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Adsorption and Activation of Water on Cuboctahedral Rhodium and Platinum Nanoparticles

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Abstract

Rh and Pt are widely used as the components in heterogeneous catalysts for multiple industrial applications. Since the metals are typically in the form of nanoparticles in real catalysts, it is important to carefully select models for the computational prediction of the catalytic properties. Here, we report a first-principle study on the water activation, an important step in numerous catalytic reactions, using the finite-size Rh and Pt nanoparticle models and compare them to the extended surface models. We show that regardless of the model, adsorption and activation of water is practically identical for both metals, whereas the dissociation is energetically more favorable on Rh. The experimentally observed difference must be thus attributed to stronger interaction of dissociated water with the metal surfaces and/or to the presence of the oxide support. Through a selection of descriptors, we demonstrate that the extended surface models cannot fully represent the atomic and electronic structures of the small nanoparticles of less than 2 nm in size.

1 Introduction

Heterogeneous transition metal catalysts are widely exploited nowadays in major industrial processes to obtain renewable fuels and sustainable chemicals. In particular, Rh and Pt have been found active in the water–gas shift and multiple reforming reactions, as highlighted in the recent reviews.\textsuperscript{1–6} Activation of water has been reported to play a key role in the networks of the aforementioned reactions, in some cases being acknowledged as the rate-limiting step for the water–gas shift cycle.\textsuperscript{7,8} Since the industrial catalysts operate at the nanoscale regime, deeper atomic-level understanding of the water activation in the heterogeneous transition metal-based catalytic systems could provide important information for systematization and rational design of advanced catalysts with higher activity and selectivity.

Along with other transition metals, adsorption and dissociation of water on the surfaces of Rh and Pt have been computationally addressed by numerous research groups during
the last decade.\textsuperscript{9–18} Calculation reports have been primarily focused on representing metal surfaces with flat and stepped surface slabs to mimic different structural features of realistic nanoparticles at relatively low computational expenses. The flat (111) and (100) surfaces of Rh and Pt have been shown to adsorb and activate water equally well.\textsuperscript{11} The (211) and (221) stepped surfaces, mimicking the edges of realistic nanoparticles, have been found to be more reactive towards water, which is likely due to the lower coordination of the surface sites.\textsuperscript{10,11} These findings are in line with the computational reports by Calle-Vallejo et al.\textsuperscript{19–21} and Fajín et al.,\textsuperscript{12} where the authors have demonstrated the stronger tendency of water to bind the undercoordinated edge and corner sites over the facets using the finite-size Pt nanoparticles. Moreover, the degree of interaction has been observed to depend on the size of Pt nanoparticles,\textsuperscript{12} making the extended surface models insufficient in describing the finite-sized facets of small nanoparticles.

To our knowledge, computational studies on water activation by Pt and, especially, Rh nanoparticles are limited, and the comparison between finite-sized Rh and Pt nanoparticles in their tendencies to bind and activate water remains incomplete. Here, we report a first-principles study, where we seek answers to the following questions: (1) what is the difference between Rh and Pt in their activity towards adsorption and activation of water, especially, in the nanoscale regime; (2) how well do the extended surfaces represent the finite-size features of small Rh and Pt nanoparticles; and (3) are the electronic structure and surface coordination sufficient descriptors that could determine quantitatively the ability of Rh and Pt metals to bind and activate water?

