroxide TiO(OH)$_2^-$ geometry based on density functional theory calculations, demonstrating that for the reaction of the anionic TiO$_2^-$ monomer with a single water molecule, the dissociative adduct (where the water is split) is energetically preferred over a molecularly adsorbed geometry. This work represents a significant improvement in resolution over previous measurements, yielding an electron affinity of 1.2529(4) eV as well as several vibrational frequencies for neutral TiO(OH)$_2$. The energy resolution of the current results combined with photoelectron angular distributions reveals Herzberg-Teller coupling-induced transitions to Franck-Condon forbidden vibrational levels of the neutral ground state. A comparison to the previously measured spectrum of bare TiO$_2^-$ indicates that reaction with water stabilizes neutral TiO$_2$ more than the anion, providing insight into the fundamental chemical interactions between titania and water. Published by AIP Publishing. https://doi.org/10.1063/1.5018414

I. INTRODUCTION

Titania (TiO$_2$) is an inexpensive, extensively studied, and environmentally benign semiconducting material with widespread applications in photovoltaics, pollution management, chemical sensing, and heterogeneous catalysis. The landmark discovery of photosensitization of water on a TiO$_2$ electrode sparked a decades-long pursuit to harness the photocatalytic properties of TiO$_2$ as a practical means of solar-powered hydrogen fuel production. However, the success of this endeavor has been limited in part by a lack of the mechanistic understanding necessary to design better catalysts. Here, we present high-resolution photoelectron spectra of the TiO$_3$H$_2^-$ anion accompanied by theoretical analysis, in order to probe the nature of the interaction between TiO$_2^-$ and a single water molecule.

The most natural starting point for understanding water oxidation by TiO$_2$ is a consideration of how water adheres to bulk titania surfaces, an active and complex area of research. The extent to which water dissociates on a TiO$_2$ surface is known to be dependent on surface structure, with varying propensities for dissociative versus molecular adsorption for different crystal phases and planes as well as a dependence on the extent of surface coverage by water. While dissociation tends to play a minor role in water adsorption on stoichiometric TiO$_2$ surfaces, it is found to be strongly preferred at point defects such as steps, edges, and in particular, oxygen vacancies. Thus, the conceptual key to understanding the surface chemistry of water on titania from a molecular level is the chemistry that occurs at these defect sites.

It is challenging to design a bulk experiment that is uniquely sensitive to the chemistry occurring at a specific surface defect. Defect sites typically make up a small fraction of total surface area and are difficult to reproducibly generate. Gas phase metal oxide clusters have been shown to be useful model systems for gaining mechanistic insight into complex catalytic processes, as these species show structural motifs such as dangling moieties and undercoordinated atoms that mimic the geometries at common defect sites. The control afforded by gas-phase experiments provides the ability to systematically manipulate reactivity-related factors such as particle size, charge, and stoichiometry. Fast-flow laser-ablation ion sources allow the production of both bare and reacted clusters, enabling the characterization of reactants, products, and potentially intermediates or transition states of model catalytic reactions using gas-phase spectroscopic techniques. These species have the added benefit of being computationally tractable, enabling experimentalists and theorists to develop a clear, molecular-scale understanding of catalytic reaction mechanisms which is difficult to obtain from bulk experiments alone.

The experimental spectroscopic characterization of bare (TiO$_2$)$_n$ clusters constitutes a growing body of work, including contributions from our laboratory. Less work has been done to probe species formed from the reaction of small (TiO$_2$)$_n$ clusters with a discrete number of water molecules,
which can either adsorb molecularly or dissociatively as on the bulk surface. The hydration of cationic TiO\(^+\) by up to 60 water molecules has been studied by mass spectrometry, though this measurement does not provide insight into the structure of the resultant clusters.\(^{50}\) These solvated cations have been structurally characterized using infrared action spectroscopy by Zheng and co-workers,\(^{51}\) who have also performed anion photoelectron spectroscopy (PES) on the anionic (TiO\(_2\))\(^-\)(H\(_2\)O)\(_{0-7}\) clusters.\(^{52}\) In both cases, their spectra indicated dissociative adsorption of water to form hydroxide species. Weichman and co-workers\(^{53}\) have recently used infrared action spectroscopy to systematically characterize the anionic (TiO\(_2\))\(^-\)(D\(_2\)O)\(_m\) clusters for \(n = 2-4\) and \(m = 1-3\), finding that the dissociative geometries are preferred for these clusters as well.

