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### Single-Atom Catalysis

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# **Observing Single-Atom Catalytic Sites During Reactions with Electrospray Ionization Mass Spectrometry**

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Dedicated to Professor Maria Flytzani-Stephanopoulos

Abstract: Single-atom catalysts (SACs) have become a prominent theme in heterogeneous catalysis, not least because of the potential fundamental insight into active sites. The desired level of understanding, however, is prohibited due to the inhomogeneity of most supported SACs and the lack of suitable tools for structure-activity correlation studies with atomic resolution. Herein, we describe the potency of electrospray ionization mass spectrometry (ESI-MS) to study molecularly defined SACs supported on polyoxometalates in catalytic reactions. We identified the exact composition of active sites and their evolution in the catalytic cycle during CO and alcohol oxidation reactions performed in the liquid phase. Critical information on metal-dependent reaction mechanisms, the key intermediates, the dynamics of active sites and even the stepwise activation barriers were obtained, which would be challenging to gather via prevailingly adopted techniques in SAC research. DFT calculations revealed intricate details of the reaction mechanisms, and strong synergies between ESI-MS defined SAC sites and electronic structure theory calculations become apparent.

#### Introduction

Single-atom catalysts (SACs) are sometimes superior to their particle counterpart because of enhanced metal dispersion, more variable oxidation states and intimate metalsupport interactions.<sup>[1]</sup> Various combinations of metal and support have been employed for numerous catalytic applications like CO<sup>[2]</sup> and alcohol oxidation,<sup>[3]</sup> selective hydrogenation,<sup>[4]</sup> and electrochemical reactions<sup>[5]</sup> among others.<sup>[1b,e,6]</sup> Although parallels are drawn between homogeneous and heterogeneous catalysts,<sup>[7]</sup> substantial questions remain about SACs regarding: (i) the exact molecular structure of the active site, (ii) the dynamics of that active site under reaction

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conditions,<sup>[8]</sup> as well as (iii) activation barriers between various intermediates in the catalytic cycle. Experimental techniques employed so far for model SACs have only been able to provide limited and imprecise information on the active site hampered by their intrinsic constraints. Spectroscopic methods like X-ray photoelectron (XPS), X-ray absorption (XAS) or Mößbauer spectroscopy provide averaged information of observable species, and thus are markedly affected by the presence of spectators. Further, they are element-specific so only provide limited and biased information from the point of view of a particular component of the catalyst.<sup>[9]</sup> Although state-of-the-art microscopy offers atomic resolution of active sites, changes in the support structure and adsorption of intermediates are difficult to discern. Furthermore, microscopy-based techniques do not allow the acquisition of statistic information of a large number of active sites.<sup>[10]</sup> Infrared spectroscopy is also commonly used but only offers indirect deductions on the active sites from the perspective of infrared light-responsive surface adsorbates. Those shortcomings highlight the urgent need to develop new methods to interrogate SAC active sites under reaction conditions.

Another reason for the limited understanding of active sites is the complexity of most SACs which commonly use inherently inhomogeneous supports like metal oxides with many different adsorption sites where each single-atom site may exhibit dissimilar structure and activity. Recently, we found that atomically dispersed noble metals supported on polyoxometalates (POMs) are active catalysts for hydrogenation and CO oxidation reactions.<sup>[11]</sup> POMs are considered to be models for metal oxides with structural simplicity, molecularly defined active sites and solubility in polar solvents. Many different classes of inorganic compounds comprising transition metals and POMs have been reported before with applications ranging from medicine  $^{\left[ 1\bar{2}\right] }$  and electronic devices<sup>[13]</sup> to catalysis.<sup>[14]</sup> A variety of POMs have been synthesized composed of many different elements and exhibiting diverse geometries.<sup>[15]</sup> Keggin-derived lacunary POMs of the type  $[PM_{11}O_{39}]^{n-}$  have been among the earliest to be reported, featuring high charge states and high stability. Most transition metal-based complexes comprising POMs have been based on those lacunary POMs with Pd, Rh, or other transition metals.<sup>[16]</sup> They serve as excellent homogeneous catalysts predominantly for oxidation reactions but also enable the in-depth study of reaction mechanisms and active sites in the more commonly used heterogenous POM-based catalysts.<sup>[17]</sup> As most POMs are symmetric and thus only offer a few discernible and sufficiently strong adsorption sites (in some cases only one), they also serve as a platform to

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investigate reaction mechanisms and parameters governing catalytic activity.<sup>[9b]</sup> We envisage, that solvent dispersible, POM-supported SACs will be an ideal model system to study the active sites and their dynamics during catalytic reactions with techniques capable of providing structural information with high sensitivity and accuracy, such as mass spectrometry.

