CO oxidation at Pd(100): A first-principles constrained thermodynamics study

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The possible formation of oxides or thin oxide films (surface oxides) on late transition-metal surfaces has recently been recognized as an essential ingredient when aiming to understand catalytic oxidation reactions under technologically relevant gas phase conditions. Using CO oxidation at Pd(100) as an example, we investigate the composition and structure of this model catalyst surface over a wide range of \((T,p)\) conditions within a multiscale modeling approach where density-functional theory is linked to thermodynamics. The results show that under the catalytically most relevant gas phase conditions a thin surface oxide is the most stable “phase” and that the system is actually very close to a transition between this oxidic state and a reduced state in terms of a CO-covered Pd(100) surface.

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I. INTRODUCTION

Catalytic oxidation using transition metals (TM)s as the active material is an important technological process, of which we have still only limited microscopic insight. Much valuable microscopic information that has been obtained under ultrahigh-vacuum (UHV) conditions is not directly transferable to the conditions of practical catalysis (with pressures of several bars and elevated temperatures). And in-situ techniques that would operate under these conditions are still struggling to achieve atomic-scale surface sensitive information. A key factor that hinders the direct transfer of the insight gained under UHV conditions is that under reaction conditions the entire structure and composition of the catalyst surface might be changed (often called the “materials gap”). And if a new material is created during the induction period of the catalytic process, it will obviously exhibit a very different chemical activity; it will be quantitatively and even qualitatively different. Under the oxygen-rich gas phase conditions and elevated temperatures of oxidation catalysis the surface of the transition metal catalyst might, e.g., be oxidized, so that the actual catalytically active state is not the pristine metal, but rather the formed oxide. One example for this is the CO oxidation reaction over ruthenium. Here it was found that under UHV conditions the Ru(0001) model catalyst shows almost no catalytic activity, whereas under high oxygen pressures its catalytic activity exceeds even the one of the frequently used palladium and platinum catalysts. This increase in the catalytic activity was traced back to the formation of a RuO2 film at the surface. For the ruthenium catalyst a clear distinction could thus be made between the different states: namely, the weakly catalytically active metal under low oxygen pressures and the highly active oxide film under high oxygen pressures.

For the model system discussed here, the CO oxidation at Pd(100), recent in-situ experimental measurements also indicate that an oxidic structure at the Pd(100) surface might be formed under gas phase conditions applied in industrial oxidation catalysis. Using surface x-ray diffraction (SXRD) measurements a \((T,p)\) diagram showing the detected structures in a pure oxygen atmosphere over a temperature range of \(T=600–1000\,\text{K}\) and pressures of \(p_{O_2}=10^{-9}–1\,\text{atm}\) could be inferred. A thin surface oxide structure was measured over an extended \((T,p)\) range, suggesting that this structure might also appear under catalytic reaction conditions. In reactor scanning tunneling microscopy (STM) experiments by Hendriksen et al. the structure of the Pd(100) surface could be directly monitored during the catalytic oxidation of CO. Here, the surface was exposed to both oxygen and CO at a total pressure of \(p_{\text{tot}}=1\,\text{atm}\) and a temperature of \(T=400\,\text{K}\). The partial pressures of the reactant gases CO and \(O_2\), as well as the reaction product \(CO_2\), were measured simultaneously with the STM images. Depending on the partial pressure ratio of \(O_2\) and CO the authors observed a change in the reaction rate, which was accompanied by a significant change in the morphology of the surface. This morphology change (albeit not atomically resolved) was interpreted as a change from the adsorbate-covered Pd(100) surface to an oxidic state.

To address this problem from a theoretical point of view we employ a multiscale modeling approach where we use density-functional theory (DFT) to describe the system on an electronic (microscopic) level. By linking these results to thermodynamics it becomes possible to address much larger system sizes and to compare the stability of different surface structures in contact with the surrounding gas phase. We use in particular a constrained equilibrium approach, where the surface is considered to be in full thermodynamic equilibrium with two separate gas phase reservoirs of \(O_2\) and \(CO\), and not allowing that \(O_2\) and \(CO\) can react. The key result obtained is a surface “phase diagram,” which provides first insight into possible surface structures over a wide range of temperature and pressure conditions of the \(O_2\) and \(CO\) gas phase. Focusing on gas phase conditions relevant for oxidation catalysis we find the system on the verge of either stabilizing a thin surface oxide structure or a CO-covered Pd(100) surface. It is thus well possible and even likely that the surface oxide structure contributes to the active state of the Pd(100) model catalyst under reaction conditions. However, to verify this, the kinetic effects of the ongoing catalytic \(CO_2\) formation need to be considered. This is done in a second step of our hierarchical approach by performing kinetic Monte Carlo (kMC) simulations on the surface structures identified here, on which we report in a consecutive paper.
II. THEORY

A. Gibbs free energy of adsorption

To determine the stability of a surface in contact with a gas phase reservoir we use the surface free energy $\gamma$. For a multicomponent system in equilibrium with atomic reservoirs (e.g., a surrounding gas or liquid phase environment or a macroscopic bulk phase) a general expression for the surface free energy is given by

$$\gamma(T,\{p_i\}) = \frac{1}{V} \left[ G_{\text{surf}} - \sum_i N_i \mu_i(T,p_i) \right].$$

Here, $G_{\text{surf}}$ is the Gibbs free energy of the solid including the surface, $A$ is the surface area, and $\mu_i(T,p_i)$ are the chemical potentials of the various species $i$ in the system. To compare the stability of different adsorption structures of oxygen and CO on the Pd(100) surface depending on the surrounding gas phase conditions one has to evaluate the Gibbs free energy of adsorption, $\Delta G_{\text{ads}}$.