A number of groups have used theoretical treatments—most commonly density functional theory (DFT)—to assess the extent to which H\(_2\)O molecules dissociate on small (TiO\(_2\))\(_m\) clusters.\(^{54-56}\) The most comprehensive work in this area has been carried out by Dixon and co-workers,\(^{57,58}\) who used a hybrid genetic algorithm to determine the lowest-energy structures for (TiO\(_2\))\(_n\)(H\(_2\)O)\(_m\) \((n = 1-4, m = 1-2n)\) clusters.\(^{59}\) Geometry optimizations and single point calculations were carried out using both DFT and coupled cluster methods, yielding similar results for all model chemistries used. For the simplest stoichiometric TiO\(_2\)/water reaction, TiO\(_2\) + H\(_2\)O, they found that the di-hydroxyl TiO(OH)\(_2\) structure, in which the water is split, is preferred over molecular adsorption (TiO\(_2\)-H\(_2\)O) by over 40 kcal mol\(^{-1}\). Of the two dissociatively adsorbed structures reported, the planar C\(_2\)v cis-hydroxyl isomer was found to be more stable than the C\(_2\)v trans-OH isomer, though the energy difference was less than 3 kcal mol\(^{-1}\).

The only experimental work on this system comes from the aforementioned anion PES study by Zheng and co-workers,\(^{52}\) who reported the photodetachment spectrum for TiO\(_2\)H\(_2\)\(^-\) formed by a laser ablation reactor source. This spectrum showed a single broad electronic band spanning \(\sim 1\) eV in electron binding energy (eBE), reflecting extended unresolved vibrational progressions. From this spectrum, they estimated an electron affinity (EA) of 1.15 \(\pm\) 0.8 eV and a vertical detachment energy (VDE) of 1.51 \(\pm\) 0.08 eV. By comparison to their photoelectron spectrum of bare TiO\(_2\)\(^-\) and DFT calculations, the anion geometry was assigned to the TiO(OH)\(_2\) dissociative adduct. Other than the general assignment based on the position of the detachment feature, little information regarding the structure of the TiO\(_2\)H\(_2\)\(^-\) species can be gleaned from this work.

Slow electron velocity-map imaging of cryogenically cooled anions (cryo-SEVI) is a high-resolution variation of traditional anion PES which provides vibrationally resolved detachment spectra reflecting the geometric and vibronic structure of the anion and neutral. Previously, we have used cryo-SEVI to characterize the unreacted TiO\(_2\)\(^-\) monomer.\(^{48}\) Here, we probe the TiO\(_2\)H\(_2\)\(^-\) species, corresponding to a single TiO\(_2\)\(^-\) reacted with one water molecule. These spectra represent a significant improvement in resolution compared to the work of Zheng and co-workers, showing an extensive vibrational structure with typical peak widths of 10 cm\(^{-1}\) full-width at half-maximum (fwhm). With the assistance of DFT calculations, we assign the anion structure to the dissociative di-hydroxide TiO(OH)\(_2\)\(^-\) isomer identified as the lowest-energy neutral geometry by Dixon and co-workers. Several vibrational frequencies for the neutral TiO(OH)\(_2\) species are extracted, as well as its EA. A comparison to the unreacted TiO\(_2\)\(^-\) cryo-SEVI spectra elucidates the energetic effect of charge on the TiO\(_2\)+ + H\(_2\)O reaction, highlighting the utility of small metal oxide clusters as tools for understanding catalytic reactions.

II. EXPERIMENTAL METHODS

The cryo-SEVI technique and apparatus have been previously described in detail.\(^{50-62}\) For the current work, modifications to the previously described cryo-SEVI laser ablation ion source\(^{62}\) were carried out to provide access to species formed by the reaction of gas phase clusters with reactant molecules of interest, introduced via a pulsed Series 9 General Valve. The design implemented is loosely based on the Smalley fast flow cluster reactor\(^{42}\) and is similar to that used by Jarrold and co-workers.\(^{53}\) As in previous experiments, the frequency-doubled output of a 20 Hz Nd:YAG laser (2-3 mJ/pulse) impinges on a rotating and translating titanium target, forming a plasma. A pulse of helium gas (150 psi backing pressure, pulse width 60 \(\mu\)s) from an Even-Lavie valve\(^{64}\) entrains the plasma and the resultant mixture condenses and cools to form charged and neutral molecules and clusters in an 80 mm long, 2 mm diameter growth channel. These then pass through a 1 mm slit and into a second channel (90 mm long, 2 mm diameter), where a second pulsed valve introduces helium bubbled through room temperature H\(_2\)O (15 psi backing pressure, pulse widths of ~160 \(\mu\)s). This second channel serves as a reactor channel in which the ablation products cool further, react with H\(_2\)O, and then expand into vacuum.

Anions formed in the source are directed through a radiofrequency (rf) hexapole ion guide and rf quadrupole mass filter, which are used to select the relevant mass window from the distribution of species formed in the ablation channels. This ion packet is then confined in a rf octupole ion trap formed by the reaction of gas phase clusters with reactant molecules (TiO\(_2\))\(_m\)/(H\(_2\)O)\(_n\) clusters for \(n = 2-4\) and \(m = 1-3\) mass spectrometer.\(^{54-56}\) The mass-separated ions are focused into the interaction region of a velocity-map imaging (VMI) electrostatic lens assembly.\(^{67,68}\) The ions of interest are photodetached by a pulse of vertically polarized light. Here, the pump laser is 98 kHz, which should contain some contribution from 50TiO\(_2\)\(^-\) in addition to the target 48TiO\(_2\)H\(_2\)\(^-\) species. However, given that the electron affinity of TiO\(_2\) is quite high (4.2 eV)\(^{48}\) relative to the detachment energies used in this work (<2 eV), this species does not contribute to the spectra.