Electrospray ionization mass spectrometry (ESI-MS) is based on the formation of an aerosol which is de-solvated and directed by an electric field to yield isolated ions in the gas phase. Due to the mild ionization, intact ions from the liquid phase can be separated by a mass filter and analyzed by a detector without fragmentation. In the ionization process, negative charges are compensated by ubiquitous cations like  $H^+$ , Na<sup>+</sup> or K<sup>+</sup>. The intact adduct ions can then be separated and fragmented in a controlled manner by accelerating and colliding them with argon atoms. Some of the transferred kinetic energy is used to induce bond breaking and the resulting fragmented ions are further separated and detected (Figure 1). Ion separation allows the identification of very low concentrations of intermediates within a complex reaction mixture, providing information on the exact composition of the molecular structure<sup>[18]</sup> of the active site including the binding of adsorbates, changes in the support structure<sup>[19]</sup> as well as the charge state of the metal center and the metalsubstrate complex (intermediates).<sup>[20]</sup> Fragmentation provides further confirmation of the composition, the binding strength of adsorbates as well as the relative stability of intermediates. ESI-MS also shows relative concentrations of intermediates, revealing information on kinetically relevant steps and enabling the quantitative analysis of reaction barriers between intermediates. As we are aware, those



*Figure 1.* ESI-MS as a tool for the elucidation of single-atom catalyzed reaction. Reaction solution including the solubilized SAC, solvents and reaction intermediates is directly injected into the ionization chamber. The nebulized reaction mixture is heated and accelerated by an electric potential while the solvent droplets are shrinking until the isolated ions are in the gas phase. Mass filters (quadrupolar magnets are shown as an example) then separate the different species which are determined by a mass detector. If further information about the reaction intermediates should be obtained, ions passing through mass filter #1 can be fragmented in a collision cell and the resulting ions undergo another separation using mass filter #2 before detection.

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advantages of ESI-MS have not been used to investigate SAC systems.

In the present work, we use ESI-MS to study active sites and catalytic reaction mechanisms of POM-supported SACs. During the testing of dozens of POM-metal combinations, we identified three main criteria for the ESI-MS study of stable POM-SACs for catalytic reactions: (i) POM-metal mixtures must be completely soluble, (ii) POM-metal has to be strongly bonded together in a stoichiometrically defined manner, (iii) POM has well-established, reliable synthetic protocol (Figure S1). Based on these criteria, we identified  $PW_{11}O_{39}^{7-}$ derived SACs to be the most promising ones for this study. CO and benzyl alcohol oxidation reactions were conducted with SACs based on different metals in the liquid phase. Not only all relevant intermediates hinting at a Mars-van Krevelen (MvK) mechanism were observed by ESI-MS, but also the rate-limiting step in  $PW_{11}O_{39}Rh_1^{\,4-}$  catalyzed CO oxidation was determined. Since ESI-MS does not provide direct information on the structure and charge density distribution in the active site, we also employed DFT calculations to confirm the oxidation state changes during the reaction cycle as well as the exact geometry of the experimentally observed intermediates. The POM-based SACs are molecularly well defined with known composition during reaction and are thus ideal systems for DFT calculations. For the liquid phase alcohol oxidation, we determined different reaction mechanisms over PW<sub>11</sub>O<sub>39</sub>Fe<sub>1</sub><sup>4-</sup> and PW<sub>11</sub>O<sub>39</sub>Pd<sub>1</sub><sup>5-</sup> catalysts and measured the reaction barriers between different intermediates. This study exemplifies the effectiveness of ESI-MS to examine molecularly defined SACs containing different active metal species in a broad range of reactions.

