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H₂ Oxidation over Supported Au Nanoparticle Catalysts: Evidence for Heterolytic H₂ Activation at the Metal–Support Interface

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Supporting Information

ABSTRACT: Water adsorbed at the metal–support interface (MSI) plays an important role in multiple reactions. Due to its importance in CO preferential oxidation (PrOx), we examined H₂ oxidation kinetics in the presence of water over Au/TiO₂ and Au/Al₂O₃ catalysts, reaching the following mechanistic conclusions: (i) O₂ activation follows a similar mechanism to that proposed in CO oxidation catalysis; (ii) weakly adsorbed H₂O is a strong reaction inhibitor; (iii) fast H₂ activation occurs at the MSI, and (iv) H₂ activation kinetics are inconsistent with traditional dissociative H₂ chemisorption on metals. Density functional theory (DFT) calculations using a supported Au nanorod model suggest H₂ activation proceeds through a heterolytic dissociation mechanism, resulting in a formal hydride residing on the Au and a proton bound to a surface TiOH group. This potential mechanism was supported by infrared spectroscopy experiments during H₂ adsorption on a deuterated Au/TiO₂ surface, which showed rapid H−D scrambling with surface hydroxyl groups. DFT calculations suggest that the reaction proceeds largely through proton-mediated pathways and that typical Brønsted−Evans Polanyi behavior is broken by introducing weak acid/base sites at the MSI. The kinetics data were successfully reinterpreted in the context of the heterolytic H₂ activation mechanism, tying together the experimental and computational evidence and rationalizing the observed inhibition by physisorbed water on the support as blocking the MSI sites required for heterolytic H₂ activation. In addition to providing evidence for this unusual H₂ activation mechanism, these results offer additional insight into why water dramatically improves CO PrOx catalysis over Au.

INTRODUCTION

The global chemical industry produces over 50 million tons of hydrogen for several important processes including ammonia and methanol synthesis, petroleum refining, and hydrogenation reactions.1,2 Industrial H₂ production (predominantly by methane steam reforming and water−gas shift units) results in H₂ feeds containing about 1% CO. Many downstream uses of H₂, particularly ammonia synthesis catalysts and fuel cells, are highly sensitive to CO, so it must be removed. The scale of hydrogen production and the potential for preparing fuel cell grade hydrogen make hydrogen purification an enormously impactful process: ammonia production in particular accounts for 3% of total global energy consumption.1 Methanation (CO + 3H₂ → CH₄ + H₂O) and pressure swing adsorption are currently used to purify H₂, but each method has its limitations.2

Another option for hydrogen purification is the preferential oxidation of CO with O₂ (PrOx reaction). In PrOx, a small amount of O₂ (typically ~1%) is added to the feed; the goal is to find catalysts that can oxidize all of the CO without oxidizing any H₂. A typical benchmark goal for this reaction is to reduce the CO concentration at the reactor outlet to 50 ppm with O₂ selectivity to CO₂ ≥ 50%.4−6 This places enormous selectivity demands on the catalyst, which must oxidize CO ~10⁶ times faster than H₂. Supported Au nanoparticle catalysts are notoriously slow hydrogenation catalysts,7 yet are highly active for CO oxidation;8−13 thus, they should be well-suited for the CO PrOx reaction.

Several research groups have investigated CO PrOx over Au in the past two decades,5,14−21 with mechanistic studies performed by the Behm,5,14−17 and Piccolo and Rousset groups18,19 as well as computational investigations by Mavrikakis and co-workers.20,21 In most cases, the presence of H₂ was found to increase CO oxidation activity. Water plays an important role as a co-catalyst in CO oxidation,13,22−29 so these observations are consistent with the in-situ production of water. Early studies by Behm showed water to improve CO PrOx performance by suppressing H₂ oxidation and at least partially prevent carbonate poisoning.14

We recently showed that PrOx performance can be dramatically improved by orders of magnitude when the surface coverage of physisorbed water is controlled.30 This improvement was greater than expected based on weakly adsorbed water’s role as a co-catalyst in CO oxidation,13,31 in order to better understand the performance-enhancing ability
of water, we undertook a more detailed examination of effects of weakly adsorbed water on the undesirable half of the PrOx reaction: \( \text{H}_2 \) oxidation. Our reaction kinetics study, carried out at low conversions, indicates that \( \text{H}_2 \) coverage is rate limiting and that physisorbed water is a strong inhibitor for the reaction. Both kinetics data and density functional theory (DFT) calculations indicate that \( \text{H}_2 \) is selectively activated at the metal support interface (MSI) and suggest that the water inhibition is primarily due to the physical blocking of these MSI sites.

The reaction kinetics, DFT results, and supporting infrared spectroscopic characterization of \( \text{H}_2 \) activation on a \( \text{D}_2 \text{O} \)-exchanged catalyst indicate that \( \text{H}_2 \) oxidation occurs at the MSI through a heterolytic \( \text{H}_2 \) activation pathway. This is a surprising discovery. Hydrogen adsorption and activation is one of the most studied reactions in all of chemistry; early investigations into the interactions between hydrogen and various metals date back to the mid-19th century.\(^{32-34}\)

Hydrogenation reactions over heterogeneous catalysts are widely used in industry and have been studied for over a century.\(^{35-38}\) Similarly, the mechanism of hydrogen activation by inorganic complexes, dating back to seminal work by Wilkinson and Vaska, has been widely studied for more than 50 years.\(^{39}\) The overwhelming majority of these studies show that hydrogen activation occurs via similar mechanisms: oxidative addition for transition metal complexes\(^{40,41}\) (and homolytic) dissociative chemisorption for metals and supported metal catalysts.\(^{42,43}\)

There are exceedingly few reports of heterolytic \( \text{H}_2 \) activation by heterogeneous catalysts; Coperet’s work on \( \text{Al}_2 \text{O}_3 \) defect sites provides several notable examples.\(^{44-46}\) and Boudart proposed such a mechanism at paramagnetic centers of \( \text{MgO} \).\(^{47}\) Examples of heterolytic \( \text{H}_2 \) activation are well-known in biological systems, particularly hydrogenase enzymes and their synthetic models.\(^{48,49}\) Recent studies on frustrated Lewis pairs (FLPs) show similar \( \text{H}_2 \) reaction pathways.\(^{50-53}\)

The studies reported herein provide strong evidence that supported metal nanoparticle catalysts can also operate via similar Lewis acid–base mechanisms, highlighting the mechanistic similarities between biological, homogeneous, and heterogeneous catalysts.

### EXPERIMENTAL SECTION

**Materials.** Gases (\( \text{H}_2, \text{N}_2, \text{O}_2, 20 \text{ vol \% } \text{O}_2/\text{He}, \) and 5 vol \% \( \text{CO}/\text{He} \)) were 5.0 grade supplied by Praxair and used without further purification. Water was purified to a resistivity of 18.2 MΩ with a Barnstead Nanopure System; no additional purification methods were employed. Commercial catalysts were purchased from STREM Chemicals. The catalysts have been fully characterized elsewhere;\(^{54} \) the basis for the \( \text{Au}/\text{TiO}_2 \) work on \( \text{Au}/\text{TiO}_2 \) defect sites provides several notable examples.\(^{44-46} \) and Boudart proposed such a mechanism at paramagnetic centers of \( \text{MgO} \).\(^{47} \) Examples of heterolytic \( \text{H}_2 \) activation are well-known in biological systems, particularly hydrogenase enzymes and their synthetic models.\(^{48,49} \)

**Reaction System.** The \( \text{H}_2 \) oxidation reaction consisted of a home-built laboratory-scale single-pass plug-flow microreactor operated at atmospheric pressure (760 Torr). Gas flows were controlled with four electronic low-pressure mass flow controllers (Porter Instruments). Water was added to the feed using a two-stage water saturator after the reactant gases were mixed; feedwater pressure was determined by adjusting the temperature of the second stage. The composition of the feed and reactor effluent (\( \text{CO}_2, \text{CO}, \) and \( \text{O}_2 \)) was determined using a Siemens Ultramat 23 IR gas analyzer with electrochemical \( \text{O}_2 \) sensor. The feed concentration was determined via a reactor bypass loop. The reaction zone consisted of finely ground fresh catalyst (5–100 mg) diluted in 1 g of \( \text{SiC} \). The catalyst powder was mixed thoroughly with the \( \text{SiC} \) and finely chopped using a spatula until homogeneous.