The detachment laser configuration is based on a tunable dye laser pumped by the second harmonic of a 20 Hz Nd:YAG laser. For photon energies above \(\sim\)1.3 eV (10 500 cm\(^{-1}\), 950 nm), the output of the dye laser is used without further modification. For photon energies below 1.3 eV, the output of
the dye laser is focused into a 63-cm long Raman cell containing 400 psi of H₂. The Q(1) (1) line in the H₂ Raman spectrum is used to red-shift the dye light by 4155 cm⁻¹, 69 producing tunable photon energies ranging from ~1.0 to 1.3 eV (8000-10 500 cm⁻¹, 1200-950 nm). Residual dye light as well as anti-Stokes shifted light is separated from the Stokes-shifted light by a long-pass dichroic mirror (900 nm cutoff). The Raman-shifted light is very slightly depolarized, 70 and thus is passed through a linear polarizer to ensure vertical polarization in the interaction region.

Photoelectrons ejected in the interaction region of the VMI assembly are projected onto a 2D detector comprising two chevron-stacked microchannel plates coupled to a phosphor screen, which is photographed using a CCD camera. 71 Each photo is analyzed for individual electron events, the centroids of which are calculated and binned in a 2200 × 2200 grid. 72 Slight deviations from circularity in the accumulated images are corrected using the circularization algorithm described by Gascooke and co-workers. 73 The electron velocity distribution is then reconstructed from the circularized images using the Maximum Entropy Velocity Legendre Reconstruction (MEVELER) algorithm. 74 The radial position features in the reconstructed image is related to electron kinetic energy (eKE) by acquiring VMI images for detachment from atomic Ni⁺ and O⁺ at several photon energies. 75,76

In addition to the eKE distributions, the MEVELER reconstruction of VMI images provides the photoelectron angular distribution (PAD) associated with each detachment transition, given by 77

\[
\frac{d\sigma}{d\Omega} = \frac{\sigma_{\text{tot}}}{4\pi} \left[ 1 + \beta P_2(\cos \theta) \right],
\]

where \(\sigma_{\text{tot}}\) is the total detachment cross section, \(P_2(x)\) is the second-order Legendre polynomial, \(\theta\) is the angle of the outgoing photoelectron velocity vector with respect to the laser polarization axis, and \(\beta\) is the anisotropy parameter. The anisotropy parameter, which ranges from −1 (perpendicular detachment) to +2 (parallel detachment), reflects the angular momenta of the detached electrons and thus provides information regarding the electronic character of each transition. 78

Due to the roughly constant resolving power, AeKE/eKE, of the VMI spectrometer, 57,68 the best eKE resolution is obtained for slow electrons. Thus, a SEVI spectrum is acquired by first taking an overview spectrum at a relatively high photon energy, then tuning the detachment laser to energies slightly above features of interest. This results in a collection of narrow, high-resolution spectral windows that are then scaled to match overview intensities, which are less sensitive to the variation of photodetachment cross section with photon energy. The high-resolution windows are spliced together to yield a complete photoelectron spectrum with sub-meV resolution. Spectra are plotted as a function of electron binding energy (eBE), given by eBE = hν − eKE.

III. COMPUTATIONAL METHODS

Possible minimum-energy structures for both anionic and neutral TiO₂ + H₂O were explored using a variety of DFT-based model chemistries. The preliminary results suggested meaningful differences between model chemistries. Therefore, initial benchmarking calculations were carried out to compare six DFT-based model chemistries with the results of coupled-cluster singles and doubles (CCSD) calculations, 79,80 as summarized in Sec. S1 of the supplementary material. Based on comparisons of these CCSD results to DFT-predicted energetic ordering of candidate anion and neutral minimum energy structures as well as vertical and adiabatic detachment energies (ADEs) (Tables S1 and S2 in the supplementary material), the Becke-93 exchange and Yang-Parr-Lee correlation functionals (B3LYP) were employed with the Def2TZVP basis set 81-85 and this model chemistry has been employed for the calculations described below.

Both doublet and quartet states were considered for anion candidates; singlet (closed-shell and open-shell) and triplet states were considered for the neutral candidates. For all identified structures, doublet anions are more stable than quartet species and the closed-shell singlet state was found to be more stable than the open-shell singlet or triplet state. Excited state calculations were carried out using the same model chemistry within the time-dependent DFT (TDDFT) formalism. 86-88 Analysis of these excited state calculations was facilitated by Martin’s Natural Transition Orbital (NTO) model. 89

All calculations were carried out using a local development version of the Gaussian suite of electronic structure programs. 90 Converged Kohn-Sham determinants were tested for stability. 91,92 Molecular geometries were optimized using standard methods 93 and the reported potential energy minima were verified using analytical second-derivative calculations. Franck-Condon (FC) spectra were simulated using the implementation by Blinov, Barone, and co-workers. 94,95 Care was taken to determine appropriate scaling factors for the DFT force constants calculated for the lowest-energy neutral state (Table S3 of the supplementary material) such that simulated FC progressions aligned well with the experimental spectra. Further details of the FC simulations can be found in Sec. S2 of the supplementary material.