#### **Results and Discussion**

The synthesis of the lacunary POM was performed following a known procedure starting from sodium tungstate.<sup>[21]</sup> The purity of the sample was confirmed by infrared spectroscopy (Figure S2)<sup>[22]</sup> by comparison to phosphotungstic acid (H<sub>3</sub>PW<sub>12</sub>O<sub>40</sub>) and sodium tungstate. <sup>31</sup>P NMR was further used to verify the absence of P-containing impurities in the POM precursors as well as catalyst samples (Figure S3). As reasoned before, due to their high negative charge, POMs like  $PW_{11}O_{39}^{7-}$  are able to bind metal cations strongly although the bonding includes both ionic as well as covalent character. When mixing molar ratios of 1:1 (POM:Rh<sup>3+</sup>), all the peaks for free  $PW_{11}O_{39}^{7-}$  in the solution disappeared and thus quantitative conversion of PW<sub>11</sub>O<sub>39</sub><sup>7-</sup> into PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> could be assumed (Figure 2a). Rh likely coordinates to four oxygen in the POM (inset in Figure 2a), as determined by single crystal X-ray crystallography before,<sup>[23]</sup> and corroborated by DFT calculations. Although varying compositions for PW11O39Rh14- derivatives have been reported including the Cl-containing species, as well as the formation of PW<sub>11</sub>O<sub>39</sub>Rh<sub>2</sub> or the respective dimeric species  $(PW_{11}O_{39}Rh_1)_2$ , [16a, 24] these compounds were not observed by ESI-MS in our system.

We investigated the CO oxidation mechanism for those structurally defined catalysts, using ESI-MS. When dilute CO



Figure 2. ESI-MS measurements during the CO oxidation reaction on  $PW_{11}O_{39}Rh_1^{4-}$ . a)  $PW_{11}O_{39}^{7-}$  (black) with the respective cation envelope for z = 3.  $PW_{11}O_{39}Rh_1^{4-}$  (orange) with a highly reduced cation envelope due to the adsorption of trivalent Rh<sup>III</sup>. The inset shows the structure of PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> as determined by single crystal X-ray crystallography before. Since the POM can contain x protons, y sodium and zpotassium cations, the following nomenclature will be used in the following for the unequivocal description of ionic species:  $\{x, y, z\}$ . b) ESI-MS spectra of  $PW_{11}O_{39}Rh_1^{4-}$  after treatment under (black) air, (orange) 2 min CO, (blue) 1 h CO, (green) 1 h CO and subsequently 1 h O<sub>2</sub> and (yellow) 1 h CO and subsequently 24 h O<sub>2</sub>. Reactions were carried out in a  $V_{\text{total}} = 0.02 \text{ L}$  deionized water with  $C_{\text{PW11039}} = 0.5 \text{ mM}$ and  $C_{\rm Rh}$  = 0.5 mM and  $F_{\rm total}$  = 100 cm<sup>3</sup> min<sup>-1</sup> of each gas (5:95 CO:Ar and 5:95  $O_2$ :Ar) at a  $p_{total} = 1$  bar and room temperature. Samples were collected out of the reaction mixture and immediately analyzed by ESI-MS.

was bubbled through a solution of  $PW_{11}O_{39}Rh_1^{4-}$  for 2 min, {[ $Rh_1^{III} _{3^+}^{3+}(CO)_2$ -( $PW_{11}O_{39}$ )<sup>7-</sup>] + H<sup>+</sup> + K<sup>+</sup>}<sup>2-</sup> (**a-I**, calc.: 1438.54, expt.: 1438.52) with two geminal CO molecules appeared without the reduction of  $Rh^{3+}$  (Figure 2b). In contrast, bubbling CO through a solution of  $PW_{11}O_{39}^{7-}$  did not yield mass signals consistent with CO adsorption on the POM (Figure S4). This confirms that the adsorption of CO and its further oxidation is associated with the formation of  $PW_{11}O_{39}Rh_1^{4-}$  as active site.

As mentioned above, the mass-selected intermediates can be fragmented to obtain further information about the composition. At a collision energy of 10 eV, a peak accounting for the desorption of one CO molecule becomes apparent but it seems that the CO oxidation reaction also occurred in the collision cell and species a-I was converted into species  $\{[Rh^{III}_{1}^{3+}(CO)-(PW_{11}O_{38})^{7-}]+H^{+}+K^{+}\}^{2-}$  (**a-II**, calc.: 943.70, expt.: 943.68) (only observed during fragmentation of a-I) after desorption of the produced CO<sub>2</sub> (Figure 3a and Figure 4). Other peaks were ascribed to a cation exchange reaction in the gas phase that commonly occur during ESI-MS experiments.<sup>[25]</sup> Reaction of a-I at relatively low collision energy indicated that the reaction barrier towards the oxygen vacant species was low. When the solution was treated with CO for 1 h, the previously observed species (a-I) disappeared and two species: { $[Rh_1^{1+}(CO)(CO_2)-(PW_{11}O_{38})^{5-}] + 2K^+$ }<sup>2-</sup> (calc.: 1456.53, expt.: 1456.51) and  $\{[Rh_1^{1+}(CO)_2 - (PW_{11}O_{38})^{5-}]\}$ 