Immediately prior to kinetics experiments, the diluted catalyst was pretreated in a mixture of 10 vol \% \( \text{H}_2, 10 \text{ vol \% } \text{O}_2, \) and balance \( \text{N}_2 \) at 100 °C for 1 h. This treatment was employed to ensure a consistent degree of surface hydroxylation on the catalyst and to remove impurities (e.g., surface organics, carbonates). The reactor was then cooled to the reaction temperature under flowing gas consisting of 19 Torr of \( \text{H}_2 \text{O}/\text{N}_2 \). The system was allowed to equilibrate for 30 min at constant reactor and water saturator temperature whenever the \( \text{H}_2 \text{O} \) pressure was changed. The catalyst water coverage was calculated from volumetric \( \text{H}_2 \text{O} \) adsorption isotherms.\(^{50} \)

**H2 Oxidation Kinetics.** Conversions were measured 5 min after steady state was achieved by collecting gas composition data every 10 s for 2 min. Steady state was defined as \( \text{O}_2 \) slip being constant with a range of 0.02 vol \% \( \text{O}_2 \) over 5 min. During kinetic experiments, conversions were held below 15% in order to maintain differential reaction conditions and keep \( \text{H}_2 \text{O} \) generation low with respect to the added \( \text{H}_2 \text{O} \). Gases (3–60 vol \% \( \text{H}_2, 0.9–10 \text{ vol \% } \text{O}_2, \) with 5–18 Torr \( \text{H}_2 \text{O} \) added via saturator) were fed to the reactor at weight hourly space velocities (WHSV) of (0.2–3.2) \times 10^3 \text{ g} \cdot \text{g}_{\text{cat}}^{-1} \cdot \text{h}^{-1}. \) The reaction temperature was 60 °C. \( \text{H}_2 \text{O} \) reaction orders were measured with 60 vol \% \( \text{H}_2 \) and 1 vol \% \( \text{O}_2 \); reaction orders were measured with 60 vol \% \( \text{H}_2 \) at three different \( \text{H}_2 \text{O} \) pressures (6.8, 11, and 18 Torr). \( \text{H}_2 \) reaction orders were measured with 10 vol \% \( \text{O}_2 \) at four different \( \text{H}_2 \text{O} \) pressures (5, 6, 9, and 12 Torr).

**DFT Calculations.** Plane wave density functional theory (DFT) calculations with periodic boundary conditions were performed using the Vienna Ab Initio Simulation Package (VASP).\(^{53-55} \) Exchange and correlation were described with the BEEF-vDW functional,\(^{58} \) and the projector augmented wave (PAW) method was used to approximate the core electronic structure.\(^{59,60} \) Spin polarization was used wherever necessary, i.e., the adsorption and activation of \( \text{O}_2 \). A plane wave energy cutoff of 400 eV was used for all the calculations. The same energy cutoff of 400 eV was used previously for studying CO oxidation on \( \text{Au}/\text{TiO}_2 \).\(^{61} \) The gas phase \( \text{H}_2 \) and \( \text{H}_2 \text{O} \) energies were calculated in a \( 10 \times 10 \times 10 \text{ Å} \) simulation box, and Brillouin zone sampling was restricted to the \( \Gamma \) point. For phase specific studies, we employed a Gaussian smearing with \( k_T = 0.01 \text{ eV} \), and geometries were optimized using a force convergence criterion of 0.01 eV/Å.

For bulk and slab models, we employed Gaussian smearing with a Fermi temperature of \( k_T = 0.1 \text{ eV} \), and the total energy was extrapolated to \( k_T = 0.0 \text{ eV} \). Residual forces on equilibrium geometries were converged to below 0.05 eV/Å. The reaction energy for the bulk oxidation from \( \text{TiO}_2 \) to \( \text{TiO}_2 \) was reproduced within an error of 0.04 eV with this arrangement; consequently, implementation of the DFT+\( \text{U} \) approach by Dudarev et al. was not necessary.\(^{62,63} \) The computationally optimized lattice constants are \( a = 4.654 \text{ Å}, \) \( a / c = 1.561 \text{ for } \text{TiO}_2 \) and \( a = 4.223 \text{ Å} \) for \( \text{Au} \). These values agree well with experimentally observed lattice constants of \( a = 4.682 \text{ Å}, \) \( a / c = 1.574 \text{ for } \text{TiO}_2 \) and \( a = 4.08 \text{ Å} \) for \( \text{Au} \). For slab models, we used a \( 3 \times 2 \) \( \times 1 \text{ Monkhorst–Pack } k \)-point mesh to sample the Brillouin zone, and a dipole correction was applied to the electrostatic potential in the \( z \) direction.

Transition states were located using the climbing image nudged elastic band (NEB) method and refined as necessary with the dimer method with a convergence criterion of 0.1 eV/Å. All transition states were confirmed as true saddle points with a single imaginary frequency mode along the reaction coordinate. Vibrational frequencies were obtained using the atomic simulation environment (ASE) module in the harmonic oscillator approximation with a displacement of 0.01 Å along each positive and negative Cartesian direction. Atomic charges were estimated based on a Bader analysis.\(^{64-66} \)

**Au/\text{TiO}_2 \) Computational Model.** The basis for the \( \text{Au}/\text{TiO}_2 \) interface is formed by a rutile \( \text{TiO}_2(110) \) (\( 5 \times 3 \)) unit cell separated by 20 Å of vacuum space in the \( z \) direction perpendicular to the surface. The bottom two bilayers of \( \text{TiO}_2 \) were fixed in their bulk positions, while all other degrees of freedom were relaxed. We did not
consider oxygen vacancies on the TiO2 surface because the presence of significant amounts of O2 and H2O in the experimental feed is likely to heal or passivate surface defects quickly.69 Next, a three-layer gold nanorod was placed along the [110] direction of TiO2 with its (111) facet exposed at the interface. We refer to this model as Au(111)/TiO2(110). The lattice constant mismatch between Au and TiO2 was minimized with a nanorod length of seven Au atoms, leaving a residual compressive strain of 5.53% in the Au nanorod along the [110] direction of the TiO2 unit cell. A similar level of strain was reported by Henkelman et al. with the gold nanorod oriented in the [210] direction.70 Compressive strain is known to lower the d-band center of metals and, in turn, decrease their reactivity.70 We have quantified the effect of 5.53% compressive strain on H2 dissociation on Au(211) step sites in the Supporting Information (SI). These estimates indicate that compressive strain alters the activation energies by less than 0.07 eV and the dissociation energies by only 0.02 eV. We consider this error negligible within the context of our study, and our qualitative conclusions are robust with respect to the effect of strain.

The Au nanorod model on TiO2 used in this study has been improved from previous nanorod models11,61,71 to accommodate possible sites for H2 activation and allow for comparisons between reactions on the metal and at the MSI. The effect of surface hydroxylation was approximated by creating bridge-hydroxyl groups (bridge-OH) at all available bridging oxygens on the TiO2 surface and hydroxyl groups at coordinatively unsaturated (cus) Ti atoms (cus-OH). The hydroxylation state of the Au/TiO2 model shown in Figure 1 can also be thought of as a model having a monolayer equivalent (MLE) of dissociated water molecules on the exposed TiO2 surface. When comparing reaction energetics between Au sites away from the MSI to activity near or at the MSI, we refer to sites within the highlighted regions of Figure 1. The Au sites are composed of atoms in local (111), (100), and (211) geometries and have a coordination number (CN) of 7. The edge atoms at the MSI have a CN of 6+, where 6 strictly counts Au neighbors and “+” accounts for bonds made with the TiO2 support. As we demonstrate in the Results and Discussion section, the activation barrier for homolytic H2 activation at these AuMSI sites with CN = 6+ is in fact slightly higher than on the (211)-type Au sites with CN = 7. Thus, the small difference in coordination number alone cannot account for a large difference in reactivity when interfacial reactions are studied.

