Solvation Effects on Methanol Decomposition on Pd(111), Pt(111), and Ru(0001)

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ABSTRACT: Solvation is crucial in many chemical and electrochemical processes related to alcohol conversion on metal surfaces. Particularly, understanding the dehydrogenation mechanism of methanol on solvated Pd, Pt, and Ru surfaces could allow the design of efficient methanol fuel cells. The large computational cost related to adopting an explicit solvation approach into density functional theory can be reduced drastically by using implicit solvation methods. In this study, we use our recently developed continuum solvation model (MGCM) to elucidate the minimum number of explicit water molecules to add to the solvated methanol/metal surface systems to reproduce experimental data with an optimized balance between time and reliability. Our results stress the importance of adding two explicit water molecules, especially for the case of Ru surfaces. For this later system, we provide a first insight into the decomposition mechanism of methanol using first-principles calculations. Our predictions can be then a useful reference for future studies that aim at designing more efficient heterogeneous catalysts with solvents.

INTRODUCTION

Biomass-derived compounds have received increasing attention in recent years as an alternative source of energy to fossil fuels.¹ Commonly biomass resources contain significant amounts of water. To avoid the complexity of long carbon polyalcohols, studies have been centered in surrogates, small alcohols, that already are of technological interest. Among such compounds, methanol, the simplest alcohol, has many advantages as energy carriers due to its high hydrogen content, availability, low cost, and ease of transport and storage. The use of methanol as a fuel is based on the so-called fuel-cell technology with potential applications in portable power and electric vehicles.² Direct methanol fuel cells (DMFCs) are used to decompose methanol to produce electricity. On the anode side of the DMFC methanol is oxidized on a metal catalyst such as platinum (Pt), palladium (Pd), or ruthenium (Ru) in an aqueous environment.³ Water plays then a key role on the surface chemistry in DMFCs as it interacts with the adsorbate through hydrogen bonds, short-, and long-range interactions, and it modifies the number of available active sites on the metal catalyst. Therefore, should an active and selective catalyst be designed, the effect of the environment is to be considered carefully.

Density functional theory (DFT) has been employed extensively in methanol decomposition on metal surfaces.³ Studies of methanol adsorption on metal/vacuum and metal/water interfaces suggest that the methanol dehydrogenation pathway depends strongly on the environment. While experiments in ultrahigh vacuum (UHV) show that the adsorption of methanol on Pt and Ru surfaces starts with the O–H scission,⁴⁵ theoretical studies under the same conditions lead to different reaction pathways. For instance, Desai et al.⁶ studied the adsorption of methanol over Pt(111) surfaces in vacuum and observed that dehydrogenation pathways starting via the activation of the C–H bond are favored. A similar pathway for methanol decomposition on Pt(111) was suggested by Greeley and Mavrikakis through microkinetic modeling based on first-principles results.⁷ DFT-based results for the case of methanol adsorbed on Pd, Pt, and Ru surfaces in vacuum show that several C–H bond breakings occur before O–H scission. However, the formation of adsorbed methanol clusters is favored, and this promotes the O–H bond breaking.⁸ A good agreement is found between the results obtained from experiments and theoretical studies when aqueous methanol decomposition is considered. In this case, the initial step in the reaction pathway is the C–H bond cleavage.⁹¹¹ Neurock et al.¹² studied the dehydrogenation pathway from CH₃OH to CO on Pt(111) in the presence of explicit water molecules through DFT calculations. All dehydrogenation steps were more exothermic than in vacuum. In this case, CH₃OH intermediates are stabilized by hydrogen bonding with nearby H₂O and H₂O⁺ species, and then the C–H bond cleavage is favored over the O–H bond breaking during the reaction path. Similar results were reported by Hartnig and Spohr¹³ who observed that O–H bond dissociation does not occur when methanol is hydrogen-bonded to a water molecule. However,
these studies mainly consider explicit molecules, and long-range solvent effects were not introduced.