**Figure 3.** Fragmentation pattern for a) species **a-I** {[Rh<sup>III</sup><sub>1</sub>(CO)<sub>2</sub>-PW<sub>11</sub>O<sub>39</sub><sup>7-</sup>] + K<sup>+</sup>}<sup>3-</sup> and b) species **a-V** {[Rh<sup>I</sup><sub>1</sub>(CO)<sub>2</sub>(O<sub>2</sub>)-PW<sub>11</sub>O<sub>38</sub><sup>6-</sup>] + Na<sup>+</sup> + K<sup>+</sup>}<sup>3-</sup> at 10 and 6 eV collision energy, respectively. Reaction conditions are the same as in Figure 2.

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 $+ H^+ + K^+$ <sup>2-</sup> (**a-III/a-IV**, calc.: 1432.06, expt.: 1432.07) were formed where one of the oxygen atoms from the support was removed (Figure 2b). This strongly suggested the involvement of a MvK type mechanism with the rapid formation of an oxygen vacancy at room temperature concomitant with the reduction of Rh<sup>3+</sup> to Rh<sup>+</sup>. Although MvK mechanisms have been among the many debated proposed reaction mechanisms for CO oxidation, we present here solid experimental evidence of stoichiometric ratios for atomically precise active sites.

In order to close the catalytic cycle, O<sub>2</sub> was bubbled through the reaction system following 1 h CO treatment. After 1 h, a different species appeared with a composition consistent with  $\{[Rh_1^{I_1}(CO)_2(O_2)-(PW_{11}O_{38})^{5-}] + H^+ +$  $Na^{+}$ <sup>2-</sup> (**a-V**, calc.: 1440.08, expt.: 1440.06) including an O<sub>2</sub> molecule bound to the oxygen vacancy. Fragmentation of that intermediate led to the formation of two species with one or two desorbed CO molecules, respectively. Furthermore, fragmentation also led to the splitting of O<sub>2</sub>, and the oxidation and desorption of CO<sub>2</sub> while the oxygen vacancy was refilled and Rh<sup>+</sup> reoxidized. The formed species lost CO to yield PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> in the gas phase. No desorption of O<sub>2</sub> was observed indicating the strong adsorption of O2 into the oxygen vacancy (Figure 3b). Bubbling O<sub>2</sub> through a solution containing PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> without prior reduction by CO did not show O<sub>2</sub> binding on Rh (Figure S5). This further confirms that the oxygen vacancy formation was necessary for strong O<sub>2</sub> binding.

Reoxidation of Rh and refill of the oxygen vacancy did not occur to a significant extent after 1 h although a small but distinguishable new peak was observed (Figure S6a) which we assumed to be the reoxidized species {[Rh<sup>III 3+</sup><sub>1</sub>(CO)(CO<sub>2</sub>)- $(PW_{11}O_{39})^{7-}]$  +  $H^+\}^{3-}$  (calc.: 951.38, expt.: 951.39). The low intensity of the peak indicates a high reaction barrier from its predecessor species and a low relative stability compared to the catalyst state after CO<sub>2</sub> desorption. That structure,  $\{[Rh^{III}_{1}{}^{3+}(CO)-(PW_{11}O_{39})^{7-}] + K^{+}\}^{3-} (a-VI, calc.: 951.38,$ expt.: 951.39), may be observable (Figure S6a) but an overlap with one of the other intermediates formed upon CO bubbling limits the fidelity of the assignment. Either treatments at room temperature for 24 h or at 70 °C for 4 h were necessary for complete catalyst reoxidation and CO desorption in order to regenerate the PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> species (Figure S6). Overall, the reaction cycle determined by ESI-MS followed an MvK mechanism with the formation of an oxygen vacancy followed by its refill by O<sub>2</sub>. The subsequent splitting of O<sub>2</sub>, and oxidation of another CO molecule is ratedetermining (Figure 4). The findings obtained for a solubilized SAC, including the overall reaction mechanism and the ratedetermining step, matched the results obtained before by combined use of operando XAS, in situ XPS and in situ infrared techniques over the heterogeneous Rh-POM SAC.<sup>[9b]</sup> Concrete evidence about the reaction stoichiometry and active site composition, which were not available previously, is reported here with less experimental and computational efforts, and without the need for synchrotron facilities.<sup>[9b]</sup>