FTIR Spectroscopy. FTIR spectra during H2 adsorption were collected on a Thermo Nicolet Nexus 470 FTIR spectrometer in a heated (20–300 °C) transmission flow cell. H2O in the feed gases was removed by a dry ice–iso-propyl alcohol (IPA) moisture trap. For the exchange experiment, 30 mg of catalyst sample was pressed (5 tons of pressure for 1 min) in a 13 mm circular pellet, which was mounted in the flow cell. D2O (99.0%, Cambridge Isotope Laboratories) was flowed through the pellet for 30 min using a two-stage saturator. The complete deuteration of the support was monitored by collecting scans over the course of the treatment. The weakly absorbed D2O was removed by flowing N2 at 120 °C for 1 h before cooling to 70 °C. This ensured no weakly adsorbed D2O remained and only OD and strongly adsorbed D2O were present. H2 was then flowed over the catalyst at a WHSV of 40 L/gcat/h for 30 min, with scans collected every 5 min.

RESULTS AND DISCUSSION

Kinetic measurements on CO oxidation have been previously reported13,31; these studies showed that weakly adsorbed H2O plays a number of important mechanistic roles during CO oxidation. Protons from water help to activate O2 by generating Au-OOH, which quickly reacts with CO, yielding Au-COOH. Physisorbed water also plays a role in the rate-determining decomposition of Au-COOH acting as a proton acceptor as the catalyst releases CO2. During PrOx, the primary feed component is H2 and some (undesirable) H2 oxidation occurs.70 Note that Behm has demonstrated that CO and H2 compete for the same O intermediate in PrOx, so understanding H2 activation may help to understand and control this competition.16 Goodman’s work used inelastic neutron scattering experiments to characterize a reactive O species during H2 oxidation as an −OH species on Au.72 We therefore aim to understand water’s role in H2 oxidation for both fundamental and practical reasons.14

Reaction Kinetics. Figure 2 shows how added water affects H2 oxidation under typical PrOx conditions (1 vol % O2, 60 vol % H2, 60 °C, 50 mg of catalyst); the data clearly show that H2O inhibits H2 oxidation. Water is the reaction product, so under PrOx conditions, the water produced from the reaction impacts the reaction kinetics. The red diamonds in Figure 2A show the total amount of water in the system, including the water produced from H2 oxidation calculated from the O2 conversion data. Under these conditions, in-situ water production is greater than the H2O added to the system.

The strong inhibitory effect of water complicates these measurements because the total water pressure in the system (including the water produced by the reaction) must be held approximately constant for a given measurement. Total O2...
conversions were held below 15% to maintain differential reactor conditions; additionally, the amount of water generated from the reaction was small relative to the amount of water intentionally added (<5 vol % for high water pressures and <25 vol % at the lowest water pressure). This puts significant constraints on the reactor operating conditions, requiring water coverages of >0.5.

Figure 2 shows kinetic data for Au/Al₂O₃ and Au/TiO₂; the extracted H₂O reaction orders are −0.70 and −0.64, respectively. It is important to clarify that we previously identified weakly bound or physisorbed water as the mechanistically important proton donor in O₂ activation over both Au/TiO₂ and Au/Al₂O₃ catalysts. Any strongly bound water, e.g., to exposed Ti atoms, is unlikely to be sufficiently mobile to participate in the fast proton donation associated with O₂ activation (and Au-COOH decomposition in CO oxidation). Further, the water pressures and reaction temperatures used in the current study affect the amount of water physisorbed on the support hydroxyl groups. Therefore, equilibrium adsorption isotherm measurements for water on the two catalysts were used to estimate the surface coverage of weakly adsorbed water (θH₂O). While this is an imperfect comparison relative to the flow system, the errors introduced are partially compensated by the additional water produced from the reaction. Further, errors in the coverage estimates are inherently included in the overall errors associated with the rate measurements. Figure 2B shows the water dependence data plotted with respect to θH₂O. The fits are quite good, indicating that any errors in the estimation of θH₂O are not likely to have a significant influence on the results. The reaction orders with respect to θH₂O are about −1.5, further showing the strong inhibition of H₂ oxidation by weakly adsorbed water.

Figure 3 shows the H₂ pressure dependence data over a large range of H₂ pressures (1−60 vol %); Figure 4 shows H₂ order plots at different water pressures. The extracted reaction orders for H₂, O₂, and H₂O are compiled in Table 1 and compared to the kinetic data for CO oxidation. Hydrogen oxidation is approximately first order in H₂ suggesting that H₂ activation is a kinetically important step. As Table 1 shows, the key difference between CO and H₂ oxidation is the water dependence: weakly adsorbed water promotes CO oxidation but inhibits H₂ oxidation. Since weakly adsorbed water is required for fast O₂ activation, this suggests that the physisorbed water inhibits H₂ activation. Water does not adsorb to Au under these conditions, and the physisorbed water resides on the support. Therefore, if physisorbed water
inhb starts.

Traditional Models for Hydrogen Activation on Metals. The widely accepted mechanism for H₂ adsorption/activation on metals (Pt, Pd, Rh, Ru, Ni, etc.) involves H₂ dissociation on the metal surface, followed by reaction with an activated hydrogen species. This reaction is largely equivalent to the classical mechanism of oxidative addition on transition metal complexes. In both cases, H₂ dissociation formally oxidizes the nanoparticle by two electrons, and the adsorbed H atoms can be considered formal hydrides. This process is largely equivalent to the classic mechanism of oxidative addition on transition metal complexes. In both cases, H₂ dissociation formally oxidizes the nanoparticle by two electrons, and the adsorbed H atoms can be considered formal hydrides. This process is largely equivalent to the classic mechanism of oxidative addition on transition metal complexes.

Table 1. Summary of Reaction Orders during H₂ and CO Oxidation Catalysis

<table>
<thead>
<tr>
<th>Reactant</th>
<th>P_H₂O (Torr.)</th>
<th>H₂ Oxidation</th>
<th>CO Oxidation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Au/TiO₂</td>
<td>Au/Al₂O₃</td>
</tr>
<tr>
<td>H₂</td>
<td>12</td>
<td>0.60 ± 0.06</td>
<td>0.90 ± 0.10</td>
</tr>
<tr>
<td></td>
<td>9</td>
<td>0.58 ± 0.06</td>
<td>0.77 ± 0.09</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>0.70 ± 0.08</td>
<td>0.92 ± 0.07</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>0.66 ± 0.06</td>
<td>0.74 ± 0.05</td>
</tr>
<tr>
<td>CO</td>
<td>18.2</td>
<td>0.21 ± 0.01</td>
<td>0.37 ± 0.09</td>
</tr>
<tr>
<td>O₂</td>
<td>10.8</td>
<td>0.12 ± 0.01</td>
<td>0.29 ± 0.08</td>
</tr>
<tr>
<td></td>
<td>6.4</td>
<td>0.28 ± 0.07</td>
<td>0.40 ± 0.10</td>
</tr>
<tr>
<td>H₂O</td>
<td>---</td>
<td>-0.64 ± 0.02</td>
<td>-0.70 ± 0.03</td>
</tr>
</tbody>
</table>

**Scheme 1. Elementary Steps for H₂ Oxidation on Au via Homolytic H₂ Activation on Au Sites Away from the MSI**

- **MSI + O₂ + TiOH → OOH_MSI + TiO⁻** (S1-1)
- 2(Au + H₂O) + 2H₂O → 2Au + 4H₂O (S1-2)
- 2H₂O_MSI + 2H₂O + Au → H₂O_MSI + + Au (S1-3)
- 2H₂O_MSI + 2H₂O + Au → H₂O_MSI + + Au (S1-4)
- 2H₂O_MSI + 2H₂O + Au → H₂O_MSI + + Au (S1-5)
- 2H₂O_MSI + 2H₂O + Au → H₂O_MSI + + Au (S1-6)

Overall Reaction: 2H₂ + O₂ → 2H₂O

**MSI** sites are Au atoms at the metal support interface. Au sites are considered to be away from the MSI, and TiOH sites are support hydroxyl groups. Steps S1-2 and S1-3 are both considered to be kinetically important; all subsequent steps are considered fast.