For this, the stability of the different adsorption structures is compared with respect to the clean metal surface. For a Pd(100) surface in a constrained equilibrium with an O$_2$ and CO gas phase $\Delta G_{\text{ads}}$ is thus given by

$$\Delta G_{\text{ads}}(\Delta \mu_O, \Delta \mu_{CO}) = \gamma_{\text{Pd}(100)} - \gamma_{\text{O,CO@Pd}(100)}$$

$$= -\frac{1}{A} \left[ G_{\text{O,CO@Pd}(100)} - G_{\text{Pd}(100)} \right]$$

$$- \Delta N_{\text{Pd}} \mu_{\text{Pd}}^{\text{bulk}} - N_O(1/2E_{\text{O}_2}^\text{tot} + \Delta \mu_O) - N_{\text{CO}}(E_{\text{CO}}^\text{tot} + \Delta \mu_{\text{CO}})$$

$$= -\frac{1}{A} E_{\text{O,CO@Pd}(100)}^{\text{bind}} + \frac{N_O}{A} \Delta \mu_O + \frac{N_{\text{CO}}}{A} \Delta \mu_{\text{CO}},$$

where $G_{\text{O,CO@Pd}(100)}$ is the Gibbs free energy of the Pd(100) surface, with $N_O$ adsorbed O atoms and $N_{\text{CO}}$ adsorbed CO molecules, and $G_{\text{Pd}(100)}$ is the Gibbs free energy of the clean palladium surface. If the number of palladium atoms per surface area $A$ in the adsorption structure, $N_{\text{Pd}}$, and in the clean surface, $N_{\text{Pd}}^\text{bulk}$, are not equal (i.e., due to a surface reconstruction)—i.e., $\Delta N_{\text{Pd}} = N_{\text{Pd}}^\text{bulk} - N_{\text{Pd}} \neq 0$—then the excess (deficiency) atoms are taken from (put into) a bulk reservoir, represented by the chemical potential of the bulk solid phase, $\mu_{\text{Pd}}^{\text{bulk}}$. The chemical potentials of the two gas phase reservoirs of oxygen, $\mu_O$, and CO, $\mu_{\text{CO}}$, have been separated into a total energy contribution and the remaining part containing all the temperature- and pressure-dependent terms—i.e., $\mu_O = 1/2 \mu_{\text{O}_2} + \mu_{\text{O}}(T,p)$, setting $\Delta \mu_O(T,p) = 1/2 \mu_{\text{O}_2}(T,p) + \mu_{\text{O}}(T,p)$ and $\mu_{\text{CO}} = E_{\text{CO}}^\text{ads} + \mu_{\text{CO}}(T,p)$.

In the last line of Eq. (2) the difference of the Gibbs free energies of the clean and adsorbate covered surfaces, and of the bulk system, has been approximated by the DFT total energies $E_{\text{CO@Pd}}$.

A detailed discussion of the thereby neglected contributions to the Gibbs free energy of adsorption and their influence on the presented results will be given below.

In the constrained equilibrium approach leading to Eq. (2) the two gas phases are assumed to be noninteracting. The Gibbs free energy of adsorption $\Delta G_{\text{ads}}$ depends then linearly on the chemical potentials of both the oxygen and CO gas phases. The slope is respectively determined by the surface coverage of oxygen and/or CO. This is most apparent in the last line of Eq. (2), where we have introduced the average binding energy of O and CO as

$$E_{\text{O,CO@Pd}(100)}^{\text{bind}} = \left( \frac{E_{\text{O,CO@Pd}(100)}^{\text{tot}} - E_{\text{Pd}(100)}^{\text{tot}}}{N_O} \right) - \frac{E_{\text{CO}}^{\text{tot}}}{2}.$$  

Since the most stable structure will be the one with the lowest surface free energy, an adsorbate structure will be stable with respect to the clean surface if $\gamma_{\text{O,CO@Pd}(100)} < \gamma_{\text{Pd}(100)}$—i.e., if $\Delta G_{\text{ads}} > 0$. If two structures contain an equivalent amount of oxygen and CO, they will also show the same dependence on $\Delta \mu_O$ and $\Delta \mu_{\text{CO}}$. The ratio of how these two structures contribute to the stable structures is governed by the law of mass action, which enables us to directly exclude a less favorable structure if its binding energy $E_{\text{O,CO@Pd}(100)}^{\text{bind}}$ differs by much more than $k_B T$ from the one of the most favorable structure.

For the adsorption of O/CO on the Pd(100) surface the average binding energy defined in Eq. (3) is equivalent to the commonly used binding energy, which is often given per O atom or CO molecule, respectively—e.g., in the case of oxygen as

$$E_{\text{O@Pd}}^{\text{bind}(100)} = \frac{1}{N_O} \left( E_{\text{O,CO@Pd}(100)}^{\text{tot}} - E_{\text{Pd}(100)}^{\text{tot}} - \frac{N_O}{2} E_{\text{CO}}^{\text{tot}} \right).$$

In the case of additional adsorption of O (CO) on the reconstructed $\sqrt{3}$ surface oxide (which will be introduced below) the binding energy of the atom (molecule) with respect to this $\sqrt{3}$ structure is then given by (here exemplified for CO)

$$E_{\text{CO@Pd}}^{\text{bind}(\sqrt{3})} = \frac{1}{N_{\text{CO}}} \left( E_{\text{CO@Pd}}^{\text{tot}(\sqrt{3})} - E_{\text{Pd}}^{\text{tot}(\sqrt{3})} - N_{\text{CO}} E_{\text{CO}}^{\text{tot}} \right),$$

where $E_{\text{CO@Pd}}^{\text{tot}(\sqrt{3})}$ is the total energy of the $\sqrt{3}$ surface oxide. We note that $E_{\text{bind}}$ is in this case different from the above defined $E_{\text{bind}}$ in Eq. (3), since the latter also contains the energetic changes due to the formation of the $\sqrt{3}$ surface oxide, i.e., it is referenced with respect to the clean Pd(100) surface.

B. Bulk oxide stability

When considering a metal surface in contact with an O$_2$ and CO gas phase complete conversion into an extended bulk oxide is also a thermodynamic possibility. The stability of the corresponding bulk oxide therefore has to be evaluated with respect to the two gas phase components and compared to the various stable structures forming at the metal surface. A sufficient oxygen content in the gas phase will lead to the formation of bulk palladium oxide, PdO, whereas a sufficient CO content will favor the decomposition of PdO into CO$_2$ and Pd metal. In a pure oxygen gas phase the thermodynamic stability of the bulk oxide is given by

$$\mu_{\text{PdO}}^{\text{bulk}} < \mu_{\text{Pd}}^{\text{bulk}} + \mu_O,$$

whereas in a pure CO environment the stability condition for the oxide is
CO OXIDATION AT Pd(100): A FIRST-PRINCIPLES...

TABLE I. Computed heat of formation $\Delta H_{\text{f}}^{\text{PdO}}(0,0)$ and binding energies of O$_2$, CO and CO$_2$ (see text). The quoted experimental values are extrapolated to $T=0$ K, and the zero-point vibration energy (ZPE) has been removed. All values are in eV.