Considering solvation effects in first-principles calculations is, nevertheless, a delicate issue. Explicit solvent modeling involves long-time scale molecular dynamics (MD) simulations as physical quantities should be averaged to compute the thermodynamic properties of interest. This results in an unaffordable computational cost for most of the systems of interest. However, the hydrogen bond stabilization of the adsorbed methanol comes mainly from the water molecules surrounding it and from those solvating the metal surface. In fact, water molecules located in the bulk, far away from the adsorption site, will have a residual effect on the methanol decomposition pathway. Thus, an alternative approach is to treat the solvent as a structureless continuum medium characterized by the value of its dielectric permittivity, while the solute and the nearby water molecules are treated through quantum mechanics. This strategy, known as implicit solvation, reduces drastically the computational cost of the calculations without affecting the accuracy of the computed physical quantities. In this context, we have recently developed a continuum solvation model into the Vienna ab Initio Simulation Package (VASP), a plane-wave based electronic structure code, VASP-MGCM (VASP-Multigrid Continuum Model). The model solves the electrostatic problem through a multigrid solver and has been tested for a large number of isolated and adsorbed molecules on metal surfaces with a good agreement between experimental and simulation data. The key point in this type of calculations is to determine the number of solvent molecules to add to the simulation box. On the one hand, if no explicit solvent molecules are considered, the solute adsorption geometry can differ from the true configuration as the short-range polarization of the adsorbate is not properly described. This leads to wrong estimates for thermodynamic...
and kinetic parameters. On the other hand, adding a large number of solvent molecules results in a high computational cost and does not necessarily improve substantially the accuracy of the results with respect to the case where we consider a few number of explicit molecules.\textsuperscript{21} Several studies have adopted a combined explicit/implicit solvation approach to study different reactions on Pt(111) and its alloys. This is the case of the oxygen reduction reaction (ORR),\textsuperscript{29,30,32} the adsorption of H, O, and OH,\textsuperscript{29} and the H\textsubscript{2}O interactions with CO and sugar alcohols.\textsuperscript{33,34} Although these studies provide relevant information about the reaction mechanisms of organic molecules on Pt(111) under solvation, the suitability of the explicit/implicit solvent approach has not been extensively studied for other metal surfaces like Pd(111) or Ru(0001).

In the present work, we have studied the effect of the solvent on the decomposition of methanol to formaldehyde on Pd(111), Pt(111), and Ru(0001) through DFT calculations. Our aim is not only to report the mechanisms of methanol dehydrogenation on metal surfaces under solvation, thus closer to the DMFC working conditions, but more importantly, to identify a minimum system setup to describe solvation effects accurately at the lowest computational cost. First of all, we demonstrate that the addition of two water molecules on the Pd and Pt surfaces permits to reproduce experimental and simulation results where the methanol C–H bond breaking is the preferred first dehydrogenation step under solvation. In addition, we report for the first time the mechanism that governs methanol dehydrogenation on Ru surfaces in an aqueous environment. This mechanism, which can only be observed when explicit water molecules are bonded to the adsorbate, consists on a proton transfer from the hydroxyl group to nearby water molecules that dissociate afterward. Experimental and computational studies involving methanol dehydrogenation on Ru(0001) have only been done in vacuum.\textsuperscript{35,36,37} Our results are a step forward in describing the methanol decomposition pathway at the metal-water interface and can serve as a reference for further studies.