To describe the nature of the detached electron we have employed the Natural Ionization Orbital (NIO) model of Thompson, Harb, and Hratchian. 96 The NIO model provides a compact orbital representation of ionization processes by utilizing one-particle difference densities. Natural orbital analysis involving this difference density yields a simplified interpretation of electronic detachment processes. The NIO model has recently been shown to provide a convenient means to distinguish between one-electron transitions and those where the one-electron process is accompanied by excitation of a second electron into the virtual orbital space. 97 The current system is Koopmans-like, evidenced by the strong resemblance between the NIO and the canonical highest-occupied molecular orbital (HOMO) of the 1-1a anion (Fig. S1 of the supplementary material), and thus the detachment transitions considered here are one-electron transitions that can be equivalently described by considering the NIO or the anion HOMO. As such, the method of Liu and Ning 98 was applied to the HOMO of the lowest-energy anion to calculate the eKE-dependent PAD expected for the removal of an electron from this orbital; the results are shown as the solid lines in Fig. 2 and discussed further in Sec. V A.
IV. EXPERIMENTAL RESULTS

The cryo-SEVI spectrum for detachment from TiO$_2$H$_7^-$ is shown in Fig. 1. In this figure, the blue trace corresponds to an overview scan taken with a relatively high detachment energy and the black traces are higher resolution SEVI scans taken with variable photon energies. The overview spectrum spans 10 000-13 500 cm$^{-1}$ in eBE and exhibits considerable vibrational structure, revealing increasing spectral congestion as the eBE increases. Due to this increased complexity, our analysis will be focused on the first ~2000 cm$^{-1}$ of structure. In this region, the high-resolution scans reveal a number of transitions (A1-11 and B1-11) with typical peak widths of ~10 cm$^{-1}$ fwhm, corresponding to detachment to different vibronic levels of the neutral TiO$_2$H$_2$ species. The sharp onset of structure at peak A1 gives an EA of 1.2529(4) eV for TiO$_2$H$_2$. The remainder of the spectrum is dominated by a ~675 cm$^{-1}$ progression (A1-3-7-11), modulated by several weaker patterns. Peak positions, widths, and shifts from the origin are summarized in Table I.

The EA provided by cryo-SEVI is larger than the value of 1.15(8) eV reported by Zheng and co-workers. This discrepancy is understandable given the lower resolution of their experiment. The remainder of the spectrum is dominated by weaker patterns. Peak positions, widths, and shifts from the origin are summarized in Table I. The cryo-SEVI spectrum for detachment from TiO$_2$H$_7^-$ shows the Franck-Condon simulation for detachment with an anisotropy parameter (+ or 0), and vibrational assignments for features in the cryo-SEVI spectrum of TiO$_2$H$_7^-$.

TABLE I. Peak positions, shifts from the vibrational origin, qualitative anisotropy parameters (+ or 0), and vibrational assignments for features in the cryo-SEVI spectrum of TiO$_2$H$_7^-$.

<table>
<thead>
<tr>
<th>Peak</th>
<th>eBE (cm$^{-1}$)</th>
<th>Shift (cm$^{-1}$)</th>
<th>Assn.</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>10 106(3)</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td>B1</td>
<td>10 165(3)</td>
<td>60</td>
<td>0</td>
</tr>
<tr>
<td>A2</td>
<td>10 519(2)</td>
<td>413</td>
<td>+</td>
</tr>
<tr>
<td>B2</td>
<td>10 576(2)</td>
<td>471</td>
<td>0</td>
</tr>
<tr>
<td>A3</td>
<td>10 784(3)</td>
<td>678</td>
<td>+</td>
</tr>
<tr>
<td>B3</td>
<td>10 846(3)</td>
<td>741</td>
<td>0</td>
</tr>
<tr>
<td>A4</td>
<td>10 926(6)</td>
<td>820</td>
<td>+</td>
</tr>
<tr>
<td>B4</td>
<td>10 987(7)</td>
<td>881</td>
<td>0</td>
</tr>
<tr>
<td>A5</td>
<td>11 121(5)</td>
<td>1016</td>
<td>+</td>
</tr>
<tr>
<td>A6</td>
<td>11 199(4)</td>
<td>1093</td>
<td>+</td>
</tr>
<tr>
<td>B6</td>
<td>11 258(4)</td>
<td>1152</td>
<td>0</td>
</tr>
<tr>
<td>A7</td>
<td>11 459(3)</td>
<td>1354</td>
<td>+</td>
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<tr>
<td>B7</td>
<td>11 524(3)</td>
<td>1418</td>
<td>0</td>
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<tr>
<td>A8</td>
<td>11 607(13)</td>
<td>1501</td>
<td>+</td>
</tr>
<tr>
<td>B8</td>
<td>11 673(12)</td>
<td>1568</td>
<td>0</td>
</tr>
<tr>
<td>A9</td>
<td>11 798(5)</td>
<td>1692</td>
<td>+</td>
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<tr>
<td>A10</td>
<td>11 874(6)</td>
<td>1769</td>
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<td>B10</td>
<td>11 936(4)</td>
<td>1831</td>
<td>0</td>
</tr>
<tr>
<td>A11</td>
<td>12 133(4)</td>
<td>2027</td>
<td>+</td>
</tr>
<tr>
<td>B11</td>
<td>12 200(6)</td>
<td>2095</td>
<td>0</td>
</tr>
</tbody>
</table>