<table>
<thead>
<tr>
<th></th>
<th>PBE</th>
<th>RPBE</th>
<th>LDA</th>
<th>Expt.</th>
<th>ZPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta H_{\text{f}}^{\text{PdO}}(0,0)$</td>
<td>−0.87</td>
<td>−0.62</td>
<td>−1.42</td>
<td>−0.97</td>
<td></td>
</tr>
<tr>
<td>$E_{\text{bind}}^{\text{O}_2}$</td>
<td>−6.22</td>
<td>−5.75</td>
<td>−7.56</td>
<td>−5.30</td>
<td>(0.10)</td>
</tr>
<tr>
<td>$E_{\text{bind}}^{\text{CO}}$</td>
<td>−11.65</td>
<td>−11.20</td>
<td>−12.93</td>
<td>−11.33</td>
<td>(0.13)</td>
</tr>
<tr>
<td>$E_{\text{bind}}^{\text{CO}_2}$</td>
<td>−17.99</td>
<td>−17.09</td>
<td>−20.43</td>
<td>−16.98</td>
<td>(0.27)</td>
</tr>
<tr>
<td>$\Delta E_{\text{mol}}^{\text{bulk}}$</td>
<td>−3.24</td>
<td>−3.02</td>
<td>−3.72</td>
<td>−3.00</td>
<td></td>
</tr>
</tbody>
</table>

Reference 20.

If we approximate the chemical potential of the free CO$_2$ molecule by its total energy only, Eqs. (6) and (7) can be combined, yielding the stability criterion for PdO in an oxygen- and CO-containing gas phase:

$$\Delta \mu_{\text{CO}} - \Delta \mu_{\text{O}} \leq -2 \Delta H_{\text{f}}^{\text{PdO}}(T=0 \text{ K}) + \Delta E_{\text{mol}}^{\text{bulk}}.$$ (8)

Here, $\Delta H_{\text{f}}^{\text{PdO}} = E_{\text{PdO}}^{\text{tot}} - E_{\text{Pd}}^{\text{tot}} - 1/2E_{\text{CO}_2}^{\text{tot}}$ is the heat of formation of PdO at $T=0$ K and $\Delta E_{\text{mol}}^{\text{bulk}} = E_{\text{bind}}^{\text{CO}_2} - E_{\text{CO}_2}^{\text{tot}} - 1/2E_{\text{O}_2}^{\text{tot}}$ is the difference in binding energies between the three gas phase molecules. Table I compiles these binding energies, as well as the heat of formation of PdO, as obtained by previously published highly converged DFT calculations$^{19}$ and using the computational methodology described below. In addition the experimental values are quoted. These values have been extrapolated to $T=0$ K, and the zero-point vibration energy (also listed in Table I) has been removed.

Substituting the chemical potential of CO$_2$, $\mu_{\text{CO}_2}$, by only the total energy term $E_{\text{CO}_2}^{\text{tot}}$ in Eq. (7) is a rather crude approximation. The most important contributions that have been neglected arise from the vibrational and translational free energy. The assumption that the CO$_2$ formed at the surface is readily transported away motivates us to disregard the translational free energy contribution as we discussed in more detail in Ref. 18. The vibrational free energy contribution will be of the order of the ZPE for the temperature range discussed here. As can be seen in Table I the ZPE value for CO$_2$ is approximately 0.3 eV. Variations of $\Delta E_{\text{mol}}^{\text{bulk}}$ in Eq. (8) of this order of magnitude do not affect any of the conclusions discussed below, which justifies the rather crude approximation for the present purpose.

C. Computational setup

With the approximations made in the last two subsections, the crucial quantities determining the stability of surface structures are the total energies of the extended surfaces and of the involved gas phase molecules. These total energies are obtained by DFT calculations which have been performed within the full-potential (linearized) augmented plane wave + local orbital (L)APW+lo method$^{21,22}$ as implemented in the WIEN2k code.$^{23}$

All surfaces are simulated within the supercell approach using inversion-symmetric slabs consisting of five Pd(100) layers with adsorption of oxygen and/or CO or the reconstructed surface oxide plus additional O/CO on both sides. The vacuum between consecutive slabs is at least 14 Å. The adsorption layers and two outermost palladium layers have been fully relaxed. The muffin-tin radii have been set to $R_{\text{MT}}^{\text{Pd}}=2.0$ bohr for palladium, $R_{\text{MT}}^{\text{O}}=1.0$ bohr for oxygen, and $R_{\text{MT}}^{\text{C}}=1.0$ bohr for carbon. Inside the muffin-tins the wave functions are expanded up to $l_{\text{max}}=12$ and the potential up to $l_{\text{max}}=6$. For the Pd(100)-(1×1) structure a [10×10×1] Monkhorst-Pack (MP) grid has been used to integrate the Brillouin zone (BZ). For larger surface unit cells the MP grid has been reduced accordingly to assure an equivalent sampling of the BZ. Since calculations with different MP grids are not fully comparable, the energies of the clean metal surface, $E_{\text{Pd}}^{\text{tot}}$, and of bulk palladium, $E_{\text{Pd}}^{\text{tot}}$, need to be evaluated at different temperatures. The influence of these different xc functionals on the results will be discussed below.

III. O AND/OR CO ADSORPTION AT Pd(100)

In this section we discuss the different adlayer structures of oxygen and/or CO on Pd(100) that will be compared within the constrained thermodynamic equilibrium approach. In addition to the on-surface adsorption of O/CO on Pd(100), the formation of a reconstructed (\(\sqrt{5} \times \sqrt{5}\))R27° (abbreviated with \(\sqrt{5}\) in the following) surface oxide has experimentally been observed at higher oxygen exposure.$^{27-30}$ Since the structure of the \(\sqrt{5}\) surface oxide is quite different from the clean Pd(100) surface, also exhibiting different adsorption sites, we will discuss the two surfaces separately.

A. Adlayers on Pd(100)

1. Pure oxygen adsorption

Experimentally, four different adsorption structures have been observed when exposing the Pd(100) surface to
TABLE II. Calculated binding energies of on-surface adlayers of O/CO on Pd(100). For the pure adlayer structures the binding energies are given per O atom and CO molecule, respectively. For the mixed structures the total binding energies are given. All values are in eV.