\section*{THEORETICAL METHODS}

Simulated Systems. For each metal we consider two different situations: (1) no water molecules are added to the simulation box and (2) two water molecules are bonded to both methanol and the metal surface (see Figure 1a–f). For simplicity, we refer to these systems as CS (clean surface) and S2W (surface with two water molecules), respectively. The configuration of the two molecules for the later system is a chain so that the HOMO (basicity) of water is close enough to a water cluster value.\textsuperscript{29} In both cases, the systems are also immersed in a continuum water medium simulated using the VASP-MGCM. As the water dissociation ratio on Ru(0001) is about 40 to 50%,\textsuperscript{29} one water molecule is dissociated on the Ru S2W (Figure 1f). For Pt, we have also studied the decomposition pathway of methanol when it belongs to a solvated explicit ice-like water ring as that described in ref 29 (Figure 1g), both in vacuum and under solvation. We refer to this system as SSWR (surface with a five-water ring). The SSWR configuration is present in long-time simulations of liquid water (Born–Oppenheimer molecular dynamics, BOMD, at 300 K),\textsuperscript{29} although other less structured 7-, 6-, and 5-membered rings are also present at the interface. At the same time, we have extracted a snapshot from the BOMD in ref 29 for the system containing the Pt surface (labeled as Pt4 in that reference) solvated by 72 water molecules. We replaced a flat water molecule by methanol, preserving the hydrogen bonds that the replaced molecule had with its neighbors. Then, the adsorbate was relaxed along its vicinal water molecules (see Figure 1 h). This system is called S71W (surface with 71 water molecules). For the SSWR, the water configuration is reminiscent of that found around the methanol molecule in the S71W. Notice that unlike a water ring model coming from ice, the SSWR presents some molecules with the dipole moment slightly tilted with respect to the Pt surface. The SSWR retains then a certain degree of liquid-like behavior in the water configuration. Both the results for the SSWR and the S71W serve as a benchmark to assess the accuracy of the two strategies described above (CS and S2W).

Computational Details. DFT calculations were performed using the Vienna ab Initio Simulation Package (VASP),\textsuperscript{16} version 5.3.3, with the Perdew–Burke–Ernzerhof (PBE) GGA exchange-correlation functional.\textsuperscript{30} Dispersion interactions were calculated through the semiemirical Grimme’s DFT-D2 method,\textsuperscript{31} with our reparameterized values for the metal atoms.\textsuperscript{32} Core electrons were described using the projector augmented wave (PAW) formalism,\textsuperscript{33} while valence electrons were represented by plane waves with a cutoff energy of 450 eV. Solvent effects were treated through the VASP-MGCM. In this case, the value of the dielectric permittivity $\epsilon$ in the space is calculated using the Fattetbert and Gygi functional form:\textsuperscript{34}

$$
\epsilon(r) = \frac{\epsilon_0 - 1}{1 + (\rho_d(r)/\rho_0)^{\beta}}
$$

Here, $\rho_d$ stands for the electronic charge density, $\epsilon_0$ is the permittivity of the bulk phase of the solvent (78.5 for water), and $\rho_0$ and $\beta$ are parameters controlling the shape of $\epsilon$. The adopted values for these two parameters are $\beta = 1.7$ and $\rho_0 = 6 \times 10^{-4}$ a.u. The electronic convergence threshold was set to $10^{-7}$ and $10^{-6}$ eV for calculations involving metal surfaces and isolated molecules, respectively. In the case of gas-phase systems and the CS, the initial configurations were relaxed until forces acting on ions were lower than 30 meV/Å (15 meV/Å for solvated and unsolvated isolated molecules). In this case, a conjugate-gradient (CG) algorithm\textsuperscript{35} was used to relax the ions into their instantaneous ground state. When explicit water molecules were adsorbed on the metal surfaces, the convergence criterion for the forces was slightly changed to 50 meV/Å and a quasi-Newton algorithm\textsuperscript{36} was used to relax the ions for solvated systems. The choice of such algorithm was based on the fact that, when considering a CG algorithm for ion relaxation, we observed that some water molecules tended to rotate too much due to the interaction with the implicit aqueous environment. The CS and S2W metal surfaces were modeled by a four-layers $2\sqrt{3} \times 2\sqrt{3} - R30^\circ$ slab separated by at least 13 Å of vacuum along the z-axis, with dipole corrections included along this direction. For the S5WR and the S71W, we used a $3\sqrt{3} \times 3\sqrt{3} - R30^\circ$ reconstruction. The two topmost layers were relaxed, while the bottom ones were fixed to the bulk distances. The lattice parameters for Pd, Pt, and Ru were 3.939, 3.968, and 3.629 Å, respectively, and $[c/a]_{\text{CS}} = 1.581$, in good agreement with the experimental values: 3.878, 3.924, and 3.865 Å, and $[c/a]_{\text{S2W}} = 1.584$. The Brillouin zone was sampled by $3 \times 3 \times 1$ (CS and S2W systems) and $2 \times 2 \times 1$ (SSWR and S71W systems) Γ-centered k-points meshes constructed through the Monkhorst–Pack scheme.\textsuperscript{36} A Gaussian smearing function with a width of 0.03 eV was used to broaden the energy levels around the Fermi level. Finally, we performed...
transition state (TS) searches through the improved dimer method (IDM). A data set collection of computational results, including all the structures, is available in the ioChemBD repository.