Measurement of the PADs of features labeled in Fig. 1 (Fig. S2 of the supplementary material) reveals that all transitions fall into one of two groups with distinct angular distributions. These two groups are represented in Fig. 2, which plots the anisotropy parameter (β) versus eKE for peaks A1-3 and B1-3. Features A1-3 show positive values of β, whereas features B1-3 show isotropic (β ≈ 0) angular distributions. The remainder of features in Fig. 1 shows PADs that are qualitatively similar to one of these two groups, and are labeled accordingly, where peaks labeled A correspond to parallel (β > 0) detachment transitions and peaks labeled B have isotropic (β ≈ 0) PADs. These anisotropy parameters are summarized qualitatively in Table I. Each B-series transition lies ~60 cm$^{-1}$ above a peak in the A1-11 series, and has been numbered accordingly.

These two groups also show distinctly different dependencies of the photodetachment cross section on eKE. Far from threshold, the A1-11 series dominates the spectrum, as can be seen in the overview spectrum in Fig. 1. As the photon energy is lowered, transitions in the B1-11 series become more important.
apparent, resulting from the relative attenuation of the A1-11 peaks. This trend is illustrated in Fig. 3, where peaks A1 and B1 are shown for several photon energies and the intensities are normalized to the peak intensity of B1. The relative scaling of detachment cross sections for low-eKE detachment is given by the Wigner threshold law:

$$\sigma \propto (eKE)^{l+1/2},$$  \hspace{1cm} (2)$$

where $\sigma$ is the detachment cross section and $l$ is the angular momentum of the detached electron. According to this law, a sharper decrease in detachment signal as the photon energy is lowered reflects higher $l$ detachment channels; it can therefore be inferred that transitions A1-11 correspond to higher $l$ detachment than transitions B1-11.

V. DISCUSSION

A. Structural assignment of TiO$_3$H$_2^-$

Figure 4 shows optimized minimum-energy structures located on the anionic and neutral TiO$_3$H$_2$ poten-tial energy surfaces, as well as zero-point corrected energies. Cartesian coordinates for these structures are provided in Secs. S3 and S4 of the supplementary material. In agreement with previous work by Dixon and co-workers, the $C_2v$ dissociative structure (1-1a') is found to be the lowest-energy neutral geometry; likewise, the 1-1a geometry is the lowest-energy anion isomer. Two additional minimum-energy structures, 1-1b and 1-1e, were found on the anion potential energy surface to lie 0.08 and 0.19 eV above 1-1a, respectively, and differ from this isomer by rotation of the hydroxide groups. A local minimum corresponding to the trans-OH 1-1b' geometry was discovered at an energy of 0.11 eV above the neutral 1-1a' structure, in reasonable agreement with the relative energy reported by Dixon. No neutral minimum corresponding to the 1-1e isomer was identified. The lowest-energy molecularly adsorbed TiO$_3$H$_2$ adduct reported by Dixon was also identified as a local minimum for both anion (1-1c) and neutral (1-1c'), though these structures lie 2.34 and 3.13 eV above the anion and neutral global minima, respectively.

Given the relative energies of the anion geometries shown in Fig. 4, the most likely structures contributing to the cryo-SEVI spectrum of TiO$_3$H$_2^-$ correspond to the dissociative geometries 1-1a and 1-1b. The assignment of a dissociatively adsorbed anion geometry is consistent with the assessment of Zheng and co-workers.$^{32}$ Predicted adiabatic detachment energies (ADEs) for 1-1a and 1-1b are 1.23 and 1.26 eV, respectively, both in good agreement with the experimental electron affinity (1.25 eV). In both cases, NIO analysis clearly indicates that the detached electron resides in the anion HOMO, which strongly resembles the Ti d$_{3z^2}$ orbital (see Fig. 5 and Fig. S3 of the supplementary material). Analysis of the NIO for either 1-1a or 1-1b gives an orbital angular momentum of $L = 2$, such that detachment from either of these orbitals would be expected to primarily yield outgoing $p$- ($l = 1$) and $f$-wave ($l = 3$) electrons. This high-$l$ detachment is consistent with the observed attenuation of the signal near threshold for peaks A1-11.