2 Computational Details

Spin-paired periodic and cluster density functional theory (DFT) calculations were performed using the grid-based projected augmented wave approach as implemented in the GPAW.\textsuperscript{22,23} The Perdew–Burke–Ernzerhof (PBE) functional\textsuperscript{24,25} was employed throughout the study.
The wave-functions were represented on a direct lattice grid with the maximal grid density of 0.20 Å. Atomic projected augmented wave (PAW) setups were employed to every atom. The core electrons were treated in a frozen-core approximation in the case of Rh, Pt, and O atoms. The PAW setups included 15 valence electrons for Rh, 16 valence electrons for Pt, 6 valence electrons for O, and 1 valence electron for H. The reciprocal space was sampled using the (6 × 6 × 1) Morkhorst–Pack-type mesh of $k$-points in the 2D-periodic calculations, while the non-periodic systems were treated at the Γ-point alone. The optimized lattice constants of 3.857 Å and 3.981 Å were used for the bulk Rh and Pt metals, respectively. Extended surfaces were represented by either (2 × 2) flat or (3 × 2) stepped slabs of four atomic layers thick, with the two bottom layers fixed to the original positions. Cuboctahedral nanoparticles were generated via the Wulff’s construction as implemented in the ASE code. The required values for the surface energies were taken from the Crystalm database. Two different size models included 55 and 147 metal atoms, resulting in a set of nanoparticles with the diameter of approximately 1.0–1.7 nm. Only the atoms in the topmost layers were allowed to relax. Complete relaxation of the nanoparticle models results in the changes in the considered energy quantities of no more than ±0.05 eV and may be neglected, thereby reducing the computational burden without significant loss in accuracy. Extended surface models were surrounded by at least 5 Å of vacuum below and 10 Å above the slabs. The nanoparticle models were surrounded by at least 7.5 Å of vacuum in each direction. Each structure was minimized until the maximal residual force was below 0.05 eV Å$^{-1}$. The transition states of water activation were located using the constraint optimization technique with a 0.025 Å step for the cleaving bond and confirmed by the presence of a single imaginary mode that corresponds to the reaction coordinate. The gas-phase water molecule was calculated in a 12.8 × 12.8 × 12.8 Å non-periodic computational cell on a 64 × 64 × 64 points uniform grid (grid spacing 0.20 Å). The vibrational analysis was thereby performed only for the adsorbate atoms. In the dissociated states, OH and H species were separated as far away from each other as possible by placing them either into individual unit cells in the case of the extended
surfaces or at the diametrically opposite sides of the nanoparticles to exclude or minimize their mutual interactions.

3 Results and Discussion

Figure 1: Structural models exploited throughout the study (bulk metal atoms in dark blue, surface metal atoms in light blue, unit cells of the extended surfaces in green). In the designation, T stands for the terrace, H for the hollow fcc, E for the edge, and C for the corner sites; T–T and E–E stand for the bridging sites (represented by the dots) at the terraces and edges, respectively.

Figure 1 represents all structural models exploited throughout the study with the relevant for discussion surface sites highlighted. The cuboctahedral metal nanoparticles are terminated by two families of facets, namely hexagonal (111) and square (100), which differ in the coordination of metal atoms. The former features 9-coordinated 111T sites and can be thus mimicked with the extended (111) surface, whereas the latter features 8-coordinated 100T sites and is mimicked with the extended (100) surface. The edges between the (111) and (100) facets consist of 7-coordinated E sites, and we represent those with the step edge.
sites on the extended (211) surface. The nanoparticle edges further intersect to form the C sites, at which the metal atoms are 5-coordinated. The C sites can be, in principle, represented by high-index extended surfaces (as shown, for example, in Ref. 29); however, this representation lies beyond the scope of the current study.

Although the surface sites in the extended surfaces have identical coordination numbers as the corresponding sites in the nanoparticles, the \textit{xy}-periodic surfaces lack mimicking finite size effects of the structural motifs in the nanoparticles. To overcome such a misalignment, Calle-Vallejo et al.\textsuperscript{19} suggested a descriptor, the so-called generalized coordination number, \( \overline{\text{CN}} \), which is calculated as

\[
\overline{\text{CN}} = \frac{\sum \text{CN}_i n_i}{\text{CN}_{\text{max}}},
\]

where \( n_i \) is the number of atoms with the conventional coordination number \( \text{CN}_i \) located in the first coordination sphere of the surface site of interest and \( \text{CN}_{\text{max}} \) is the maximal value of the coordination number in the system, which equals 12 for the considered “atop” sites on the fcc metals. Table 1 gives the generalized coordination numbers for the relevant adsorption sites in the studied systems (Figure 1). The values of \( \overline{\text{CN}} \) for the sites in the nanoparticles approach those for the corresponding sites in the extended surfaces with the increasing size of the nanoparticles, making the generalized coordination number a more descriptive quantity, which reflects better the size-dependent nature, as has been successfully shown by the authors in a comprehensive evaluations of the interaction of water and other small species with various sites of truncated octahedral Pt nanoparticles.\textsuperscript{19–21}

We evaluated adsorption, activation, and dissociation of molecular water on the sites specified in Figure 1. Water was found to bind each examined site preferably in the “atop” position (Figure 2). The transition states of activation, illustrated in Figure 3, were located at the region of 1.5–1.7 Å for the length of the cleaving O–H bond. The adsorption, \( \Delta E \), and transition state, \( \Delta E^\dagger \), energies (Table 2) were calculated with respect to the infinitely separated species as \( \Delta E^{(4)} = E^{(1)}_\Sigma - (E_S + E_W) \), where \( E^{(1)}_\Sigma \), \( E_S \), and \( E_W \) are the DFT energies
Table 1: Generalized coordination numbers of the relevant sites on the nanoparticles and extended surfaces.