■ RESULTS AND DISCUSSION

Pt(111). In Figure 2 we show the energy profile for methanol decomposition on Pt(111) for the different systems in vacuum and under solvation. The first step is the adsorption of methanol to the metal surface, described by the replacement energy, $E_r$, the energy cost of replacing a water molecule by methanol.

$$E_r = E_m + E_w - E_m - E_w$$

Here, $E_m$ is the energy of the system with adsorbed methanol, $E_w$ stands for the same system with a water molecule located at the methanol position, while $E_m$ and $E_w$ are the energies of the isolated methanol and water molecules, respectively. The superscript "0" denotes vacuum, while $\epsilon$ stands for the continuum-solvated. Adsorption energies for methanol on the CS in vacuum are provided in Section S1. Except for the S5WR, where the replacement is more exothermic in solution than in vacuum, the replacement energies increase when solvating the systems. Nevertheless, the values of $E_r$ fall in the range ($-0.14, 0.11$) eV, making the replacement of water by methanol almost thermoneutral. We have also calculated $E_r$ for the S71W in vacuum. In this case, $E_r = 0.09$ eV, a value that falls in the range of the error for all our measured replacement energies.

Solvent effects are found in the first dehydrogenation step. From now on, we call $\Delta E_a$ the difference in the height of the first activation barrier between the paths involving $\text{O}^-\text{H}$ and $\text{C}^-\text{H}$ bond breaking. That is, $\Delta E_a = E_{TS, \text{OH}} - E_{TS, \text{CH}}$, with $E_{TS, \text{OH}}$ and $E_{TS, \text{CH}}$ standing for the first transition state energies, respectively.

In the three studied systems, there is an increase in $\Delta E_a$ under solvation with respect to what we observe in vacuum. This result agrees with both experimental and simulation data for methanol decomposition on Pt(111), where hydroxymethyl ($\text{H}_2\text{COH}$) becomes the most stable product in liquid.

A commonly adopted explanation is that, under solvation, the methanol OH group has a larger number of water molecules around it as compared to the CH$_3$ group. This situation reduces the probability for the binding of the OH group to the Pt surface catalytic sites and favors the C$^-\text{H}$ bond breaking. To account for this fact, it is specially important to include explicit water molecules near the OH group in order to reproduce part of its first solvation shell. This strategy, adopted in the S2W, limits the methanol OH group mobility and its further binding to the Pt surface. Increasing the number of explicit molecules further does not improve the model, as $\Delta E_a$ is very small (0.06 eV under solvation and 0.03 eV in vacuum), and thus, both C$^-\text{H}$ and O$^-\text{H}$ bond breakings are possible in contrast with the experimental observation. This result is due to the strong rigidity imposed by the S5WR model.

![Figure 2](image-url) Reaction profiles for the dehydrogenation of methanol to CH$_2$O on Pt(111). $\epsilon$ stands for solvated systems. For clarity, all the intermediates are labeled.

![Figure 3](image-url) Reaction profiles for the dehydrogenation of methanol to CH$_2$O on Pd(111). The dashed line for the CS corresponds to the path starting with the O$^-\text{H}$ bond breaking but with the oxygen atom from the adsorbed methoxy tricoordinated to the Pd surface. $\epsilon$ stands for solvated systems. On the right panels we show the methoxy configurations on the CS (one of the H atoms is hidden due to the perspective); tricoordinated (top) and monocoordinated (bottom). For clarity, all the intermediates are labeled.
We have simulated a methanol trimer adsorbed on the CS to address cooperative effects. For the isolated molecules, the methanol cluster is more stable than the separated methanol molecules (both in vacuum and solvated by water). When the trimer is adsorbed on Pt, the methanol trimer is only more stable in vacuum but solvation renders separated methanol molecules more favorable (see Section S2). Therefore, O−H bond breaking cannot be favored through cooperative interactions on the surface if water is present.