Additionally, the eKE-dependence of the anisotropy parameter calculated$^{98}$ for detachment from the 1-1a HOMO (Fig. 2, solid lines) is in good agreement with the measured $\beta$ values for peaks A1-11, showing positive values in the eKE region of interest. Given the similarity between the 1-1a and 1-1b NIOs, detachment from the 1-1b isomer is expected to have a similar PAD. The agreement between experiment and theory further supports the conclusion that the global minimum of the TiO$_3$H$_2^-$ anion takes the dissociative TiO(\OH)$_2^-$ dihydroxide geometry and that detachment from one of these isomers to the ground state of the corresponding neutral accounts for the most intense structure (A1-11).

To determine definitively which of these species is responsible for the experimentally observed detachment transitions, we consider the Franck-Condon profiles for detachment from both isomers, as the FC profile is highly sensitive to the anion and neutral geometries. Figures S4 and S5 of the supplementary material show the experimental overview spectrum overlaid with FC simulations for detachment from the ground electronic state of the 1-1a and 1-1b anion, respectively, to the closed-shell singlet state of the corresponding neutral. To provide a better comparison to experiments, Figs. S6 and S7 of the supplementary material show FC profiles calculated using neutral vibrational frequencies that have been scaled as described in Table S3 of the supplementary material. Of these two
isomeric candidates, the FC profile for detachment from the \( \tilde{X}^2A_1 \) state of the 1-1a anion (Figs. S4 and S6 of the supplementary material) provides better agreement with experiment than detachment from the \( \tilde{X}^2A' \) 1-1b anion (Figs. S5 and S7 of the supplementary material). The calculation in Fig. S6 of the supplementary material using scaled frequencies is shown in red in Fig. 1 and reproduces the dominant vibrational structure observed in the experimental spectrum. In particular, the first several intense features (A1-3) are reproduced in the FC simulation for 1-1a, whereas the 1-1b simulation shows extra structure between A1 and A3. Thus, we assign the 1-1a cis-OH geometry as the lowest-energy isomer of TiO\(_3\)H\(_2\), and the observed spectral features reflect the vibronic structure of the corresponding 1-1a' neutral.

**B. Vibrational assignments**

The agreement between the simulated and experimental spectra in Fig. 1 enables straightforward assignment of the major structure (A1-11) as \( \tilde{X}^1A_1 \) \( \leftrightarrow \tilde{X}^1A_1 \) detachment transitions terminating in totally symmetric vibrational levels of the neutral ground state (Table I). The less intense B1-11 series of peaks are not fully described by this FC simulation, and the previously described PADS indicate that these transitions do not have the same electronic character as A1-11. As such, we will first address the vibrational assignments of A1-11.

The dominant progression (A1-3-7-11) is assigned to the \( v_3 \) vibrational mode, which corresponds to a symmetric Ti—OH stretching motion (Fig. S8 of the supplementary material). The shift of peak A3 from the vibrational origin (A1) gives a frequency of \( 678(5) \) cm\(^{-1} \) for this mode. There is also significant participation of the \( v_4 \) totally symmetric OH wagging mode in the spectrum (A2, A6, A10), as well as some activity in the \( v_2 \) terminal Ti=O stretch (A5, A9). The shifts of peaks A5 and A2 give neutral vibrational frequencies of \( 1016(8) \) and \( 413(3) \) cm\(^{-1} \) for \( v_2 \) and \( v_4 \), respectively. Table II summarizes the energetic quantities for neutral TiO(OH)\(_2\) extracted from experiment and compares them to the 1-1a' B3LYP/Def2TZVP results, showing reasonably good agreement for these totally symmetric modes.

The Franck-Condon activity of different normal modes can be rationalized by considering the geometrical changes that occur upon photodetachment, which are summarized for the B3LYP/Def2TZVP 1-1a \( \tilde{X}^2A_1 \) anion and 1-1a' \( \tilde{X}^1A_1 \) neutral equilibrium geometries in Table S4 of the supplementary material. The most significant change is a 13.6\(^\circ \) increase in the H—O—Ti bond angle, which results in the FC activity of the \( v_4 \) OH-wagging mode. The involvement of the titanium-oxygen stretching modes \( v_2 \) and \( v_3 \) is a reflection of the change in bond lengths upon photodetachment for both the terminal Ti=O bond as well as the hydroxide Ti—O bonds. The hydroxide and terminal Ti—O bonds of the B3LYP geometries decrease by 0.08 and 0.05 Å, respectively, corresponding to modest 3%-4% decreases in these bond lengths. These constitute the

**TABLE II.** Electronic and vibrational energies for neutral 1-1a' TiO(OH)\(_2\) extracted from the TiO\(_3\)H\(_2\) cryo-SEVI spectrum, and compared to the (unscaled) results obtained from B3LYP/Def2TZVP calculations.