Double-reciprocal (e.g., Lineweaver–Burk) plots, which, in this case, graph 1/rate vs 1/P₄, can be used to evaluate changes to heterogeneous catalysts. When coupled with appropriate kinetic models, double reciprocal plots can provide a useful framework for evaluating the viability of a reaction mechanism, the intrinsic activity at an active site, and a means of evaluating changes to the number of active sites on a catalyst. Hydrogen oxidation over Au is generally described as limited by hydrogen activation in the literature. Homolytic H₂ activation on the Au surface and subsequent reaction with Au-OOH at the MSI can be considered with a kinetic model similar to the H₂ oxidation mechanism on Pt proposed by Dumesic and co-workers. This mechanism involves hydrogen activation on the metal followed by reaction with an activated oxygen species (see Scheme 1). Applied the steady state approximation, which also assumes a constant coverage of Au-OOH at a particular water pressure, yields the following rate law in eq 2 (full derivation available in the SI):

$$ν_{homo} = \frac{k_1 k_{H₂} K_{H₂} P_{H₂} [Au]_T [MSI]_T}{k_3 K_{MSI} P_{MSI} [MSI]_T + k_2} (1 + K_{MSI} P_{MSI})$$

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where $k_2$ is the rate constant for step S1-2 and $k_3$ is the rate constant of step S1-3. Note that the derived rate law has a maximum $H_2$ dependence of 0.5; our kinetic data show this value to be 0.64 ± 0.05 for Au/TiO$_2$ and 0.83 ± 0.08 for Au/Al$_2$O$_3$. This rate law has the associated double-reciprocal form (eq 3):

$$\nu = \frac{1}{1 + \frac{k_{\text{homo}}[\text{Au}]}{k_2[H_2]}} + \frac{(1 + K_{\text{R}}[P_{\text{O}_2}])}{k_3K_{\text{O}_2}[\text{MSI}][\text{Au}]}$$

Two kinetic parameters, $v_{\text{max}}$ and $K_R$, can be extracted from eq 3 (eqs 4 and 5):

$$v_{\text{max}} = \frac{1}{\text{intercept}} = \frac{k_3K_{\text{O}_2}[\text{Au}][\text{MSI}][\text{Au}]}{1 + K_{\text{R}}[P_{\text{O}_2}]}$$

$$K_R = \frac{\text{slope}}{\text{intercept}} = \frac{k_3K_{\text{O}_2}[\text{P}_{\text{O}_2}]}{k_2[1 + K_{\text{R}}[P_{\text{O}_2}]]}$$

Figure 5 shows the $H_2$ dependence data (Figure 4) fitted to eq 3. While the data are reasonably linear, all of the lines have $y$-intercepts < 0, implying physically meaningless negative values for the extractable kinetic parameters ($v_{\text{max}}, K_R$). Thus, the traditional homolytic $H_2$ adsorption model, where the rate-determining step is either homolytic $H_2$ adsorption or the reaction of the adsorbed Au-H with an adsorbed O species, is inconsistent with the kinetic data. In light of this incomplete understanding of the reaction, alternative mechanistic interpretations must be considered. These mechanistic pathways should account for the following experimental observations from this study: (i) near first-order dependence on $H_2$ pressure; (ii) $O_2$ activation through the proton-mediated generation of Au-OOH, as indicated by the $O_2$ reaction order; (iii) strong inhibition by physisorbed water on the support, which implicates the MSI as an important reaction site.

Density Functional Theory Calculations. Given the inconsistencies between the traditional homolytic $H_2$ activation model and our kinetic measurements, we performed DFT calculations to better understand the interactions between $H_2$, water, and the catalyst. To this end, we used a sophisticated model of the Au/TiO$_2$ interface in order to capture the key features of the real system. Notably, the nanorod model shown in Figure 1 exposes two distinct hydroxyl groups at the MSI: a bridge hydroxyl, bridge-OH, and a terminal (cus = coordinatively undersaturated) hydroxyl, cus-OH. These surface hydroxyls, which have different acid/base properties, are necessary to represent the experimental conditions where water is present in the system. We note that we examine the limiting case of full hydroxylation, which is equivalent to dissociatively adsorbing 1 MLE of water onto the stoichiometrically terminated rutile TiO$_2$(110) surface. The high surface hydroxyl coverage destabilizes individual OH species and increases their reactivity, but affords the advantage of eliminating artificial charge imbalances and maintains the key electronic structure features of stoichiometric TiO$_2$. The SI contains a density of states analysis showing that hydroxylation induces minimal differences in the band gap and electronic states near the Fermi level that determine chemical reactivity. These important features set this model apart from previous interface models $^{99-102}$ and are shown below to improve our ability to model interface reactivity.

The initial exposure of water to stoichiometrically terminated TiO$_2$(110) leads to the strongly exothermic
formation of cus-OH and bridge-OH sites ($\Delta E = -1.01$ eV/H$_2$O). When additional water physisorbs to the hydroxylated support, it preferentially adsorbs on the bridge-OH at the Au/TiO$_2$ interface with a binding energy of $-0.53$ eV/H$_2$O. In comparison, physisorption of water near the bridge-OH sites away from the interface is exothermic by $-0.30$ eV/H$_2$O. Water adsorption onto the Au sites ($-0.19$ eV/H$_2$O) is weaker still. This binding preference is consistent with earlier work using a Au$_{10}$/TiO$_2$ nanocluster model, and the generally weaker binding on Au sites for the nanorod model (relative to the Au$_{10}$/TiO$_2$ model) can be attributed to the higher Au coordination number in the nanorod model.

From the experimental H$_2$ oxidation kinetics, we can infer that the reaction is limited by hydrogen availability on the surface and that O$_2$ is not involved in H$_2$ activation. We report here two possible mechanisms for H$_2$ activation on Au/TiO$_2$. The dominant homolytic H$_2$ dissociation pathway on nanorod Au–Au sites resembling a (211) step geometry (CN = 7) away from the MSI is thermodynamically unfavorable ($\Delta E = +0.59$ eV) and is associated with a large activation barrier ($E_a = 1.16$ eV, Figure 6A). The absence of a significant support effect is corroborated in Table S1, showing that control simulations on a Au(211) slab yield nearly identical values for $\Delta E$ and $E_a$.

Homolytic activation of H$_2$ at Au sites near the MSI, where the edge Au atoms are also coordinated to support Ti and O atoms (CN = 6+), is even less favorable ($\Delta E = 0.79$ eV, $E_a = 1.27$ eV). Our calculated values for the activation barrier for H$_2$ dissociation at the Au sites afforded by the nanorod model agrees with the value of 1.1 eV reported for a periodic Au(111) slab, but is significantly higher than the barrier of only 0.27 eV over an unsupported 12-atom cluster. These values indicate that homolytic H$_2$ activation on Au sites even along the interface is expected to be slow, unless highly under-coordinated sites are present. This interpretation is consistent with the widely reported observation that Au is a poorly active hydrogenation catalyst.

We also examined H–H bond activation across the MSI as depicted in Figure 6B and C; the reaction energies and associated activation barriers are summarized in Table 2. H$_2$ adsorption at the MSI is essentially thermoneutral ($\Delta E = -0.03$ eV), and the associated activation barrier ($E_a = 0.70$ eV) is almost half that of the barrier at Au step sites on the nanorod ($E_a = 1.16$ eV). Figure 6B also shows that H$_2$ activation at the MSI proceeds through more complicated interactions with both the metal and support cus-OH sites, resulting in heterolytic H–H bond cleavage, with a proton residing on the support hydroxyl and a formal hydride on the Au. We note that heterolytic H$_2$ dissociation involving a bridge-OH site (which has weak Brønsted acid character) has a moderate thermodynamic barrier ($\Delta E = 0.44$ eV) and a correspondingly higher activation barrier ($E_a = 1.15$ eV). Both of these values are comparable to the homolytic cleavage at (211)-type Au sites away from the MSI. Further, we note that H$_2$ activation on cus-OH is affected by hydroxyl stability, and the saturation of the TiO$_2$ surface with dissociated water increases the cus-OH reactivity. These effects are quantified in the SI and limit the maximum barrier for heterolytic H$_2$ activation on cus-OH to $E_a = 0.91$ eV across the MSI when the TiO$_2$ surface is incompletely hydroxylated.