The second dehydrogenation step under solvation is rather similar for the S2W and the S5WR systems. In particular, for the CH\textsubscript{3}O dehydrogenation, the height of the barrier equals 0.17 and 0.39 eV for the CS and S2W, respectively, being this quantity for the S5WR equal to 0.33 eV. With regard to the CH\textsubscript{2}OH dehydrogenation, both the CS and S2W yield similar results as compared to the S5WR.

**Pd(111).** The analysis of the energy profiles on Pd(111) yields similar results to those for the Pt surface (see Figure 3). Regarding the replacement energies, \(E_r\) equals −0.19 and 0.01 eV for the CS and S2W, respectively, meaning that methanol adsorption is not much more preferred than desorption, even in an aqueous environment. The value of \(\Delta E_a\) for the CS is the same for both solvated and unsolvated systems (\(\Delta E_a = 0.33\) eV). However, \(\Delta E_a\) increases by 0.16 eV for the S2W when solvating the system (from 0.24 eV in vacuum to 0.40 eV under solvation). The first activation barrier for the C−H bond breaking decreases from 0.61 eV (vacuum) to 0.53 eV (solvation). These results indicate that, like in Pt, the C−H bond breaking becomes the preferential mechanism in aqueous environment.

With respect to the configuration of the adsorbed CH\textsubscript{3}O there is a difference between the Pd CS and S2W (see Figure 3). For the CS, the configuration with the oxygen atom coordinated to a fcc site (tricoordinated structure) is 0.34 eV more stable in vacuum than that with the oxygen atom coordinated to a top site (monocoordinated structure, see dashed line in Figure 3). When solvating the tricoordinated geometry, the energy decreases by 0.19 eV with respect to the flat situation due to the three available hydrogen atoms that the implicit solvent has to interact with. The scenario is quite different for the S2W as the water molecules maintain the adsorbate configuration flat and close to the surface. This adsorption reordering is the most serious reason to include explicit water molecules in any calculation under solvation.

**Ru(0001).** Water dissociates on Ru surfaces (dissociation ratio of about 40 to 50%), which might have an effect on methanol decomposition. For instance, explicit water molecules have been taken into account to study the conversion of levulinic acid on this surface.\textsuperscript{42,43} In Figure 4 we show the energy profile for methanol dehydrogenation on the Ru surfaces. For the CS \(E_f^0 = −0.22\) eV (in vacuum, \(E_f^0 = −0.09\) eV). Solvating the system increases \(\Delta E_a\) in 0.20 eV with respect to the vacuum situation, and therefore, the C−H bond breaking becomes the most favored methanol first decomposition mechanism. The exothermic character of the methanol O−H bond breaking depends, however, on the orientation of the C−O bond with respect to the surface plane. This issue, which

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**Figure 4.** Reaction profiles for the dehydrogenation of methanol to CH\textsubscript{2}O on Ru(0001). The dashed line for the CS corresponds to the path starting with the O−H bond breaking but with the oxygen atom from the adsorbed methoxy tricoordinated to the Ru atoms. \(\epsilon\) stands for solvated systems. On the right panels we show the methoxy configurations on the CS (one of the H atoms is hidden due to the perspective); tricoordinated (top) and monocoordinated (bottom). For clarity, all the intermediates are labeled.

**Figure 5.** Adsorbate configuration for the S2W before methanol replaces a dissociated water molecule (a) and after methanol is adsorbed (b).
is inherent to the CS, becomes more clear for the solvated system where the reaction is exothermic if the C–O bond is perpendicular to the surface plane and endothermic when the C–O bond is parallel to it. The net effect of taking the tricoordinated CH₂O structure is an increase in the height of the activation barrier for its dehydrogenation of 0.53 and 0.38 eV for the unsolvated and solvated systems, respectively.