<table>
<thead>
<tr>
<th>site</th>
<th>site</th>
<th>M&lt;sub&gt;55&lt;/sub&gt;</th>
<th>M&lt;sub&gt;147&lt;/sub&gt;</th>
<th>M&lt;sub&gt;∞&lt;/sub&gt;</th>
<th>CN&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>111T</td>
<td>—</td>
<td>6.50</td>
<td>7.50</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>100T</td>
<td>6.33</td>
<td>6.50</td>
<td>6.66</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>4.66</td>
<td>5.00</td>
<td>5.50</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>3.33</td>
<td>3.33</td>
<td>—</td>
<td>5</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> See Figure 1 for notation of the adsorption sites.

<sup>b</sup> Conventional coordination numbers.

of the metal–water (transition state) complex, plain metal, and gas-phase water molecule, respectively. Table 3 shows the energy of the dissociated state, ΔE<sup>D</sup>, which was calculated similarly for the nanoparticles and as ΔE<sup>D</sup> = (E<sub>Σ</sub><sup>OH</sup> + E<sub>Σ</sub><sup>H</sup>) − (2E<sub>S</sub> + E<sub>W</sub>) in the case of the extended surfaces, where E<sub>Σ</sub><sup>OH</sup> and E<sub>Σ</sub><sup>H</sup> are the respective DFT energies of the metal–OH and metal–H complexes. In such formulation, more negative values indicate stronger interaction and/or greater stability.

Table 2: Adsorption (∆E) and transition state (∆E<sup>‡</sup>) energies (in eV) and corresponding imaginary frequencies (ω, in cm<sup>−1</sup>) for the most favorable configurations of water at the representative sites on Rh and Pt nanoparticles and extended surfaces.

<table>
<thead>
<tr>
<th>site</th>
<th>model</th>
<th>∆E (Rh)</th>
<th>∆E (Pt)</th>
<th>∆E&lt;sup&gt;‡&lt;/sup&gt; (Rh)</th>
<th>ω (Rh)</th>
<th>∆E&lt;sup&gt;‡&lt;/sup&gt; (Pt)</th>
<th>ω (Pt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>111T</td>
<td>M&lt;sub&gt;∞&lt;/sub&gt;</td>
<td>−0.34</td>
<td>−0.29</td>
<td>+0.53</td>
<td>450.0&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.59</td>
<td>366.2&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>M&lt;sub&gt;147&lt;/sub&gt;</td>
<td>−0.19</td>
<td>−0.16</td>
<td>+0.72</td>
<td>610.6&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.80</td>
<td>368.1&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td>100T</td>
<td>M&lt;sub&gt;∞&lt;/sub&gt;</td>
<td>−0.34</td>
<td>−0.25</td>
<td>+0.55</td>
<td>488.3&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.55</td>
<td>236.6&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>M&lt;sub&gt;147&lt;/sub&gt;</td>
<td>−0.24</td>
<td>−0.23</td>
<td>+0.61</td>
<td>456.3&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.56</td>
<td>230.8&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>M&lt;sub&gt;55&lt;/sub&gt;</td>
<td>−0.22</td>
<td>−0.19</td>
<td>+0.56</td>
<td>428.8&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.57</td>
<td>324.8&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td>E</td>
<td>M&lt;sub&gt;∞&lt;/sub&gt;</td>
<td>−0.39</td>
<td>−0.37</td>
<td>+0.44</td>
<td>446.4&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.39</td>
<td>279.9&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>M&lt;sub&gt;147&lt;/sub&gt;</td>
<td>−0.37</td>
<td>−0.36</td>
<td>+0.36</td>
<td>500.1&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.30</td>
<td>499.8&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>M&lt;sub&gt;55&lt;/sub&gt;</td>
<td>−0.38</td>
<td>−0.34</td>
<td>+0.42</td>
<td>306.0&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.41</td>
<td>307.7&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td>C</td>
<td>M&lt;sub&gt;147&lt;/sub&gt;</td>
<td>−0.56</td>
<td>−0.66</td>
<td>+0.26</td>
<td>699.2&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.19</td>
<td>305.7&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>M&lt;sub&gt;55&lt;/sub&gt;</td>
<td>−0.57</td>
<td>−0.59</td>
<td>+0.20</td>
<td>542.6&lt;sup&gt;i&lt;/sup&gt;</td>
<td>+0.10</td>
<td>632.4&lt;sup&gt;i&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> See Figure 1 for notation of the adsorption sites.