Based on the well-defined catalyst stoichiometries from the MS experiments, we performed DFT calculations to investigate the intermediate structures found by MS as well as



*Figure 4.* Reaction mechanism of the Rh single-atom catalyzed CO oxidation based on ESI-MS and DFT calculations; the calculated and experimentally determined masses of representative ions are displayed, the rate-determining step is highlighted in red. In the center, the DFT computed energy profile of the CO oxidation reaction is displayed. The numbers in parenthesis are the calculated reaction barriers for each elementary step. In this Figure, the PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> is simulated as a charged species in gas phase.

the interconnecting reaction barriers. We considered two different models as two limiting cases for the charging state of the catalyst: the first one regards PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> as isolated, fully deprotonated species, and the second model considers the impact of counterions (protons) rendering the catalyst system neutral. The results for the charged model are shown in Figure 4 and Figure S7. The first CO adsorption on PW<sub>11</sub>O<sub>39</sub>Rh<sub>1</sub><sup>4-</sup> is very strong with an adsorption energy of -2.38 eV, compared with the adsorption of O<sub>2</sub> (-0.99 eV) or H<sub>2</sub>O molecule (Table S2), producing the stable structure **a-VI** (Figure 4). The second CO molecule reacts directly with one bridging oxygen forming a carbonate species (a-I) through a small barrier (TS1,  $E_a = 0.20 \text{ eV}$ ). The calculated desorption of  $CO_2$  from **a-I** is moderately endothermic (+0.68 eV) with a barrier of 1.10 eV (TS2), creating an O vacancy. Considering the increase of entropy for  $CO_2$  in gas phase, this step will still be exergonic. Intermediate a-II is unstable and can transform to **a-III** with a very small barrier ( $E_a = 0.07 \text{ eV}$ ). This implies that the concentration of a-II is low and may explain why it is not easily observed in the ESI-MS experiments. Based on a charge density analysis, Rh bound to the lacunary POM is significantly reduced after the formation of the oxygen vacancy which is consistent with a formal oxidation state change from  $Rh^{III}$  to  $Rh^{I}$  (Table S1 and Figure S7) as expected from our previous in situ spectroscopic studies on a comparable Rh SAC.<sup>[9b]</sup> Concomitant with the reduction of Rh<sup>III</sup>, the charge density of the W atom adjacent to the oxygen vacancy

is also reduced noticeably. With the change of Rh oxidation state, the ligand structure around Rh rearranges from a slightly distorted octahedron to a square-planar binding mode (a-IV). Calculations hence suggest that the active site structure changes during the reaction with alternating binding of Rh to five, four, three or two oxygen atoms of the POM support. From square planar a-IV, O2 adsorbs, both re-oxidizing Rh to 3 + and forcing it into an octahedral environment again. This step is rather difficult with a barrier of 1.33 eV. The last step forms the second CO<sub>2</sub> molecule and regenerates the PW11O39Rh14- structure in a highly exothermic manner.

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Although calculations on this charged model appear to be valid for the investigation of reactions occurring in the gas phase after stripping most of the counterions, the solution structure of the catalyst must be assumed to be different. Therefore, we considered the catalyst structure with protons as counterions in order to neutralize the system. We first optimized 138 structures with different proton locations, and found that protons prefer to

occupy the bridging oxygen atoms near Rh<sup>III</sup> (Figure S9 and Table S2). Starting with this optimal structure, we further analyzed the reaction intermediates as well as the interconnecting transition states. With counterions, most reaction intermediates show similar geometries as their counterparts in the charged model (a'-I to a'-VI in Figure S10 comparing with a-I to a-VI in Figure S7). The pathway for the first CO adsorption on a'-VI, however, is different: after CO adsorption, a'-VII is firstly formed and then it can isomerize to a'-I with a small barrier of 0.58 eV, or it directly forms a'-II with a barrier of 0.71 eV. Although the isomerization (from a'-VII to  $\mathbf{a'-I}$ ) is faster, the CO<sub>2</sub> formation (TS'-2) is slower and therefore, the two pathways may co-exist in reactions. In addition, O<sub>2</sub> adsorption is barrierless in the neutral model (from a'-IV to a'-V) showing a big difference to the charged model (Figure 4 TS5). Finally, the conversion from a'-II to a'-**III** is slower than that in the charged model, although the barrier (0.64 eV) is still relatively small. The transition state energy (TS'6) between a'-V and a'-VI is now the highest (0.89 eV) in the reaction cycle fully consistent with experimental observations that the reaction cycle can be completed at relatively low temperature and that the second CO oxidation step is rate-determining.