<table>
<thead>
<tr>
<th></th>
<th>Cryo-SEVI</th>
<th>B3LYP/Def2TZVP</th>
</tr>
</thead>
<tbody>
<tr>
<td>EA (eV)</td>
<td>1.2529(4)</td>
<td>1.2341</td>
</tr>
<tr>
<td>( v_2 ) (cm(^{-1} ))</td>
<td>1016(8)</td>
<td>1062</td>
</tr>
<tr>
<td>( v_3 ) (cm(^{-1} ))</td>
<td>678(5)</td>
<td>683</td>
</tr>
<tr>
<td>( v_4 ) (cm(^{-1} ))</td>
<td>413(3)</td>
<td>485</td>
</tr>
<tr>
<td>( v_8 ) (cm(^{-1} ))</td>
<td>60(4)</td>
<td>16</td>
</tr>
</tbody>
</table>
second- and third-largest fractional changes upon detachment for the five geometrical parameters summarized in Table S4 of the supplementary material.

While the above considerations fully assign peaks A1-11, the differing PADS and threshold behavior of peaks B1-11 indicate that these transitions have different electronic character, ruling out assignment to Franck-Condon allowed transitions within the $X^1A_1 \leftrightarrow X^2A_1$ electronic band including vibrational hot bands. These transitions are also apparently not equivalent to detachment from an excited anion state, as a TDDFT calculation for the anion (Table S5 of the supplementary material) does not reveal an excited state with sufficiently low excitation energy to be populated in the cold trap, where ions typically have temperatures on the order of 10 K. It is possible that peaks B1-11 arise from detachment to a separate electronic state of neutral TiO(OH)$_2$, and that the two groups of features correspond to overlapping electronic bands with the A1-B1 spacing providing the electronic term energy. However, this would require a neutral state with a term energy of around 60 cm$^{-1}$, and the present theoretical results do not provide evidence for such a state, with the lowest-energy neutral excited state of 1-1a lying 3.86 eV above the closed-shell singlet (Table S5 of the supplementary material).

Another possible assignment for peaks B1-11 is detachment from a different TiO$_2$H$_2^-$ geometry. In the theoretical framework, the 1-1b trans-OH TiO(OH)$_2^-$ dissociative adduct is expected to be only 0.08 eV higher than the cis-OH (1-1a) isomer. Given this low relative energy as well as its structural similarity to the global minimum, this geometry could interconvert with the slightly more stable 1-1a geometry in the ion trap, resulting in contributions from multiple isomers in the cryo-SEVI spectrum. However, the barrier for conversion of the 1-1a isomer to the 1-1b geometry is estimated to be $\sim$1300 cm$^{-1}$ from constrained optimization and single point calculations (Fig. S9 of the supplementary material), indicating that such interconversion is highly unlikely in the cold environment of the ion trap. Additionally, as discussed above, the NIOs for the 1-1a and 1-1b isomers are quite similar (Fig. 5 and Fig. S3 of the supplementary material), with both orbitals dominated by Ti $d_{z^2}$ character. Thus, if detachment features from the 1-1b geometry were observed, the transitions would not have PADS that deviate so significantly from the cis-OH detachment transitions, ruling out assignment of peaks B1-11 as detachment from the 1-1b anion. A similar argument can be made to rule out detachment from the 1-1c isomer, which also differs from 1-1a only by rotation of the hydroxyl groups.

Given the above considerations, and the observation that each of the B1-11 features lies $\sim$60 cm$^{-1}$ above a transition in the A1-11 series, we conclude that these features correspond to detachment transitions terminating in $X^1A_1$ vibrational levels with odd quanta of excitation along a low-frequency non-totally symmetric mode. These FC-forbidden transitions obtain their oscillator strength and PADS through Herzberg-Teller (HT) coupling to an excited neutral electronic state. The most likely vibrational assignment for peaks B1-11 involves the $b_1$-symmetric vibration of the Ti atom through the plane of the three oxygen atoms (Fig. S8 of the supplementary material) and has the lowest calculated frequency (Table S3 of the supplementary material) for the 1-1a' neutral. Each Bi transition in the B1-11 series is thus assigned as terminating in the state corresponding to the A1 transition, plus a single quantum of excitation along $v_8$. With this assignment, the position of peak B1 relative to A1 gives a vibrational frequency of 60(4) cm$^{-1}$ for the $v_8$ umbrella mode of neutral TiO(OH)$_2$.