Given the high pH$_{2}^*$ value for H$_2$ (36), the DFT prediction of a heterolytic H–H cleavage pathway is surprising, so we sought further insight into the potential driving forces for this reaction pathway. We examined the electron density redistribution for both heterolytic and homolytic H$_2$ activation using Bader charge analysis. H$_2$ adsorption on top of the cus-OH sites is athermal at Au sites even along the interface is expected to be slow, unless highly under-coordinated sites are present. This interpretation is consistent with the widely reported observation that Au is a poorly active hydrogenation catalyst.

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equal Bader charges for each H atom in the transition state (Figure 6A). This traditional homolytic H₂ adsorption pathway also shows little electron transfer from Au to the (formal) dihydrides. Gold is one of the few metals more electronegative than hydrogen, so the highly covalent nature of the Au–H bonding is not surprising; further, the difficulty of formally oxidizing Au is consistent with the thermodynamic unfavorability of this step.

The atomic charges from the Bader analysis are referenced to the initial states of H₂ dissociation. The transition state Bader analysis of the heterolytic MSI pathway (Figure 6B) shows a relatively early transition state with a +0.35 e⁻/Å charge on the developing proton. The developing hydride, however, has a slightly smaller negative charge, indicating that some of the negative charge is transferred to the Au nanorod. This is confirmed by the Bader analysis of the final state (Figure 6C), which shows ~0.14 e⁻/Å charge on the (formal) hydride, ~0.11 e⁻/Å charges distributed across the Au nanorod, a charge of ~0.50 e⁻/Å transferred to the oxygen in the cus-OH site, ~0.14 e⁻/Å charges distributed on Ti atoms, and the balance of ~0.11 e⁻/Å charges distributed over the remaining oxygen atoms.¹⁰⁶ The H₂ dissociation involving the bridge-OH site is also heterolytic in nature with a similar charge of +0.38 e⁻/Å on the developing proton in the transition state.

**Infrared Spectroscopy.** If the heterolytic H₂ adsorption pathway predicted by the DFT studies is correct, we should expect to see rapid exchange between gas phase hydrogen and the developing proton. To test this hypothesis, we used infrared spectroscopy to monitor H₂ adsorption on a D₂O-exchanged support protons. To test this hypothesis, we used infrared spectroscopy to monitor H₂ adsorption on a D₂O-exchanged Au/TiO₂ catalyst. Figure 7 shows the evolution of the O–H(D) region of the IR spectrum over time after the introduction of H₂; the D₂O exchanged catalyst was collected as a background spectrum to more easily observe the changes upon H₂ adsorption. The growth of the broad band at ~3300 cm⁻¹ (ν(O–H)) is consistent with the rapid addition of H-bonded protons upon introducing H₂ to the catalyst. The loss of the band at 2700 cm⁻¹, which is associated with non-H-bonded or “dangling” –OD bands similarly indicates that these –OD groups become involved in H-bonding interactions when H₂ is added to the system. The increase of the band at ~1600 cm⁻¹ (δ(H–O–H)) and 1400 cm⁻¹ (δ(D–O–D)) and the loss of the band at 1200 cm⁻¹ (δ(H–O–D)) is also consistent with proton exchange between H₂ and the deuterated support.

Heterolytic H₂ adsorption at the MSI sites requires the H–D exchange, so the observed changes in Figure 7 are consistent with this mechanism. Further, they exclude the possibility that H₂ adsorption occurs exclusively on the metal; if this were the case, no exchange with the support would be observed. The results in Figure 7 are also consistent with homolytic adsorption followed by fast exchange with the support. As discussed below, this is kinetically indistinguishable from heterolytic adsorption, although the DFT calculations suggest that heterolytic activation is the more likely pathway. Additionally, by using the molar absorptivity for weakly adsorbed water for FTIR,¹³ a rate for water formation can be extracted from Figure 7. This rate represents a lower limit for H₂ activation, considering this is not a direct measure of the activation itself. Rather, it measures the rate of water formation, which requires the formation of two O–H bonds on a single O atom and is an approach to the equilibrium distribution of all protons and deuterons in the system. The rate extracted from Figure 7 was 0.0113 ± 0.0001 mol·H₂O/molAu/s, which is generally consistent with the rates measured for H₂ oxidation.

**Hydrogen Spillover and Metal Support Proton Exchange.** The terms used to describe hydrogen spillover and proton exchange processes (generally revealed through H–D exchange experiments) are often used interchangeably in the literature. To clearly define these processes, we refer to Prins’ discussion in a recent review.¹⁰⁷ The term “spillover” was originally coined by Boudart and Vannice to describe the migration of H atoms from the metal particles to the support, because the H atoms spill over, as it were, from a hydrogen-rich to a hydrogen-poor surface.¹⁰⁵ The term is typically applied to metals that adsorb hydrogen strongly (e.g., Pt, Rh) and is most commonly employed to describe processes that occur at elevated temperatures. Given the weak binding of H₂ to Au,¹⁰⁶,¹⁰⁸ and the low H surface coverages accessible under these H₂ pressures, it is difficult to describe Au as a hydrogen-rich surface. The term “hydrogen spillover”, at least as it is classically defined, probably is not an appropriate description of the chemistry we observe. Prins also points out that Bronsted acid–base chemistry can induce H–D exchange on nonreducible supports and highlights examples where H–D exchange is observed but not attributable to hydrogen spillover.

On reducible supports, hydrogen spillover may be better described as a proton-coupled electron transfer in which the proton transfers to the metal oxide surface and an electron through the metal cations. For Au/TiO₂, there are intimate electronic interactions between the metal and the support, and adsorbing CO appears to induce a partial reduction of the underlying support.¹¹¹ While we believe that the bulk of evidence better supports heterolytic activation at the MSI, it is also consistent with homolytic activation followed by fast deprotonation by the support. Under this description, the H–D exchange that we observe could also be considered as hydrogen spillover. Presumably these processes are at work in the all-proton systems as well.

We also examined the possibility that the observed reaction kinetics and H–D exchange might be due to homolytic H₂ activation followed by fast deprotonation by the support. The potential energy diagram for this pathway is graphed in red in Figure 8 and juxtaposed against the direct heterolytic activation pathway in blue. We note that the energy values.
used in Figure 8 are strictly for the MSI. Homolytic H₂ activation sites with slightly more favorable energetics exist away from the interface as discussed earlier and shown in Table 2; however, these sites are incapable of undergoing proton transfer to the support. It is not surprising that the final states for direct heterolytic adsorption and homolytic adsorption followed by deprotonation are essentially equivalent both structurally and energetically (Table 2).

Both the potential energy diagram in Figure 8 and the activation energetics summarized in Table 2 are consistent with heterolytic dissociation across the interface as the dominant H₂ activation route with a significantly lower barrier to final states that differ only in the location of the Au–H (Figure 8, Table S3). The exothermic nature of the proton transfer step suggests that the formal oxidation of Au during homolytic cleavage makes this process unfavorable. It also indicates that heterolytic cleavage is the preferred pathway because it avoids this formal oxidation. It is also noteworthy that deprotonation of the formal Au hydride by the support is energetically favorable and has only a moderate kinetic barrier of 0.39 eV. This is consistent with the electronegativity of Au providing the driving force for deprotonation. Importantly, this facile deprotonation could result in transfer of all the hydrogens to the support via deprotonation pathways regardless of whether they resulted from homolytic or heterolytic cleavage.

**Heterolytic H₂ Activation Kinetic Model.** Since the DFT and FTIR studies suggest heterolytic H₂ activation at the MSI as the likely pathway, the H₂ dependence data in Figure 4 was re-evaluated using a kinetic model consistent with this mechanism. This pathway also allows us to carefully consider the involvement of support protons (and therefore acid–base centers) in the reaction mechanism. The DFT calculations suggest that H₂ activation occurs at MSI sites with access to basic hydroxyls, and O₂ activation at MSI sites requires acidic hydroxyls (or water). We therefore distinguish between H₂ and O₂ activation sites with the following nomenclature: O₂ activation occurs at MSIA sites and H₂ activation occurs at MSIB (A = acidic, B = basic).