<table>
<thead>
<tr>
<th>Coverage Θ</th>
<th>E_{bind}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure O structures</td>
<td></td>
</tr>
<tr>
<td>p(2 × 2)-O_{hol}</td>
<td>0.25</td>
</tr>
<tr>
<td>c(2 × 2)-O_{hol}</td>
<td>0.50</td>
</tr>
<tr>
<td>Pure CO structures</td>
<td></td>
</tr>
<tr>
<td>p(2 × 2)-CO_{br}</td>
<td>0.25</td>
</tr>
<tr>
<td>c(2 × 2)-CO_{br}</td>
<td>0.50</td>
</tr>
<tr>
<td>(2\sqrt{2} \times \sqrt{2})R45^°-CO_{br}</td>
<td>0.50</td>
</tr>
<tr>
<td>(3\sqrt{2} \times \sqrt{2})R45^°-CO_{br}</td>
<td>0.67</td>
</tr>
<tr>
<td>(4\sqrt{2} \times \sqrt{2})R45^°-CO_{br}</td>
<td>0.75</td>
</tr>
<tr>
<td>(1 × 1)-CO_{br}</td>
<td>1.00</td>
</tr>
<tr>
<td>Mixed O/CO structures</td>
<td></td>
</tr>
<tr>
<td>(2 × 2)-O_{hol}-CO_{br}</td>
<td>0.50</td>
</tr>
<tr>
<td>(2 × 2)-2O_{hol}-CO_{br}</td>
<td>0.75</td>
</tr>
<tr>
<td>(2 × 2)-O_{hol}-2CO_{br}</td>
<td>0.75</td>
</tr>
<tr>
<td>(2 × 2)-2O_{hol}-2CO_{br}</td>
<td>1.00</td>
</tr>
</tbody>
</table>

oxygen. These are a p(2 × 2), a c(2 × 2), a (5 × 5), and the \(5\) surface oxide structure. The \(5\) surface oxide will be discussed in detail in the next section. The (5 × 5) structure will not be considered in this study, since the structure and state of the oxygen atoms have not been well established so far, i.e., there exists no well-defined structural model. In addition, the (5 × 5) structure appears to be only of metastable character and its formation was found to be very sensitive to the surface preparation and oxygen exposure range, so that in the oxidation as well as in the reduction process the (5 × 5) structure can be bypassed, going directly from a (2 × 2) to a \(\sqrt{5}\) structure and vice versa.

This leaves as ordered adlayers only the p(2 × 2) at 0.25 monolayer (ML) coverage and the c(2 × 2) at 0.5 ML. Computing the binding energy of O atoms in all high-symmetry sites offered by the (100) surface (bridge, top, and hollow), we find in both structures the fourfold hollow site to be the most stable adsorption site. The binding energies are calculated with respect to the O₂ gas phase molecule [cf. Eq. (4)], and the values of the binding energies per oxygen atom are listed in Table II. It can be seen that the binding energy decreases going from the p(2 × 2) to the higher-coverage c(2 × 2) adlayer, indicating overall repulsive interactions between the adsorbed oxygen atoms.

2. Pure CO adsorption

The adsorption of CO on Pd(100) has been studied intensively. All in experimental studies it is found that CO binds upright in the bridge position via the C atom. At a coverage of Θ = 0.50 ML an ordered (2\sqrt{2} \times \sqrt{2})R45° adlayer is formed, which is compressed to a (3\sqrt{2} \times \sqrt{2})R45° structure at Θ = 0.67 ML and a (4\sqrt{2} \times \sqrt{2})R45° at Θ = 0.75 ML.

Mixed O/CO adsorption

As to the adsorption of oxygen and CO on the Pd(100) surface, experimentally no ordered overlayers have been reported so far. If the Pd(100) surface is exposed to both oxygen and CO, the two adsorbed species tend to form separate domains instead. In their low-energy electron diffraction (LEED) measurements Stuve et al. observed that for a fully developed p(2 × 2)-O/Pd(100) structure the p(2 × 2) LEED pattern vanished after exposure to CO at a temperature of \(T = 80 \text{ K}\). Assuming a barrier of \(= 1.0 \text{ eV}\) for the reaction of O and CO to form CO₂, it is rather unlikely that the adsorbed oxygen reacted with the CO at this low temperature. The disappearance of the LEED signal was thus interpreted as a CO-induced disordering of the oxygen islands. To nevertheless obtain an idea about the simultaneous adsorption of O and CO on Pd(100) we set up different models for mixed overlayer structures that seemed to be most obvious from a combinatorial point of view. For this we used (2 × 2) surface unit cells placing one to two O atoms and CO molecules in their favorite adsorption sites—i.e., O in hollow and CO in bridge sites. Excluding any structures where O and CO are closer to each other than the length \(a\) of the (1 × 1) surface unit cell we obtain the four different struc-
The average binding energies calculated using Eq. (3) are listed in Table II.

Similar to the pure adlayer structures of O or CO on Pd(100), the binding energy per adsorbate decreases with increasing coverage in the considered coadsorption structures. This trend in binding energies reflects the aforementioned repulsive interactions among the adsorbed O atoms and CO molecules. For a coverage of \( \Theta = 0.50 \) ML, the binding energy of the mixed O/CO overlayer is less favorable by \( 70 \) meV than the sum of the binding energies of the respective pure adlayers of O and CO. This indicates that the repulsive interactions among adsorbed O and CO are even stronger than between O, O and CO, CO, respectively, in the pure adlayer structures. At finite temperatures configurational entropy would tend to counteract the repulsive interactions, but in the temperature range discussed here this entropy contribution is rather small. The adsorption of both O and CO into separate domains. The adsorption of CO into a pure adlayer structure further decreases the binding strength of the adsorbates.

The even stronger repulsive interactions in these mixed adsorbate structures will thus favor the separation of oxygen and CO into separate domains. The adsorption of CO into a \( p(2 \times 2) \)-O structure can then induce a demixing of the two species and destroy the long-range order, which is consistent with the disappearance of the LEED spot intensities in the aforementioned experiment by Stuve et al.\cite{stuve1981}

### B. O and CO on the surface oxide

At a coverage of \( \Theta = 0.80 \) ML, the adsorbed oxygen induces a reconstruction of the Pd(100) surface forming the \( \sqrt{5} \) surface oxide structure. An atomistic model for the surface oxide (cf. Fig. 3) was proposed in a detailed experimental and theoretical study, revealing that the structure can actually be described as a PdO(101) trilayer on Pd(100).\cite{keast1981} The surface unit cell contains four palladium and four oxygen atoms. Two Pd atoms are fourfold and the other two are twofold coordinated by oxygen. There are also two kinds of O atoms. Two O atoms sit on top of the reconstructed Pd layer and two at the interface to the Pd(100) substrate forming the aforementioned trilayer structure.