Molecular adsorption of water was found to be practically equivalent (within 0.10 eV) for the corresponding sites on Rh and Pt. The adsorption energies on the extended surface models indicate that water favors the E site on the stepped (211) surfaces over the terrace sites on the (111) and (100) surfaces in line with the previous computational reports. 9–18
Figure 2: Adsorption modes of molecular water on each studied surface site (bulk metal atoms in dark blue, surface metal atoms in light blue, oxygen in red, hydrogen in white, unit cells of the extended surfaces in green). See Figure 1 for the notation of the surface sites.

Table 3: Dissociated state energies ($\Delta E^D$, in eV) for the most favorable configurations of dissociated water on Rh and Pt nanoparticles and extended surfaces.

<table>
<thead>
<tr>
<th>model</th>
<th>$\Delta E^D$ (Rh)</th>
<th>$\Delta E^D$ (Pt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M (111)</td>
<td>−0.28</td>
<td>+0.30</td>
</tr>
<tr>
<td>M (100)</td>
<td>−0.60</td>
<td>−0.28</td>
</tr>
<tr>
<td>M (211)</td>
<td>−0.82</td>
<td>−0.52</td>
</tr>
<tr>
<td>M_{147}</td>
<td>−0.93</td>
<td>−0.73</td>
</tr>
<tr>
<td>M_{55}</td>
<td>−0.73</td>
<td>−0.66</td>
</tr>
</tbody>
</table>
Figure 3: Structures of the transition state of water dissociation for all studied structural models (bulk metal atoms in dark blue, surface metal atoms in light blue, oxygen in red, hydrogen in white, unit cells of the extended surfaces in green).
Figure 4: Dependencies of the molecular adsorption of water (a) on the center of the d-band projected onto the corresponding sites and (b) on the generalized coordination numbers (right). Coding: Rh in red, Pt in blue, nanoparticles with circles, extended surfaces with triangles. The trendlines and correlation coefficients are given for the nanoparticle models only.

Adsorption energies on the nanoparticle models are consistent with those calculated for the extended surfaces, with an exception for the 111T site. The difference may be attributed to the incomplete formation of the (111) facets on the nanoparticles in the selected size regime. Overall, the highly undercoordinated C sites of the nanoparticles bind water the strongest, as intuitively expected, followed by the E, 100T, and 111T sites in the same order as for the extended surface models.

In the case of the nanoparticle models, the adsorption energies are found to correlate linearly ($r^2 > 0.89$) with the d-band center projected onto the corresponding adsorption sites (Figure 4a). Moreover, we observe strong linear dependencies ($r^2 > 0.94$) of the adsorption energies on the generalized coordination numbers (Figure 4b), which indicates that the coordination of the surface sites plays a dominant role in determining the strength of interaction. Calle-Vallejo et al. reported similar strong correlations for the truncated octahedral Pt nanoparticles of 0.7–1.7 nm in size. The data obtained for the extended surface models, however, dropped out from the relationships, suggesting that these models are unable to fully mimic the facets of the cuboctahedral Rh and Pt nanoparticles in the selected size regime.
As in the case of the adsorption energies, the difference in the calculated transition state energies between Rh and Pt is found to be negligibly small (within 0.10 eV). The most stable transition states on the extended surface models are identified for the E sites on the stepped (211) surfaces, followed by the terrace sites on the (111) and (100) surfaces. The transition state energies calculated for the nanoparticle models were consistent with the extended surface models, again with an exception of the 111T site, which is likely due to incomplete formation of the (111) facets on the Rh\textsubscript{147} and Pt\textsubscript{147} nanoparticles. The most stable transition states on the nanoparticle models were observed at the C sites, followed by the E, 100T, and 111T sites.

![Figure 5](image_url)

**Figure 5:** Dependencies of the transition state energy of water activation (a) on the center of the d-band projected onto the corresponding sites and (b) on the generalized coordination numbers. Coding: Rh in red, Pt in blue, nanoparticles with circles, extended surfaces with triangles. The trendlines and correlation coefficients are shown for the nanoparticle models only.

Similarly to the adsorption, the transition state energies calculated for the nanoparticle models demonstrate strong dependencies ($r^2 > 0.85$) on the generalized coordination numbers and moderate dependencies ($r^2 > 0.80$) on the projected d-band centers (Figure 5). Generalized coordination numbers of the sites occupied by OH in the transition state complexes were used in the analysis. The observed trends result in the fact that our data show strong ($r^2 > 0.85$) Brønsted–Evans–Polanyi-type (BEP) transition state scaling relation-
Figure 6: Transition state scaling for the water adsorption and activation. Coding: Rh in red, Pt in blue, nanoparticles with circles, extended surfaces with triangles. The trendlines are shown for the nanoparticle models only.