We first examine the possibility that the reaction is hydrogen coverage limited. If H₂ activation was the only important kinetic step, one would expect the kinetics to be first order in hydrogen and zeroth order in oxygen. While the experimental reaction orders are consistent with the rate law describing this mechanism (details in the SI), if the reaction was truly limited by heterolytic H₂ activation, one would expect the measured kinetics to be closer to unity. It appears more likely that the reaction is limited by H coverage, i.e., that a high H₂ activation barrier leads to sufficiently low H coverage to limit the rate of a subsequent step with a lower activation barrier. The most likely candidate for this limiting step is a slow reaction between Au−H and activated O₂ (see additional DFT calculations below).

**Scheme 2. Elementary Steps for H₂ Oxidation via Heterolytic H₂ Activation on Au**

\[
MSIA + O_2 + TiOH \xrightarrow{K_{O_2}} OOH_{MSIA} + TiO^-
\]

(S2-1)

\[
OOH_{MSIA} + TiOH \xrightarrow{E_1} O_2H_{2MSIA} + TiO^-
\]

(S2-2)

\[
2(MSIB + H_2 + TiO) \xrightarrow{E_2} H_{MSIB} + TiOH
\]

(S2-3)

\[
H_2O_{2MSIA} + H_{MSIB} \xrightarrow{k_4} [MSIA]_2 + O_{MSIB} + H_2O
\]

(S2-4)

\[
H_{MSIB} + TiO^\mathrm{fast} \xrightarrow{K_{fast}} MSIB + TiOH
\]

(S2-5)

\[
OH_{MSIA} + TiOH \xrightarrow{K_{fast}} MSIB + H_2O + TiO^-
\]

(S2-6)

Overall Reaction: \(2H_2 + O_2 \rightarrow 2H_2O\)

**Scheme 2** presents this mechanism, assuming that H₂ adsorption is quasi-equilibrated (step S2-3 can therefore be described with \(K_{O_2}\)) and that the reaction between the formal Au hydride and adsorbed H₂O is rate determining.

Initially ignoring the effects of water, a Langmuir–Hinshelwood treatment of the MSIₐ sites yields the rate law in eq 6 under constant H₂O pressure (details in the SI).

\[
\nu_{\text{hetero}} = \frac{k_{4}K^\prime_{O_2}P_{H_2}P_{O_2}[MSIA]_2[MSIB]}{(1 + K_{H_2}P_{H_2})(1 + K_{O_2}P_{O_2} + K_{O_2}^\prime P_{O_2})}
\]

(6)

The associated double-inverse form of the rate law is presented in eq 7.
where $k_{\text{obs}}$ is the combined rate constant defined in eq 8:

$$k_{\text{obs}} = \frac{k_{\text{obs}}}{(1 + K'O_2 + K'H_2'O_2)}$$

The $v_{\text{max}}$ and $K_R$ kinetic parameters can be defined and extracted using eqs 9 and 10:

$$v_{\text{max}} = \frac{1}{\text{intercept}} = \frac{k_{\text{obs}}}{\text{MLE}}$$

$$K_R = \frac{\text{slope}}{\text{intercept}} = \frac{1}{K_{H_2}}$$

This mechanism is readily evaluated with double-reciprocal (Lineweaver–Burk) plots; Figure 9 shows this treatment does a far better job of describing the $H_2$ dependence data than does the homolytic $H_2$ activation mechanism (Figure 5). For both catalysts, $v_{\text{max}}$ terms vary linearly with the feedwater content, while the $K_R$ terms show relatively little change. This is essentially “noncompetitive inhibition” in the Michaelis–Menten terminology. The extracted descriptive parameters ($K_R$ and $v_{\text{max}}$; Figure 10 and Table 3) show that water has essentially no effect on the $H_2$ binding equilibrium ($K_R$), while $v_{\text{max}}$ is inversely related to weakly adsorbed water coverage. It is unlikely that the net rate constant ($k_{\text{obs}}$) varies linearly with water pressure, so the most reasonable conclusion is that the number of available active sites ($[\text{MSI}]_T$) decreases as water is added to the system. This behavior is consistent with water physically blocking the $H_2$ activation sites, as we have observed in other reactions with other reaction poisons.93,94

The water poisoning can be added to the simplified rate law with the simple assumption that the physisorbed water binds to the support, but not preferentially at the MSI. Thus, $Au_{\text{surf}}$ site availability is proportional to the fraction of the oxide surface that is not covered by water and inversely proportional to the weakly adsorbed water coverage on the support (eq 12, more details in the SI). Additionally, CO oxidation, which may report on $O_2$ activation, shows a similar inverse first-order dependence on water at these water pressures (data in the SI). Thus, the addition of a simple water site blocking term is consistent with the loss of both $H_2$ and $O_2$ activation sites. Equations 12 and 13 show the heterolytic rate law and

$$v_{\text{hetero}} = \frac{1}{k_{\text{obs}}[\text{MSI}]_T[\text{MSI}]_T + 1} + \frac{1}{k_{\text{obs}}[\text{MSI}]_T[\text{MSI}]_T + 1}$$

**Table 3. Michaelis–Menten Parameters Extracted from Figure 9 for H$_2$ Oxidation**

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>$P_{H_2O}$ (Torr.)</th>
<th>$\theta_{H_2O}$</th>
<th>$K_R$ (atm)</th>
<th>$K_{H_2}$ (atm$^{-1}$)</th>
<th>$v_{\text{max}}$ ($s^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Au/Al$_2$O$_3$</td>
<td>12</td>
<td>0.79</td>
<td>0.29 ± 0.09</td>
<td>3.4 ± 1.1</td>
<td>0.04 ± 0.02</td>
</tr>
<tr>
<td>9</td>
<td>0.70</td>
<td>0.46 ± 0.09</td>
<td>2.2 ± 0.4</td>
<td>0.14 ± 0.03</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>0.58</td>
<td>0.5 ± 0.1</td>
<td>2.0 ± 0.4</td>
<td>0.27 ± 0.03</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>0.53</td>
<td>0.49 ± 0.07</td>
<td>2.0 ± 0.3</td>
<td>0.35 ± 0.04</td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>0.44 ± 0.09</td>
<td>2.4 ± 0.6</td>
<td>---</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Au/TiO$_2$</td>
<td>12</td>
<td>0.80</td>
<td>0.09 ± 0.03</td>
<td>11.1 ± 3.7</td>
<td>0.06 ± 0.02</td>
</tr>
<tr>
<td>9</td>
<td>0.72</td>
<td>0.16 ± 0.02</td>
<td>6.3 ± 0.8</td>
<td>0.092 ± 0.007</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>0.59</td>
<td>0.22 ± 0.03</td>
<td>4.6 ± 0.6</td>
<td>0.12 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>5</td>
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<td>0.13 ± 0.01</td>
<td>7.7 ± 0.6</td>
<td>0.15 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>0.15 ± 0.02</td>
<td>7.4 ± 2.4</td>
<td>---</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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*Reaction conditions: 60 °C, 10 vol % $O_2$, 3–20 vol % $H_2$, 5–12 Torr $H_2O$, WHSV = 2.3 × 10$^3$ L/g$_{cat}$/h.*
associated double-inverse equation, respectively, with water site blocking included:

$$\nu' = \frac{-1}{k_{P} K_{M} + k_{P} K_{M} + k_{P} K_{M} + k_{P} K_{M}}$$  \hspace{1cm} (12)$$

The mechanism shown in Scheme 2 and the associated rate law (eq 12) are broadly consistent with the kinetic data. The kinetic plots also provide several independent measures of the \(H_2\) adsorption equilibrium constant (eq 12). These values (2.4 \(\pm\) 0.6 atm^{-1} for Au/Al\(_2\)O\(_3\) and 7 \(\pm\) 2 atm^{-1} for Au/TiO\(_2\)) are reasonably small and appropriately describe the weak binding of \(H_2\) to Au catalysts. These values are entirely in line with the near-thermoneutral \(H_2\) binding predicted by the DFT model.

It is likely that the general nature of the proton-assisted pathways is similar on Au/Al\(_2\)O\(_3\) catalysts, although we expect some differences in the ability of the nonreducible support to assist in stabilizing the negative charges associated with the Au hydride. Considering the significant computational cost, we deemed a full DFT investigation of the Au/Al\(_2\)O\(_3\) catalyst too expensive to pursue.