In addition to the previous discussion on this surface oxide structure,\cite{keast1981} we note that the potential energy surface for a registry shift of the entire surface oxide trilayer over the Pd(100) substrate is very shallow. In particular, shifts of up to \( \sim 0.5 \) Å along the direction as indicated by the arrow in

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**Fig. 3.** (Color online) Top view of the \( (\sqrt{5} \times \sqrt{5})R27^\circ \) surface oxide structure (left part) and the Pd(100) substrate (right part) and their corresponding surface unit cells (black lines). Red small spheres represent the lattice oxygen atoms, large light-blue ones the reconstructed Pd atoms, and large dark-blue spheres Pd atoms belonging to the underlying Pd(100) substrate. The arrow indicates the easy shift direction of the surface oxide trilayer along the Pd(100) substrate.

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**Fig. 4.** (Color online) Top views of high-symmetry adsorption sites on the \( (\sqrt{5} \times \sqrt{5})R27^\circ \) surface oxide structure. Red small spheres represent the lattice oxygen atoms, large light-blue ones the reconstructed Pd atoms, and large dark-blue spheres Pd atoms belonging to the underlying Pd(100) substrate.
TABLE III. Calculated binding energies of CO on the \( \sqrt{5} \) surface oxide structure calculated using Eq. (5). 2CO denotes that two CO molecules are adsorbed in the \( \sqrt{5} \) surface unit cell. The adsorption of two CO molecules in top4f sites or one CO in bridge and a second in a top2f site does not lead to a stable structure. All values are per CO molecule and in eV.

<table>
<thead>
<tr>
<th></th>
<th>( E_{\text{bind}}^{\text{CO@5}} )</th>
<th>( E_{\text{bind}}^{\text{O@5}} )</th>
<th>( E_{\text{bind}}^{\text{CO@5}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO( ^{\text{br}} )</td>
<td>(-0.93)</td>
<td>(-0.75)</td>
<td>(-0.93)</td>
</tr>
<tr>
<td>CO( ^{\text{top2f}} )</td>
<td>(-0.62)</td>
<td>(-0.51)</td>
<td>(-0.51)</td>
</tr>
<tr>
<td>CO( ^{\text{top4f}} )</td>
<td>(-0.13)</td>
<td>(-0.29)</td>
<td>(-0.29)</td>
</tr>
</tbody>
</table>

The adsorption of oxygen in the high-symmetry sites of the \( \sqrt{5} \) surface oxide gives similar results. Again, the bridge site is the most stable site (\( E_{\text{bind}}^{\text{O@5}} = -0.14 \) eV) and the hollow sites are unstable upon relaxation. However, it is now not possible to adsorb any oxygen in the top2f and top4f sites in the \( \sqrt{5} \) unit cell. If one CO molecule and one O atom are adsorbed simultaneously, likewise the adsorption in two bridge sites is found to be favored.

Our calculations with one and two adsorbates per \( \sqrt{5} \) surface unit cell thus indicate noticeable overall repulsive lateral interactions. We correspondingly extended our calculations also to more sparse adlayers in \( (2 \times 1) \) and \( (1 \times 2) \) \( \sqrt{5} \) surface unit cells, but did not find significant lateral interactions extending across the \( \sqrt{5} \) surface unit cell. The obtained binding energies were always to within 25 meV per adsorbate of those calculated for the same adsorption site in a \( (1 \times 1) \) cell.

To increase the configuration space of considered overlayers based on the \( \sqrt{5} \) surface oxide, we also considered structures in which the original surface oxide structure is slightly modified. In a first step, one of the upper hollow site oxygen atoms is removed, and CO and O are again placed in the aforementioned sites. Also in this modified structure the bridge site results as the preferred adsorption site. If both upper oxygen atoms are removed from the \( \sqrt{5} \) surface oxide, we find the structure already to some extend destabilized. Due to the lower palladium density in the reconstructed layer (four Pd atoms on five Pd(100) substrate atoms per unit cell) the structure is rather open, and structural relaxation showed that the Pd atoms can now move quite easily in lateral direction on the surface. It is still possible to adsorb O/CO in any of the other adsorption sites, but we found these structures either unstable upon relaxation or in general much less stable than the corresponding structures, where the original O hollow vacancy is refilled.

In addition to simply removing the upper oxygen atoms from the \( \sqrt{5} \) surface oxide structure and filling additional on-surface adsorption sites, we also substituted the topmost O atoms by CO molecules and placed additional O atoms and/or CO molecules in top2f, top4f, and bridge sites. In the substituted hollow site the CO binds quite strongly. However, since the O atoms bind even stronger to this site by 0.6 eV, O will always preferentially occupy the hollow sites when the two adsorbates compete for this site. The situation is reversed for adsorption in the bridge sites. Regardless of whether O or CO occupies the threefold hollow sites, we always find a stronger binding by at least 0.7 eV of CO compared to O at these bridge sites.

Even if we leave out the four unstable hollow sites in Fig. 4, there is finally still a huge amount of possible overlayers structures that can be created by combinatorially placing an arbitrary number of O and CO per surface unit cell into any of the available sites. Since there is only little known about the adsorption on the surface oxide, none of these structures can be excluded a priori. Motivated by the strong repulsive interactions seen in our calculations with one or two adsorbates per surface unit cell, we nevertheless discard quite a number of these structures with the criterion that no two adsorbates may sit in directly neighboring sites (i.e., at a distance of less than 1/4 of the length of \( \sqrt{5} \) surface unit cell). This still leaves 92 “plausible” structures, and DFT calculations were performed for all of them (for more details on the calculated overlay structures see Ref. 44). Out of these only 55 are stable upon relaxation and are then considered in our constrained “phase diagram.”

IV. STABILITY IN A CONSTRAINED EQUILIBRIUM
WITH AN O\(_2\) AND CO GAS PHASE

Using the Gibbs free energy of adsorption as defined in Eq. (2) it is possible to compare the stability of all calculated adsorption structures in a constrained equilibrium with an O\(_2\) and CO gas phase. In this approach the surface is considered to be in equilibrium with two separate gas phase reservoirs of O\(_2\) and CO, i.e., the formation of CO\(_2\) is neglected in the gas phase as well as at the surface. In the gas phase this assumption seems quite justified, since the direct reaction of O\(_2\) and CO is kinetically hindered by a huge free energy barrier. At the surface, though, the reaction is actually supposed to take place at the working catalyst. Here, the assumption of a constrained equilibrium is thus only reasonable as long as the O\(_2\) and CO adsorption and desorption
CO OXIDATION AT Pd(100): A FIRST-PRINCIPLES...