Figure 6 shows the transition state scaling for the water adsorption and activation. The coding includes Rh in red, Pt in blue, nanoparticles with circles, and extended surfaces with triangles. The trendlines are displayed only for the nanoparticle models.

The analysis of the complete reaction profiles (Figure 7) revealed that the difference between Rh and Pt metals is hidden behind the structures of the dissociated states of water. As the data in Table 3 show, the OH and H species present generally more stable configurations on Rh, making the water activation there thermodynamically more favorable. In terms of energy, the difference was found to be still insignificant (<0.10 eV) in the case of the Rh$_{55}$ and Pt$_{55}$ nanoparticles; however, it increased to approximately 0.20 eV for the M$_{147}$ nanoparticles and further to >0.30 eV values for the extended surfaces. Among the extended surfaces, the stepped (211) surfaces give the closest values of $\Delta E^D$ to those for the nanoparticles.

The difference between Rh and Pt and the best performance of the (211) extended surfaces may be linked to the different preference of the adsorption sites by the OH and H...
Figure 7: Complete reaction profiles of the adsorption and activation of water on Rh and Pt nanoparticles and extended surfaces. Color coding: 111T in red, 100T in green, E in yellow, and C in blue. See Figure 1 for the notation of the surface sites.
Figure 8: Adsorption modes of dissociated water on each studied structural model (bulk metal atoms in dark blue, surface metal atoms in light blue, oxygen in red, hydrogen in white, unit cells of the extended surfaces in green).
species. In the case of Rh, both hydroxyl and hydrogen favor the E–E bridging sites of
the nanoparticles and the corresponding E–E bridging sites at the step edges of the (211)
extended surfaces. On the flat surfaces of Rh, both OH and H prefer adsorption to the
T–T bridging sites on the (100) surface and to the H sites on the (111) surface; however,
these configurations are higher in energy. On Pt, the OH and H species preferably bind to
the C and E–E bridging sites of the nanoparticles, respectively, while on the stepped (211)
surface, both species adsorb on the E–E bridging sites, which is the closest approximation
of the nanoparticle, considering that there are no C sites on any of the extended surfaces.
High-index extended surfaces might have thus provided necessary kink sites to mimic the
corners of the nanoparticles. On the flat surfaces, the hydroxyl favors the T–T bridging
sites, whereas the hydrogen prefers the T–T bridging site on the (100) surface and the H site
on the (111) surface. Figure 8 summarizes the most favorable configurations of dissociated
water on each considered structure model. These findings are qualitatively in line with the
experimental observations, which suggest that the water activation is less favorable on Pt
due to the low thermodynamical stability of PtO$_x$.\textsuperscript{34} Moreover, a systematic evaluation of Pt
catalysts supported on reducible oxides has demonstrated that the presence of the support
significantly improves performance of Pt in the water–gas shift.\textsuperscript{35}

4 Conclusions

In summary, we computationally examined and compared water adsorption, activation, and
dissociation by the Rh and Pt metals. We modeled the metal surfaces using the M$_{55}$ and
M$_{147}$ cuboctahedral nanoparticles as well as the extended slabs. For the nanoparticle models,
the adsorption of water was found to strongly depend on the nature of the surface sites in
terms of coordination and the structure of d-band, resulting in strong linear dependencies of
the adsorption energies on the considered descriptors, i. e. generalized coordination numbers
and the projected centers of the d-band. The transition state energies showed excellent
scaling relationships with the adsorption energies as well as the strong correlations with the
descriptors, enabling prediction of these values at relatively low computational expense. For
the extended surface models, the aforementioned relationships were found to be significantly
weaker, indicating that such models cannot represent the finite-size effects on the electronic
and atomic structures of the cuboctahedral Rh and Pt nanoparticles in the chosen size
regime (<2 nm) and the reaction. Nonetheless, the extended surface models do mimic well
the energetics of the water activation, and the difference between Rh and Pt metals in their
tendencies to adsorb and activate water was overall small (<0.10 eV) using the different
structural models. Regardless of the model, the dissociation of water was found to be
thermodynamically more favorable on Rh (>0.10 eV depending on the model), which is likely
due to the different affinity of the OH and H species to the Rh and Pt metals. Experimentally
observed inequalities may be thus linked to the higher stability of the OH and H species on
Rh compared to Pt and/or to the use of the oxide supporting materials, upon interaction
with which, metals nanoparticles alter their shapes and, consequently, atomic and electronic
structures. Variety of the interface sites, generated thereby, may have different properties
compared to the sites in the bare metal nanoparticles.

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