We now consider the symmetry requirements for the excited electronic state which gives rise to transitions B1-11 through Herzberg-Teller coupling. Consider a vibronic state $|a\rangle$ whose electronic and vibrational symmetries are $\Gamma^a_{\text{elec}}$ and $\Gamma^a_{\text{vib}}$, respectively. This state can undergo HT-coupling with another vibronic state, $|b\rangle$, with symmetries $\Gamma^b_{\text{elec}}$ and $\Gamma^b_{\text{vib}}$, provided

$$\Gamma^a_{\text{elec}} \otimes \Gamma^a_{\text{vib}} \otimes \Gamma^b_{\text{elec}} \otimes \Gamma^b_{\text{vib}} \supset \Gamma^\text{TS},$$

where $\Gamma^\text{TS}$ is the totally symmetric representation within the relevant molecular point group. If state $b$ is FC-allowed for detachment from the anion in question (i.e., $b | \Psi^\text{anion} \rangle \neq 0$), detachment to state $a$ will reflect the $b$-state electronic character, which will be observable in the PAD and threshold behavior.

In the current case, the observed HT-coupled levels correspond to states with odd quanta of excitation along the $b_1$-symmetric $v_8$ mode ($\Gamma^b_{\text{elec}} = b_1$) within the vibrational manifold of the ground neutral electronic state ($\Gamma^a_{\text{elec}} = A_1$). Within the $C_2v$ point group, these levels are Franck-Condon forbidden, and so can only be observed if they mix with some vibronic level $b$ that is not FC-forbidden, i.e., $\Gamma^a_{\text{vib}} = a_1$. Thus, the observed features must arise from HT-coupling with a $B_1$-symmetric electronic excited state. The lowest such singlet state identified from TDDFT (Table S5 of the supplementary material) is the relatively high-lying $\tilde{C}^1B_1$ excited state, residing 4.45 eV above neutral $\tilde{X}^1A_1$ 1-1a'. Using the NTO model, we determined that this state would involve detachment from an orbital with angular momentum $L = 1$, resulting in outgoing $s$- ($l = 0$) and $d$-wave ($l = 2$) electrons, in contrast to the $p$- and $f$-wave detachment expected from the 1-1a NIO. Thus, the assignment of peaks B1-11 as arising from Herzberg-Teller coupling to this $^1B_1$ state is consistent with the observed relative attenuation of A1-11 versus B1-11 as photon energy is lowered.

C. Charge effects on the TiO$_2$ + H$_2$O reaction

The cryo-SEVI spectrum of unreacted TiO$_2^-$ has been previously reported, giving an electron affinity of 1.5892(5) eV for the singlet ground state of TiO$_2$. The electron affinity of the TiO(OH)$_2$ dissociative adduct is lower by roughly 0.3 eV, and this difference provides insight into the energetic effects of charge on the dissociation of H$_2$O by TiO$_2$. The lower electron affinity for the reacted species implies that the neutral TiO$_2 +$ H$_2$O $\rightarrow$ TiO(OH)$_2$ reaction is more exothermic than the anionic counterpart—that is, reaction with water to form the dissociative TiO(OH)$_2$ adduct stabilizes neutral TiO$_2$ more than it does the anion. As a consequence, it can be inferred that neutral TiO$_2$, where the titanium center has a +4 oxidation state, is more reactive towards water than anionic TiO$_2^-$ where the Ti oxidation state is +3. The energies shown
The lowered electron affinity of TiO$_2$H$_2$ relative to TiO$_2$ indicates that neutral TiO$_2$ is stabilized to a greater extent than anionic TiO$_2^-$ by reaction with water, suggesting that the reaction of TiO$_2$ with water is impeded by excess negative charges. This conclusion is consistent with the existing electrochemical understanding of photocatalytic water splitting by TiO$_2$ in photoelectrochemical cells, where the sensitization of water arises from the ability of TiO$_2$ to accept electron density. This work highlights the relevance of gas-phase studies of metal oxide cluster reactions to bulk catalytic reactions, as well as the utility of cryo-SEVI as a structural probe for such systems.

SUPPLEMENTARY MATERIAL

See supplementary material for details regarding electronic structure and Franck-Condon calculations as well as Figs. S1–S9 and Tables S1–S5.

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VI. CONCLUSION

Vibrationally-resolved photoelectron spectra of the TiO$_2$H$_2^-$ anion are obtained using slow electron velocity-map imaging of cryogenically cooled anions, yielding spectra that reflect the vibronic structure of neutral TiO$_3$H$_2$. These results provide a significant improvement in resolution over prior work on this system, clearly resolving the onset of the structure at the adiabatic detachment energy. A Franck-Condon simulation for detachment from the C$_{2v}$ 1-1a TiO(OH)$_2^-$ cts-hydroxide geometry captures the dominant vibronic structure observed experimentally, enabling the assignment of this geometry as the lowest-energy anion isomer in agreement with density functional theory calculations. In addition to transitions reproduced in the FC simulations, a series of features resulting from Herzberg-Teller coupling to an excited neutral electronic state is identified in the cryo-SEVI spectrum, evidenced by noticeable differences in the angular and energy dependence of the photodetachment cross sections.

The comparison of the current spectral results to the previously reported cryo-SEVI spectrum of unreacted TiO$_2^-$ provides insight into the reactivity of TiO$_2^-$ towards water.


100 E. P. Wigner, Phys. Rev. 73, 1002 (1948).