FIG. 5. (Color online) Surface “phase diagram” of the Pd(100) surface in constrained thermodynamic equilibrium with an O$_2$ and CO gas phase. The atomic structures underlying the various stable (co)adsorption structures on the metal and the surface oxide, as well as a thick bulklike oxide film (indicated by the bulk unit cell), are also shown (Pd=large blue spheres, O=small red spheres, and C=small yellow spheres). The black bar marks gas phase conditions representative of technological CO oxidation catalysis—i.e., partial pressures of 1 atm and temperatures of 300–600 K. The dependence of the chemical potentials of the two gas phases is translated into pressure scales for $T=300$ K and $T=600$ K (upper $x$ axes and right $y$ axes).

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A. Surface phase diagram

The results of the above-described calculations concerning the aforesaid overlayers of oxygen and CO on Pd(100), also including the (modified) $\sqrt{5}$ surface oxide structure, are summarized in Fig. 5, which shows the most stable “phases” (i.e., the ones that maximize $\Delta G^{ad}$ as defined in Eq. (2)) for any given chemical potential of the oxygen and CO gas phases. The dependence on the chemical potentials has also been converted into more intuitive pressures scales for temperatures of $T=300$ K and $T=600$ K in the top two $x$ axes for oxygen and the right two $y$ axes for CO. As expected from the Gibbs phase rule, stable “phases” cover volumes in the three-dimensional space spanned by ($T$, $P_{O_2}$, $P_{CO}$) and coexistence regions between two “phases” cover areas. Correspondingly in the two-dimensional sectional plane shown in Fig. 5, this leads to areas and lines for the two cases, respectively. Out of the set of 191 tested structures only 11 turn out to be “phases” appearing in Fig. 5.

We will start the discussion of the obtained constrained surface “phase diagram” in Fig. 5 in the lower left corner. Here, both the chemical potential of oxygen and CO are very low, i.e., the oxygen and CO content in the gas phase is insignificant and consequently the clean Pd(100) surface results as the most stable system state. If we now move along the $x$ axis in Fig. 5 to the right, we reach more and more oxygen-rich conditions, while the CO content in the gas phase is kept low. The increase in the oxygen chemical potential leads to a stabilization of oxygen-containing structures with increasing coverage: First, the $p(2 \times 2)$-O overlayer on Pd(100), then the $\sqrt{5}$ surface oxide, and finally the PdO bulk oxide representing thick bulklike oxide films on the surface. This sequence of stable structures was also confirmed by in-situ surface x-ray diffraction measurements. Interestingly, the $c(2 \times 2)$ structure observed under UHV conditions does not appear in the surface “phase diagram.” Thus we conclude that the $c(2 \times 2)$ structure is most likely a metastable state produced by the exposure kinetics.
If the oxygen content is kept low in the gas phase and the CO content is gradually increased—i.e., moving from the lower left corner along the y axis to the top in Fig. 5—a series of ordered CO adlayer structures with increasing coverage is stabilized on Pd(100). At first we find the also experimentally observed (2\sqrt{2} \times 2)R45^\circ, (3\sqrt{2} \times \sqrt{2})R45^\circ, and (4\sqrt{2} \times \sqrt{2})R45^\circ structures and, finally, for a very high CO content in the gas phase a (1 \times 1) structure with 1 ML CO in bridge sites.

Starting again in the lower left corner of Fig. 5 we now move along the diagonal, which corresponds to increasing both the oxygen and CO content in the gas phase. Intuitively, one would expect coadsorption structures of O and CO on Pd(100) to become favorable. But none of the above-discussed ordered coadsorption structures (cf. Fig. 2) are found to be a most stable “phase” under any gas phase conditions. This is consistent with the already mentioned experimental findings, that O and CO prefer to form separate domains rather than ordered coadsorbed overlayers. The lower stability of such mixed structures can be explained by the strongly repulsive interactions between adsorbed O and CO, which lead to no significant decrease in binding energies (cf. Table II). However, we can of course not exclude that ordered arrangements with different periodicities than those considered here would not lead to a lowering in the repulsive interactions. To take a reasonable part in the “phase diagram,” though, the binding energies would have to increase by as much as 0.3–0.5 eV per O atom/CO molecule compared to the now proposed structures. Compared to the pure overlayer structures this would even imply the necessity of attractive interactions between the two species.

For high oxygen and CO content in the gas phase (upper right part of Fig. 5) we find instead coadsorbed structures involving the √5 surface oxide (two orange-white hatched regions) to become stable. These two mixed structures correspond to the surface oxide with one and two CO molecules adsorbed in bridge sites, respectively. In a small range of very oxygen-rich and intermediate CO gas phase conditions additional O adsorption in the bridge sites of the √5 structure leads also to a stable “phase” (dark red-white hatched region).

Looking again at the whole surface “phase diagram” in Fig. 5 the 11 “phases” can be divided into three groups: on-surface adlayer structures of O or CO on Pd(100), phases involving the √5 surface oxide structure, and the stability region of the bulk oxide as discussed in Sec. II B. Focusing on these three different groups of phases two important conclusions with respect to the relevance of oxide formation under catalytic reaction conditions can be drawn. First, the formation of a thick, bulklike oxide (gray cross hatched area) at the surface under technologically relevant gas phase conditions of O2 and CO (black bar in Fig. 5, p1 \sim 1 atm, T \sim 300–600 K) can be ruled out. This is thus in marked contrast to the much wider stability range of bulk RuO2, which does extend to these conditions. Second, the stability region of the √5 surface oxide (hatched area) does extend to such conditions. In fact, it is either this monolayer thin surface oxide or CO adlayers on Pd(100), which are neighboring “phases” around the catalytically active region in (T,p) space. As can be seen in Fig. 5 the catalytically relevant conditions are actually right at the boundary between the √5 surface oxide structures and the CO overlayer structures on Pd(100). Small changes in the Gibbs free energy of adsorption causing a shift in this boundary could thus well affect the conclusion as to which is the most stable “phase” under these conditions. Apart from the assumption of a constrained thermodynamic equilibrium, the main uncertainties, which may cause such changes in ΔGads, are the approximate DFT total energies and the neglected free energy contributions. We verified that the uncertainties in the DFT total energies due to the numerical approximations (supercell setup, finite basis set) are not significant in this respect. What needs to be scrutinized are therefore the uncertainties due to the approximate xc functional underlying the DFT total energies and the neglected free energy contributions to ΔGads. This will be done in the next two subsections.