After considering the two extreme cases with the catalyst bearing either no counterions or being completely neutralized by protons, it is important to note that the catalyst (de)protonation should be expected to be fast and highly dynamic throughout the reaction cycle.  $PW_{11}O_{39}Rh_1^{4-}$  may be bound to a varying number of different cations (e.g. H<sup>+</sup>, Na<sup>+</sup>, or K<sup>+</sup> as observed experimentally). We have demonstrated here that during the ESI-MS measurements, the structures are interconvertible, for example, the catalyst structure does not change significantly during the ionization procedure.

The selective catalytic oxidation of alcohols using environmentally benign oxidants in water is an important reaction adhering to the principles of green chemistry. Prompted by the widely known oxidation activities of POMs but general lack of mechanistic understanding, we employed ESI-MS to follow the oxidation of benzyl alcohol into benzaldehyde in aqueous hydrogen peroxide over solubilized SACs anchored on PW<sub>11</sub>O<sub>39</sub><sup>7-</sup>. In catalytic activity tests, TOF values (Table S3) indicate that among the 15 tested metals (Cr, Mn, Fe, Co, Ni, Cu, Zn, Rh, Pd, Ag, Ce, W, Pt, In, and Bi), PW<sub>11</sub>O<sub>39</sub>Fe<sub>1</sub><sup>4-</sup> seems to be the most relevant candidate regarding its comparably high activity (4.4  $h^{-1}$  at 70 °C) and low price. Conversions were generally kept below 25% to minimize the formation of benzoic acid. Good stability of the catalyst under reaction conditions was confirmed: no obvious decomposition of PW<sub>11</sub>O<sub>39</sub>Fe<sub>1</sub><sup>4-</sup> species occurred even after reaction at 70°C for 5 h (Figure S11).

ESI-MS revealed that equimolar mixtures of iron nitrate and POM led to the same active site as observed for  $PW_{11}O_{39}Rh_1^{4-}$  (Figure S12). The addition of benzyl alcohol to solutions containing PW<sub>11</sub>O<sub>39</sub>Fe<sub>1</sub><sup>4-</sup> led to the formation of the intermediate  $[Fe^{III_{1}^{3+}}(C_{7}H_{8}O)-(PW_{11}O_{39})^{7-}]^{4-}$  (b-I, calc.: 710.06, expt.: 710.05) (Figure S13) confirming the relatively weak binding of the substrate indicated by the low concentration of intermediate **b-I** relative to  $[Fe^{III}_{1}^{3+}-(PW_{11}O_{39})^{7-}]^{4-}$ (calc.: 683.30, expt.: 683.28).  $PW_{11}O_{39}^{7-}$  did not adsorb benzyl alcohol (Figure S14) thus providing evidence that the presence of Fe is essential for substrate binding. Other benzyl alcohol derivatives can also be adsorbed on PW<sub>11</sub>O<sub>39</sub>Fe<sub>1</sub><sup>4-</sup> and give rise to the intermediates  $[Fe_{1}^{III}, 3+(C_7H_7NO_3) (PW_{11}O_{39})^{7-}]^{4-}$  (calc.: 721.55, expt.: 721.57) for 4-nitrobenzyl alcohol and  $[Fe_{11}^{III_{3}}(C_{8}H_{10}O_{2})-(PW_{11}O_{39})^{7-}]^{4-}$  (calc.: 717.56, expt.: 717.57) for 4-methoxybenzyl alcohol. Differences in the adsorption strength of alcohol derivatives seem to follow the order benzyl alcohol <4-nitrobenzyl alcohol <4-methoxybenzyl alcohol based on the differences in the concentrations of the alcohol-bound intermediates after liquid phase adsorption (Figure S15).

Upon addition of hydrogen peroxide to the reaction mixture with benzyl alcohol and heating to 90°C, a range of other species assignable to various reaction intermediates appeared in the ESI-MS spectra (Figure 5a). Although the involvement of high valent Fe-oxo species in catalytic oxidation cycles is frequently reported, little evidence exists about the exact structure and dynamics of active sites even for homogeneous catalysts.<sup>[26]</sup> In the first step, the alcohol-bound  $PW_{11}O_{39}Fe_1^{4-}$  intermediate **b-I** adsorbed hydrogen peroxide upon removal of a proton, giving rise to intermediate  $\{[Fe_1^{III}_3^{+}(OOH^-)(C_7H_8O)-(PW_{11}O_{39})^{7-}] + 2H^+\}^{3-}$  (**b-II**, calc.: 958.07, expt. 958.07). At low collision energies of around 3 eV, benzyl alcohol desorbed from **b-II** and a new species with the composition  $\{[Fe_1^{III}_3^{+}(OOH^-)-(PW_{11}O_{39})^{7-}] + 2H^+\}^{3-}$  (calc.: 922.73, expt. 922.72) formed which was relatively stable