There are relatively few quantitative measures of \(H_2\) adsorption thermodynamics on supported Au catalysts, making direct comparisons with the literature difficult. Van Bokhoven and co-workers measured volumetric \(H_2\) chemisorption over several Au/Al\(_2\)O\(_3\) catalysts.\(^{112}\) Their H: Au ratios attributable to strong chemisorption were small, varying between 5% and 20%.
for different catalysts over a wide range of temperatures. These values are consistent with H₂ activation at the MSI. Thomson also calculated the H₂ heat of adsorption on a series of Au/TS-1 type catalysts and found exceedingly weak heat of adsorption values around 1−2 kcal/mol.¹¹³

There are a few additional experimental measurements of H₂ adsorption on Au model systems. Stobinski and co-workers have reported several studies of H₂ adsorption on Au films, determining a heat of adsorption of 21 kJ/mol.¹¹⁴ The weak H₂ binding sites were attributed to undercoordinated atoms at edges, corners, and kink/step sites.¹¹⁵ Madix and Campbell reported very weak H₂ adsorption on hot Au filaments at low temperatures (216 K), with a dissociative activation barrier of 51 kJ/mol.¹¹⁶ Note that adsorption on these Au surfaces was assumed to be traditional homolytic chemisorption, so the determined values may not be directly comparable to the heterolytic process that our data suggest. Rather, they suggest that traditional chemisorption on Au may be so weak that alternative H₂ activation pathways become more attractive explanations for the observed reactivity.

**Alternative Reaction Pathways.** We also considered the possibility that the reaction proceeds through a traditional homolytic cleavage mechanism away from the MSI. Dumesic’s work on H₂ oxidation over supported Pt catalysts showed that H₂ chemisorption followed by a rate-determining reaction with an active oxygen species can have a maximum H₂ reaction order of 0.5. However, if a subsequent step involving Au-H is rate determining, the observed reaction order could approach unity (derivations available in the SI). While we cannot rule such a mechanism out entirely, it is not supported by the kinetic data or the DFT calculations. Because water does not bind to Au, it is difficult to rationalize the strong water inhibition at support θ_H₂O values around 0.5 if the reaction does not occur at the metal−support interface. Additionally, if a later step was rate determining, we would expect a relatively high coverage of Au-OOH, since this species binds both rapidly and strongly.¹³ Given the very few reports characterizing the nature of O₂ binding to Au catalysts,¹¹⁷ this seems unlikely. Finally, the rapid H−D exchange demonstrated with FTIR shows that gas phase H₂ exchanges with support protons and that this rate is comparable to the overall reaction rate.

It is far more plausible that the reaction is limited by the thermodynamics and kinetics of hydrogen binding. We therefore examined the subsequent elementary steps with the DFT nanorod model; details are provided in Table S3. Table 4 compiles these results, first examining the elementary steps that occur on metal sites (i.e., without the aid of MSI protons). Note that all of the steps after homolytic H₂ activation have lower activation barriers than dissociative chemisorption. This further supports the kinetic evidence that H₂ oxidation does not proceed through this traditional pathway on Au.

The second half of Table 4 adds the elementary steps that involve proton transfer chemistry. Entry P1 shows the exchange reaction between cus-OH₂⁺ and bridge-O⁻ is
essentially thermoneutral. Consequently, the cus-OH$_3^-$ and bridge-OH protons have similar acidities, and cus-OH and bridge-O$^-$ are of similar basicity; this simplifies the surface chemistry substantially. For convenience, we consider the support as a proton source/sink and generally refer to support protons as H$_{\text{MSI}}^+$ for the remainder of this discussion: the energetics of the processes will not change significantly if an appropriate cus-OH or bridge-OH site is used. This also helps consider the reaction kinetics: particularly in the presence of water, the support can rapidly supply protons to or remove them from species adsorbed on the metal.

Next, Au-H deprotonation by the support is thermodynamically favorable and has a lower barrier (0.39 eV) than all but one of the calculated barriers for the metal-only reactions. This suggests that even if a dissociative chemisorption mechanism was at work, deprotonation would likely be fast relative to other steps that might occur on the Au. Thus, regardless of the details of H$_2$ activation, the catalysis likely proceeds through the significantly faster proton-mediated chemistries that the support affords.

We evaluate the remaining possible reactions in a stepwise fashion considering each of the likely O-containing intermediates individually. Each intermediate has three nominal reaction pathways, shown graphically in Figure 11: (i) reaction with Au-H, (ii) reaction with a MSI proton (H$_{\text{MSI}}^+$), or (iii) reaction with a free Au site (this leads to dissociation or disproportionation depending on the nature of the intermediate). For Au-OOH (Figure 11A), Au-OH (Figure 11C), and Au-O (Figure 11D), protonation reactions are nearly barrierless and are always considerably faster than reaction with Au-H or any of the alternate pathways. The exception to this is Au-H$_2$O$_2$ (Figure 11B), which has low barriers to react with either Au-H or H$_{\text{MSI}}^+$. Direct dissociation of Au-H$_2$O$_2$ is also a viable reaction pathway.

These results are illuminating and highlight the special reactivity of Au catalysts. First, the relatively fast proton transfer steps enable a variety of chemistries (e.g., heterolytic H$_2$ activation, fast Au-OH decomposition) that are unavailable when only metal-catalyzed pathways are considered. These weak Brønsted acid–base chemistries generally have lower activation barriers than the comparable metal-only pathways. Indeed, the pathways involving H$_{\text{MSI}}^+$ and Au-OOH, Au-OH, and Au-O are essentially barrierless despite the fact that the deprotonation energetics in Figure 8 indicate the H$_{\text{MSI}}^+$ species to be ca. $\sim$0.7 eV more stable than Au-H. This is an intriguing result, as it indicates that the addition of weak Brønsted acid–base chemistry to the metal system breaks the predicted transition state scaling relations that hold for many other systems. Readers are directed to a recent perspective by Kumar et al. for an excellent review on how interfacial sites can employ different chemistries to overcome the limitations of metal scaling relations.

Second, the unfavorable energetics for formal metal oxidation, along with the associated high activation barriers, are a key feature of the Au chemistry (e.g., dissociative chemisorption/oxidative addition). Consequently, pathways that deliver electrons to the electronnegative Au or avoid formal oxidation are generally more facile. Finally, the metal–support interface provides an interesting means of considering H$_2$ oxidation over these materials. Essentially, the fastest pathways involve supplying electrons to the Au (and ultimately to the adsorbed O species) through H$_2$ adsorption. Protons are concurrently delivered to the support via heterolytic H$_2$ adsorption and Au–H deprotonation. These Au catalysts function essentially via proton-coupled electron transfer mechanisms, with hydrogen electron density distributed both to the Au atoms and, at least for titania, the underlying support. In a sense, this mechanism can be considered as being closely related to electrochemical reactions, with (conceptually) hydrogen activation and deprotonation occurring at basic MSI sites and O$_2$ activation occurring at acidic MSI sites. The Au (with help from the support) then serves as electron source/sink, depending on the half-reaction. This is shown conceptually in Scheme 3.

These processes are remarkably similar to those that Wilson and Flaherty have identified in their detailed examination of the liquid phase partial reduction of O$_2$ to H$_2$O$_2$ over Pd/SiO$_2$ and PdAu/SiO$_2$ catalysts. Their work showed that protic solvents, which act as fast proton carriers whose concentration is determined by solution pH, dramatically improve catalyst activity. In our work protons are transferred across the hydroxylated and/or water-covered support; their chemical potential is determined by the pressures of H$_2$O and H$_2$. One of the key differences between the two systems is that O$_2$ activation on Au requires a proton to generate Au-OOH in a single step, while the Pd system appears to go through a series of stepwise proton–electron transfer steps mediated by a protic solvent. The considerably less favorable and site-specific activation of H$_2$ at the MSI for Au catalysts is also an important difference in these systems. These observations are consistent with the support effects Hutchings and co-workers observe in their PdAu H$_2$O$_2$ synthesis catalysts after acid pretreatments.