B. Evaluating the Gibbs free energy

To calculate the Gibbs free energy of adsorption ΔGads the Gibbs free energies of the different components entering Eq. (2) have to be evaluated. The Gibbs free energy

\[ G(T,p) = E^{\text{tot}} + F^{\text{vib}} - TS^{\text{conf}} + pV \]  

(9)

comprises contributions from the total energy E^{\text{tot}}, the vibrational free energy F^{\text{vib}} including the ZPE, the configurational entropy S^{\text{conf}}, and the pV term. In Eq. (2) we substituted the Gibbs free energy difference by only the leading term, the total energy differences, which can be directly obtained from DFT calculations. We assess the uncertainty introduced by this approximation by an order-of-magnitude estimate of the remaining contributions to ΔGads. If this first approximation reveals that the results are significantly influenced by considering all contributions to the Gibbs free energy, the respective terms have to be calculated explicitly. However, the order-of-magnitude estimate can be obtained very easily and is thus helpful to decide whether or not it becomes necessary to evaluate the entire Gibbs free energy.

Following the discussion in Refs. 13 and 16 the contributions to the Gibbs free energy of adsorption arising from the pV term and the configurational entropy are rather small for systems like the one presented here. We will thus only discuss the most crucial approximation: namely, the neglect of the vibrational free energy contribution, ΔF^{\text{vib}} to the Gibbs free energy of adsorption. In order to estimate the size of ΔF^{\text{vib}} we approximate the phonon density of states (DOS) within the Einstein model by one characteristic frequency. Following the approach outlined in Ref. 13 the vibrational contribution to the Gibbs free energy of adsorption for the (2 \times 2)-O/Pd(100) structure can then be estimated as

\[ \Delta F^{\text{ads,vib}}(T) \approx -\frac{1}{A} \left( F^{\text{vib}}(T, \omega_{\text{O-Pd}}) - \frac{1}{2} F^{\text{vib,ZPE}}(\omega_{\text{O-Pd}}) \right), \]

(10)

where \( F^{\text{vib}}(T, \omega) \) is the frequency-dependent function:
CO OXIDATION AT Pd(100): A FIRST-PRINCIPLES...

FIG. 6. (Color online) Vibrational contribution to the Gibbs free energy of adsorption for the \(p(2 \times 2)-O\) adlayer on Pd(100) within the Einstein approximation using different characteristic frequencies (see “inset” and text).

\[
F^{\text{ads, vib}}(T, \omega) = \frac{1}{2} \hbar \omega + k_B T \ln(1 - e^{-\beta \omega}),
\]

with \(\beta = 1/k_B T\). \(\bar{\omega}^\text{surf}\) is the characteristic vibrational frequency of an oxygen atom adsorbed in a fourfold hollow site on Pd(100), and \(\bar{\omega}^\text{ads, O}_2\) is the stretch frequency of the \(O_2\) gas phase molecule. The change in the vibrational contribution of the Pd atoms in the clean and oxygen-covered Pd(100) phase molecule. The change in the vibrational contribution to \(\Delta G^\text{ads, vib}\) is neglected, so that the vibrational contribution to \(\Delta G^\text{ads}\) is given by the difference in the vibrational energy of the \(O_2\) molecule in the gas phase and the oxygen atom on the surface. For the oxygen molecule only the ZPE (not included in the presented DFT total energies) has to be considered, since all remaining contributions to \(F^{\text{ads, vib}}\) are already contained in \(\Delta \mu^\text{O}\) (cf. Eq. (2)). In Fig. 6 the resulting vibrational contribution \(\Delta F^{\text{ads, vib}}(T)\) is shown for a characteristic frequency of \(\bar{\omega}^\text{surf}\) of 48 meV for the Pd-O stretch frequency of an adsorbed O atom and an O-O vibrational frequency of \(\bar{\omega}^\text{ads, O}_2\) of 196 meV. We find that for the \(p(2 \times 2)-O/Pd(100)\) structure \(A(2 \times 2) = 31.16 \, \text{Å}^2\) the vibrational contribution to \(\Delta G^\text{ads}\) stays below \(\sim 3 \, \text{meV/Å}^2\) for temperatures up to \(T = 1000 \, \text{K}\) (black solid line in Fig. 6). Since the chosen \(\bar{\omega}^\text{surf}\) is only a rough guess, we also allow for a \(\pm 50\%\) change of this value, but even then the contribution does not increase considerably (red dashed, \(-50\%\), and blue dotted, \(+50\%\), lines in Fig. 6).

Similar estimates have been performed for every structure considered for the surface “phase diagram” in Fig. 5. For CO-containing structures \(\Delta F^{\text{ads, vib}}\) comprises two different contributions when comparing the gas phase and the adsorbed state, namely, the change of the O-C stretch vibration due to the adsorption and an additional Pd-C vibration. For structures involving the \(\sqrt{3}\) surface oxide \((A, \tau = 38.95 \, \text{Å}^2)\) an additional contribution arises from the change in the vibrational energy between bulk and surface Pd atoms. The \(\sqrt{3}\) structure contains one surface atom fewer than the corresponding Pd(100) surface per unit cell which also has to be balanced by the bulk reservoir (\(\Delta N^\text{Pd} = -1\) in Eq. (2)).

FIG. 7. (Color online) Surface “phase diagram” for the Pd(100) surface in a constrained equilibrium with an \(O_2\) and CO gas phase. The top plot (a) shows the RPBE results and the bottom one (b) the LDA results. The color coding of the different “phases” is equivalent to the one used in Fig. 5. The black bar marks again gas phase conditions representative of technological CO oxidation catalysis—i.e., partial pressures of 1 atm and temperatures between 300 and 600 K. Note that the range of chemical potentials shown is larger for the LDA results, compared to the RPBE and the range in Fig. 5.

With this procedure we find that for all structures \(\Delta F^{\text{ads, vib}}\) stays always below \(10 \, \text{meV/Å}^2\) for temperatures up to \(T = 600 \, \text{K}\). The surface “phase diagram” discussed in the previous section is not significantly changed if these estimated, maximum values for the vibrational contribution are included. There are some small shifts in the boundaries between stable “phases,” but none of the unstable structures disappears or is missing. Furthermore, the boundary between surface oxide structures and CO adlayers on Pd(100) is only marginally affected and consequently remains in the vicinity of the catalytically active region in \((T, p)\) space. We are thus confident that for the results discussed here the approximation of the Gibbs free energy differences by only the total energy terms in Eq. (2) is justified.