The situation changes drastically when we add two water molecules (one dissociated) to the simulation box. On the reference S2W (that with two water molecules and an OH, see Figure 5a) the oxygen atom from the dissociated water is hydrogen-bonded to the same two hydrogen atoms. When methanol replaces the dissociated water, one of the remaining two water molecules becomes an OH group (see Figure 5b). Then, the global hydrogen bond network becomes slightly more distorted than for the previous situation making the methanol adsorption more endothermic than on the Pd and Pt surfaces ($E^f_r = 0.49$ eV and $E^f_r = 0.37$ eV). With regard to the methanol first dehydrogenation step, the O–H bond breaking is barrierless as the proton is transferred to the nearest water molecule (which is a dissociated water) and then the newly formed water molecule dissociates. This situation is not observed for the path involving the C–H bond breaking, where the activation barrier equals 0.52 and 0.46 eV for the unsolvated and solvated systems, respectively. These results give us some clues about the mechanism that governs methanol decomposition on Ru. This fact reinforces not only the need for the addition of two water molecules to the simulation box but also the importance of adopting a configuration such as that in Figure 5b. With regard to the CH₃OH dehydrogenation, it is also an inactivated process due to the proton transfer from the OH group to a water molecule. In this case, solvating the system involves that the surface with the adsorbed CH₃OH has a similar energy to that with the adsorbed CH₂O (0.13 eV of difference). For the CH₂O dehydrogenation, however, adding the solvent does not involve a large change in the barrier ($ΔE^f = 0.04$ eV).

To compile the results, a comparison between the three surfaces can be drawn. We have found that (i) for Pd and Pt the methanol C–H bond breaks first, whereas the O–H bond breaking is the first step on Ru, (ii) the number of explicit water molecules needed is the same for all surfaces, (iii) although the SSWR ring structures are slightly more flexible than the ice-derived ones, they might still be too rigid to represent the aqueous environments found in ref. 29 where 5-, 6- and 7-membered rings also appear, and (iv) the models used to study water-assisted processes on Ru need to account for the dissociation of some water molecules.

**CONCLUSIONS**

We have employed first-principles calculations to study the methanol dehydrogenation path to formaldehyde on Pd(111), Pt(111), and Ru(0001) under aqueous environment. In particular, we have adopted a combined explicit/implicit solvation approach. The results for the methanol dehydrogenation profiles on solvated Pd and Pt surfaces agree with experiments where the methanol C–H bond breaking is the preferred first dehydrogenation step. This situation is attributed to the fact that the methanol CH groups in solution are more exposed to the metal surface atoms than the OH group, which favors the C–H bond breaking. The calculated reaction profiles for the surface containing two explicit water molecules (S2W) qualitatively agree with those calculated for a benchmark system, the SSWR (surface with a five-water ring) where the methanol environment is close to that encountered on systems with several explicit water layers. The addition of two explicit water molecules also prevents the intermediates to adopt undesired configurations (especially for the case of the CH₂O in Pd and Ru in the CS) that result in overestimated values for the activation energies. Water dissociation on Ru(0001) makes the methanol decomposition very different from the case of Pd and Pt surfaces. The dehydrogenation of methanol on the Ru S2W occurs via a proton transfer from the methanol OH group to a nearby water molecule that dissociates afterward. This makes the methanol O–H bond breaking to become barrierless. No theoretical results exist for the dehydrogenation of methanol on solvated Ru(0001), and therefore, the aforementioned mechanism can serve as a reference for further studies. Notice that the water dissociation at the Ru interface is crucial to address the catalytic properties of this particular surface that is extensively used, for instance, in the conversion of levulinic acid to gamma-valerolactone.³⁵ Our results show the need to include both implicit and explicit water molecules in the computational box and thus pave the way toward faster and more reliable simulations in catalytic and electrochemical reactions on surfaces.

**ASSOCIATED CONTENT**

In Section S1, we provide the data associated with the energy profiles for the methanol dehydrogenation, that is methanol adsorption and replacement energies, and desorption energies of formaldehyde as well as activation and reaction energies; in Section S2, we give the energies for the clustered methanol species on metal surfaces (PDF).

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**Notes**

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