**Implications for PrOx.** Our initial motivation for studying H$_2$ activation was to better understand the role of water in improving the PrOx reaction. It is now clear that water plays two beneficial roles during PrOx. First, water, or another proton source, is required for fast CO oxidation, as it is involved in both O$_2$ activation and rate-determining Au-COOH decomposition. Second, water clearly poisons H$_2$ activation at the MSI. This suggests that a less volatile proton source at the MSI can accomplish two complementary goals: (i) it may be possible to suppress H$_2$ activation by selectively blocking the MSI sites without poisoning the desirable CO oxidation reaction, and (ii) faster CO oxidation rates may be achieved at higher temperatures at which physiosorbed water would readily desorb. Either of these outcomes would constitute a significant improvement to the process conditions for the PrOx reaction.

**Comparisons to Other H$_2$ Activation Systems.** There are numerous precedents for heterolytic H$_2$ activation mechanisms in the enzyme and inorganic chemistry literature. Several groups have suggested heterolytic H$_2$ activation by...
hydrogenases, and Crabtree has shown that Ni−Fe hydrogenases employ a heterolytic H₂ activation process, resulting in a formal hydride on the (unoxidized) Ni and a proton on an adjacent bridging oxo group. This conclusion has been supported by DFT calculations and substantial synthetic modeling and is fundamentally similar to what we observe for the Au catalysts. Several systems of transition metal complexes have been shown to activate H₂ heterolytically. Metal hydrides have long been known to be acidic, so these systems might be considered as operating via the traditional oxidative addition mechanism followed by rapid deprotonation. There is also now a fairly extensive literature of systems that might be considered as operating via the traditional model for homolytic H₂ adsorption and subsequent dissociation of H₂ at the surface, which effectively function as FLPs. In this case, the heterolytic activation step was suggested as an alternative to hydrogen spillover to the titania support; the homolytic dissociation of H₂ over ruthenium metal sites remains the fastest H₂ scission pathway in this system.

Heterolytic H₂ activation has also been implicated at the Ru/TiO₂ MSI during hydrodeoxygenation of phenols. In this case, the heterolytic activation step was suggested as an alternative to hydrogen spillover to the titania support; the homolytic dissociation of H₂ over ruthenium metal sites remains the fastest H₂ scission pathway in this system. The dominant heterolytic H₂ activation over a supported metal catalyst is a surprising discovery; we are aware of exceedingly few examples of solid systems that have reported convincing experimental evidence of this general pathway. Notably, Copéret and co-workers have shown that γ-Al₂O₃, when treated at appropriately high temperatures, can activate H₂ and catalyze the hydrogenation of simple alkenes. The thermal treatment generates defect sites on the alumina surface, which effectively function as FLPs. Tomishige and co-workers also found a first-order hydrogen dependence in studying hydroxyl-containing ether hydrogenolysis over Brønsted bases. They similarly argued that the hydrogen dependence was most consistent with a net heterolytic activation of H₂ with the proton being transferred to the water solvent and the hydride being stabilized by the Rh. Further, Stair and co-workers have also recently prepared single-site supported aluminum catalysts on catechol-containing porous organic polymers. These materials hydrogenate alkenes through H₂ insertion into Al−O bonds, resulting in a formal hydride on the aluminum atom.

Several groups have shown that the addition of Brensted bases increases the hydrogenation activity of supported Au catalysts. For example, Cao and co-workers reported that the addition of quinolines dramatically improved hydrogenation activity over Au while suppressing activity over Pt, Pd, and Ru catalysts. The addition of a variety of amine bases, including cycloaliphatic diamines, to several supported Au catalysts resulted in improved alkene partial hydrogenation activity. In their study, Rossi and co-workers considered these Au-amine systems to be FLPs. Similar improvements in 1,3-cyclohexadiene partial hydrogenation were found for Au particles suspended in imidazolium ionic liquids. In all these studies, heterolytic H−H bond activation was implicated in the improved activity. We also note that incorporating secondary phosphine oxide ligands into solution-phase ligand stabilized Au nanoparticles (NPs) resulted in improved activity and selectivity in cinnamaldehyde hydrogenation. These clever systems incorporated a basic o xo group into the phosphine to assist in heterolytic H₂ activation.

The heterolytic activation of H₂ over Au catalysts therefore has significant precedent in other similar chemistries. This literature also provides some important context to understand this system. The active sites at the MSI can be considered to be composed of a Brønsted base (surface hydroxyl), which stabilizes the developing proton in close proximity to a soft Lewis acid (Au NP), which stabilizes the developing hydride. The surface hydroxyls on alumina and titania are not particularly strong bases, which suggests that this reaction is driven by the activity of the Au NP to stabilize the developing hydride.

Haruta’s group originally showed that H₂ activation rates track with the number of metal—support interface sites and determined a consistent turnover frequency for H₂ dissociation across a large range of particle sizes assuming that the active sites were at the MSI. Takeda and co-worker’s DFT study found a similar decrease in the activation barrier for H₂ dissociation at the MSI on Au/TiO₂. This study also suggested H₂ dissociation through an O²−−H+−H−Au pathway at the Au/TiO₂ interface was energetically favored compared to the Au−H−H−Au pathway. In a different study using H₂−D₂ exchange reactions, Nakamura et al. also propose the Au²⁺−O⁶⁻−Ti sites at the Au/TiO₂ interface to be the active sites for H₂ dissociation. The strong evidence for heterolytic H₂ activation reported here provides a clear mechanistic understanding of these reports.

Many other supported metal catalyst systems contain soft Lewis acids in close proximity to weak Brønsted bases, yet undergo homolytic H₂ activation on the metal. This is likely due to the electronic structure and high electronegativity of Au. Dissociative chemisorption, which is analogous to oxidative addition in transition metal complexes, requires a formal two-electron oxidation of the metal NP. The high electronegativity of Au, combined with the particularly stable full d-band, make Au the most noble metal and increase the thermodynamic and kinetic barriers for dissociative chemisorption. The heterolytic pathway does not require a formal oxidation of the metal, and thus appears to be the favored H₂ activation pathway for Au.

### CONCLUSIONS

Kinetic experiments for H₂ oxidation over Au/TiO₂ and Au/Al₂O₃ catalysts show unexpectedly high reaction orders for H₂ and strong reaction inhibition by water physisorbed on the support. Combined with previous results for water-assisted CO oxidation on Au catalysts, this study shows that water drastically improves PrOx performance by promoting CO oxidation and inhibiting the undesirable H₂ oxidation reaction. However, the reaction kinetics are inconsistent with the traditional model for homolytic H₂ adsorption and subsequent hydrogenation of oxygen moieties exclusively on metal sites. DFT studies using rutile TiO₂(110)-supported Au nanorod model indicate that H₂ activation on the surface is most facile across the MSI. Surprisingly, the DFT model shows this to occur through a heterolytic H−H dissociation pathway, resulting in formation of a support hydroxyl group and a formal hydride adsorbed on the Au. This pathway does not require the formal oxidation of Au associated with the traditional homolytic activation mechanism that is commonly seen on other metals. Infrared spectroscopy experiments during H₂ adsorption on a deuterated Au/TiO₂ catalyst showed H−D scrambling with the metal oxide, lending further support for the heterolytic activation pathway. The experimental reaction kinetics were also consistent with the heterolytic H₂ activation model, and showed that the water
poisoning of H₂ oxidation was largely due to active site blocking.

The reaction network provided important fundamental insights into the nature of the catalysis. Protons near the MSI were calculated to be more stable than formal Au hydrides; yet, the MSI protons play the dominant role in fastest pathways for water formation. From the perspective of omnipresent transition state scaling or Brønsted–Evans–Polanyi relationships, this finding is counterintuitive. This system provides an example of how different types of chemistries (in this case metal sites with weak Brønsted acid/base chemistry) can be combined to overcome scaling relations and lead to significantly faster catalysis.

■ ASSOCIATED CONTENT

 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.8b04991.

O₂ reaction order plots; water effects on CO oxidation on Au/TiO₂; homolytic and heterolytic H₂ activation mechanisms and rate law derivations; effect of compressive strain and surface hydroxylation on the reactivity of Au/TiO₂; calculated energetics and configurations for each elementary step (PDF)

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Notes

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