C. Influence of the xc functional

To obtain an estimate of the uncertainty in Fig. 5 due to the choice of the PBE-GGA xc functional we reevaluate the
constrained surface “phase diagram” using the RPBE (Ref. 25) and LDA (Ref. 26) xc functionals. The corresponding surface “phase diagrams” are shown in Fig. 7 and are to be compared with Fig. 5. The “phase diagram” obtained by using the RPBE approximation for the xc energy (top graph, Fig. 7(a)) looks in fact very similar to the previously discussed one. There are some shifts in the actual “phase” boundaries, and the stability regions of the high-coverage phases of oxygen and CO, \( \sqrt{5} + \text{O}^\bullet \) and \((1 \times 1)\text{CO}^\bullet/\text{Pd}(100)\) structures, are shifted outside the range of chemical potentials shown. Yet the overall topology is fully conserved. In agreement with the PBE results the bulk oxide is not a stable “phase” anywhere near ambient gas phase conditions. Due to the smaller heat of formation (cf. Table. I), the stability region of the bulk oxide is in fact much farther away from this region. Most importantly, the boundary between the simple overlayer structures on \( \text{Pd}(100) \) and the phases based on the \( \sqrt{5} \) surface oxide structure (hatched area) is only very little influenced, so that the most interesting region for oxidation catalysis (black bar) is again right at the transition between a CO-covered \( \text{Pd}(100) \) surface and “phases” involving the surface oxide.

Even if the LDA is used as xc functional [bottom plot, Fig. 7(b)], the stability region of the bulk oxide does not extend to the catalytically relevant gas phase conditions. It covers a much larger range compared to the “phase diagrams” obtained using the two GGA functionals though. It should also be noted that in the shown LDA “phase diagram” the range of O and CO chemical potentials is enlarged to lower values to include the stability region of the clean metal surface, which in the LDA appears at much lower gas phase concentrations. This is a consequence of the strong overbinding in the LDA, which stabilizes any adsorbate structure at much lower pressures at the surface. Nevertheless, the range of technologically relevant catalytic gas phase conditions (black bar) lies again right at the boundary between the CO covered \( \text{Pd}(100) \) surface and \( \sqrt{5} \) surface oxide structures.

From these results, we conclude that the choice of xc functional does have a strong influence on the absolute values of the binding energies and thereby also on the location of most of the phase boundaries in Figs. 5 and 7. However, particularly the boundary between the CO overlayer structures on \( \text{Pd}(100) \) and the \( \sqrt{5} \) surface oxide is almost unchanged. The conclusion on the proximity of both phases to the catalytically relevant gas phase conditions seems therefore untouched by the uncertainty due to the approximate xc functional.

V. CONCLUSIONS

The stability of the \( \text{Pd}(100) \) surface has been investigated in a constrained thermodynamic equilibrium with a two-component gas phase consisting of \( \text{O}_2 \) and \( \text{CO} \). In this approach the formation of \( \text{CO}_2 \) in the gas phase and at the surface is not considered, such that the effect of the surrounding gas phase on the surface structure and composition is modeled (as a first approximation) through contact with independent reservoirs representing the reactants. To establish the actual surface “phase diagram” a large number of different structures with O and CO adsorbed in high-symmetry sites on the \( \text{Pd}(100) \) surface and on the \( \sqrt{5} \) surface oxide structure have been considered. We find that under gas phase conditions of ambient temperatures and pressures, as applied in heterogeneous oxidation catalysis, it is either the nanometer thin surface oxide structure or a CO-covered \( \text{Pd}(100) \) surface that is stable, whereas the stability region of the bulk oxide does not extend to these gas phase conditions.\(^{18}\)

To obtain an estimate of the uncertainty introduced by the choice of a specific xc functional the total energies entering the surface “phase diagram” have been calculated using the PBE, RPBE, and LDA. Comparing the “phase diagrams” for these three different xc functionals, partly dramatic shifts in the positions of the boundaries between the different stable “phases” can be observed. Yet it can also be seen that the position of the boundary between the CO-covered \( \text{Pd}(100) \) “phases” and the \( \sqrt{5} \) surface oxide structures is in fact little affected. For all three xc functionals, the catalytically active region is very close to this boundary, but just still within the stability range of the surface oxide. We also verified that this finding is not affected by the numerical uncertainties and the free energy contributions neglected in our approach.

These results suggest that the only monolayer thin surface oxide structure might indeed be a relevant structure for the CO oxidation reaction on \( \text{Pd}(100) \) at technological relevant pressures. It has to be considered, though, that there are still two notable approximations in the constrained atomistic thermodynamics approach, as it has been applied here. First, configurational entropy is not included, and at finite temperatures a first effect would be to smear out the phase boundaries in the “phase diagram” and create coexistence regions, where the energetically low-lying configurations are mixed according to the law of mass action. At the phase boundary relevant for CO oxidation catalysis, this could, e.g., lead to the formation of coexisting domains of CO-covered \( \text{Pd}(100) \) and surface oxide patches. Second, the neglected kinetics of the ongoing catalytic \( \text{CO}_2 \) formation might significantly change the stability range of the different phases as obtained in the constrained equilibrium approach. A possible coexistence between patches of metallic \( \text{Pd}(100) \) covered by the reactants and patches of the surface oxide (possibly also with adsorbed \text{CO}) could then even under steady-state conditions go hand in hand with a continuous formation and decomposition of the oxidic phase. Such oscillations in the morphology of the catalyst surface could again significantly influence the catalytic function of this surface.

Based on the insight described in this paper we thus identify either the \( \sqrt{5} \) surface oxide or CO-covered \( \text{Pd}(100) \) as surface structures most relevant under catalytically interesting gas phase conditions. It is therefore specifically the stability of the \( \sqrt{5} \) surface oxide against CO-induced decomposition, which needs to be scrutinized to assess the active state of the surface under reaction conditions. In a second paper following the present one, we correspondingly extend our
study to investigate this stability. Using kinetic Monte Carlo simulations we then explicitly scrutinize the two major uncertainties of the present approach: namely, the effect of configurational disorder and the kinetic effects due to the ongoing reactions.

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