against further fragmentation to  $PW_{11}O_{39}Fe_1^{4-}$  (Figure 5b). With the addition of a proton, a water molecule was removed and the Fe<sup>V</sup> oxo complex was formed with the composition  $\{[Fe_{1}^{V,5+}(O^{2-})(C_{7}H_{8}O)-(PW_{11}O_{39})^{7-}] + H^{+}\}^{3-}$  (b-III, calc.: 952.41, expt.: 952.42). The existence and good catalytic oxidation performance of high valence Fe-oxo species on POMs has been proposed computationally and demonstrated experimentally before.<sup>[27]</sup> Fragmentation pattern of that species are indicative of a much less stable species with decomposition even occurring in the final product. The Fe SAC needs to form a transient radical Fe<sup>IV</sup> species **b-IV** which was not observed by ESI-MS presumably due to its low stability. Although the catalyst state binding the product benzaldehyde cannot be easily discerned after short term reaction, ESI-MS analysis of the reaction mixture after 5 h reaction at 70°C (Figure S16) clearly indicated another peak belonging to { $[Fe_{1}^{III}]^{3+}(C_7H_6O)-(PW_{11}O_{39})^{7-}] + H^+$ }<sup>3-</sup> (**b-V**, calc.: 947.08, expt.: 947.07). Fragmentation revealed a clean pattern with desorption of benzaldehyde at low fragmentation energies of 2 eV (Figure S17). Excellent overlap between the experimentally measured spectrum and the sum of the predicted isotope pattern was observed (Figure 5 a) confirming the presence of all the above-mentioned intermediates. Moreover, a detailed comparison of the experimentally observed and simulated spectra further corroborated the assignment of the intermediates (Figure S18). Combined, we determined a hitherto unknown reaction mechanism for the single atom catalyzed oxidation reaction using ESI-MS.

Relative concentrations of various intermediates change as a function of temperature (Figure 6a). Determining those concentrations during catalytic reactions leads to the identification of the temperature dependence of intermediates and thus the activation energy through the Arrhenius equation. Although apparent activation barriers for overall reactions are commonly determined, there is a lack of tools that can easily quantify reaction barriers between intermediates of single-atom catalysts without computational methods. For all three identified intermediates during  $PW_{11}O_{39}Fe_1^{4-}$ catalyzed oxidation reaction, concentrations have been determined at temperatures from 60 to 90°C (Figure S19). Arrhenius plots (Figure 6b-d) revealed that the activation barriers are  $121 \pm 2 \text{ kJ mol}^{-1}$ ,  $128 \pm 4 \text{ kJ mol}^{-1}$  and  $81 \pm$  $4 \text{ kJmol}^{-1}$  for the hydroperoxo, the oxo and the benzaldehyde-bound intermediates, respectively. This is consistent with the fact, that for the oxidation reaction of 4-nitrobenzyl alcohol, a less reactive substrate confirmed by previous studies,<sup>[28]</sup> only reasonable concentrations of the hydroperoxo and 4-nitrobenzaldehyde-containing intermediate were observed while the concentration of oxo intermediate seemed to be negligible (Figure S20). Inversely, the concentration of the oxo species was relatively high when 4-methoxybenzyl alcohol-a more reactive substrate for the alcohol oxidation reaction-was used (Figure S21), indicative of a reduced reaction barrier. The comparably high activation barrier for the Fe-oxo formation is in agreement with the relatively low activity at 70°C and the previously reported strong pH dependence for the Fe-oxo formation being unfavorable under acidic conditions.<sup>[29]</sup> For the oxidation of organic compounds with Fe-based catalysts involving the formation



**Figure 5.** ESI-MS spectra for the  $PW_{11}O_{39}Fe_1^{4-}$ -catalyzed benzyl alcohol oxidation. a) Overlap between the experimental spectra during the benzyl alcohol oxidation and a linear combination analysis of three key intermediates during the catalytic cycle. The three observed intermediates carrying different cations {xH<sup>+</sup>, yNa<sup>+</sup>, zK<sup>+</sup>} are indicated by different colors at their respective *m*/*z* values. Asterisks mark the locations where species occur that are also visible when iron nitrate and  $PW_{11}O_{39}^{7-}$  are mixed (see Figures S11 and S12 for more details). Fragmentation pattern for b) the Fe-oxo (**b-III**) and c) the Fe-hydroperoxo (**b-II**) intermediates are shown at increasing collision energies (in eV).

of Fe-oxo species has been shown to be dependent on the pH with lower values accelerating the reaction. This aligns well with our assumption that the oxo intermediate formation involves the addition of a proton. A decrease in the activity was previously observed, however, when the pH dropped to values below 4, presumably due to the inversely pH-dependent hydroperoxo intermediate formation becoming ratedetermining.<sup>[30]</sup> Combining the observed intermediates with the determined activation barriers, we propose a reaction mechanism as shown in Figure 7 where Fe undergoes oxidation state changes involving Fe<sup>III</sup>, Fe<sup>IV</sup>, and Fe<sup>V</sup>, and the formation of the Fe-oxo species is rate-determining. A quantitative correlation between ESI-MS and catalytic activity as well as the analysis of the reaction mechanism for benzyl alcohol oxidation by PW<sub>11</sub>O<sub>39</sub>Pd<sub>1</sub><sup>5-</sup> are provided in the SI (Figures S22–28 and accompanying text). We found that the Pd-based SAC follows an entirely different reaction mechanism at least partly due to the lack of Pd oxo formation.

#### Conclusion

This study exemplifies the advantages of ESI-MS to examine molecularly defined SACs containing different active metal species in solution-phase oxidation reactions. Despite the time passed between injection and analysis, ESI-MS proves effective for the investigation of the structure and structural evolution of solubilized, POM-supported SAC sites during reaction. Atomically precise active sites with welldefined composition were created and identified, and their dynamics during catalytic cycles were determined. Taking advantage of the soft ionization ability of the electrospray preserving various reaction intermediates, and the high m/zresolution and sensitivity of the mass detector, even low concentrations of unstable intermediates were separated from complex reaction mixtures and analyzed thoroughly. As such, the reaction mechanism, and more importantly the stoichiometry of actives sites as well as key substrate-active site



**Figure 6.** a) ESI-MS spectra for the PW<sub>11</sub>O<sub>39</sub>Fe<sub>1</sub><sup>4-</sup>-catalyzed oxidation reaction at increasing temperatures from bottom to top: 50, 60, 70, 80 and 90 °C. Determination of the reaction barriers for the formation of different intermediates estimated by Arrhenius plots for b) the Fehydroperoxo **b-II**, c) the Fe-oxo **b-III** and d) species **b-V** containing benzaldehyde. Reaction conditions for **a**-**d**:  $V_{total} = 0.02$  L deionized water with  $C_{PW_{11}O_{39}} = 0.5$  mM,  $C_{Fe} = 0.5$  mM,  $C_{BnZOH} = 25$  mM, and  $C_{H_2O_2} = 50$  mM at T = 60 °C – 90 °C under magnetic stirring at 800 rpm. ESI-MS parameters are described in the methods section.



**Figure 7.** The proposed mechanism of the  $PW_{11}O_{39}Fe_1^{4-}$ -catalyzed alcohol oxidation reaction is shown. The experimentally determined and simulated masses, as well as the experimentally determined reaction barriers for the most relevant steps are shown. Oxidation states of Fe are indicated by different colors. All of that information was obtained from ESI-MS measurements.

complexes in the full catalytic cycle, were determined for several SACs in both CO and alcohol oxidation. Together with the unique POM-based SACs, this approach is particularly suitable for DFT calculations revealing even further insights into reaction mechanisms.

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With the wide variety of POM structures and other soluble metal oxides,<sup>[17,31]</sup> as well as the unlimited number of atomically dispersed metals across the periodic table, we believe that this mass spectrometry approach furnishes a general platform for the elucidation of structure–activity correlations of SACs in solution phase reactions with high precision. Beyond mechanistic analyses, ESI-MS proves powerful in the rapid investigation of large catalyst libraries in minimal amounts of time,<sup>[32]</sup> while POM-based SACs offer a potent platform for such high-throughput studies in the future.

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#### Conflict of interest

The authors declare no conflict of interest.

Keywords: heterogeneous catalysis · homogeneous catalysis · mass spectrometry · polyoxometalates · reaction mechanisms

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