The reconstruction of R h(001) upon oxygen adsorption

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Abstract

We report on a first-principles study of the structure of O/Rh(001) at half a monolayer of oxygen coverage, performed using density functional theory. We find that oxygen atoms sit at the center of the black squares in a chess-board c(2×2) pattern. This structure is unstable against a rhomboid distortion of the black squares, which shortens the distance between an O atom and two of the four neighboring Rh atoms, while lengthening the distance with respect to the other two. We actually find that the surface energy is further lowered by allowing the O atom to get off the short diagonal of the rhombus thus formed. We predict that the latter distortion is associated with an order–disorder transition, occurring below room temperature. The above rhomboid distortion of the square lattice may be seen as a rotation of the empty white squares. Our findings are at variance with recent claims based on STM images, according to which it is instead the black squares which would rotate. We argue that these images are indeed compatible with our predicted reconstruction pattern. © 1998 Elsevier Science B.V. All rights reserved.

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1. Introduction

The study of the physical and chemical properties of the (001) surface of rhodium is an important step towards the ambitious goal of understanding the catalytic properties of transition metals. Recently, the clean R h(001) surface has attracted experimental [1–3] and theoretical [4–9] interest because of the anomalously small interlayer relaxation at the surface, an unexpected result which is in contrast with the general behavior of transition-metal surfaces. One more reason of interest is due to the behavior of the surface upon oxygen adsorption. Oxygen adsorption on Rh(001) is known to be dissociative and to saturate at half a monolayer, independently of the adsorption temperature. At this coverage, a (2×2) reconstruction has been observed by SPA-LEED and confirmed by STM [10]. The oxygen atoms sit in the troughs formed by four first-layer rhodium atoms and fill these sites in a c(2×2) geometry. This geometry may be considered as a chess-board whose black squares are occupied by oxygen atoms, whereas the white ones are empty. Within this picture, the reconstruction observed in Ref. [10] has been described as a rotation of the black squares, resulting in a (2×2)p4g symmetry (see Fig. 1a). This distortion preserves the shape of the black squares, while the white squares become rhomboid. A similar behavior is observed for nitrogen and carbon.
distortion can then be rationalized by a simple chemical model of the re-bonding of O atoms at the surface.

2. Computational method

Our calculations are based on density functional theory within the local-density approximation [15,16], using Ceperley-Alder exchange-correlation (XC) energies [17]. The one-particle Kohn-Sham equations are solved self-consistently using plane-wave basis sets in a pseudopotential scheme. Because of the well-known hardness of the norm-conserving pseudopotentials for the O and – to a lesser extent – Rh atoms, we make use of ultra-soft pseudopotentials [18] which allow an accurate description of the O and Rh valence pseudo-wave functions with a modest basis set including plane waves up to a kinetic energy cut-off of 30 Ry. In the case of Rh, we found it convenient to treat the s and p channels using the norm-conserving potential, while the ultra-soft scheme is applied only to the hard d orbital [19–21]. Brillouin zone integrations have been performed using the Gaussian smearing [22] special-point [23] technique. We find that the structural properties of bulk rhodium are well converged of the irreducible wedge of the Brillouin zone (IBZ). The isolated surface is modeled by a periodically repeated supercell. We have used the same supercell for both the clean and the O-covered surfaces. For the clean surface, we have used five atomic layers plus a vacuum region corresponding to ~6 layers. We have used the same Gaussian smearing function as in the bulk calculations with a (12,12,2) Monkhorst-Pack mesh [23], resulting in 21 special k-points in the (1 × 1) surface IBZ. Convergence tests performed with a value of s twice as small and a correspondingly finer mesh of special points resulted in no significant changes in total energies and equilibrium geometries. The latter are found
by allowing all the atoms in the slab to relax until the force acting on each of them is smaller than $0.1 \times 10^{-3} \text{Ry/}a_0$.

### 3. Results

For the clean surface we find that – in agreement with other ab-initio calculations [4–9] – the first layer relaxes inward by $3.8 \pm 0.2\%$ (see Table 1), whereas the second layer is practically unrelaxed (the error is essentially a finite-size effect, and it is estimated by repeating the calculation using thicker slabs). As noted in Section 1, the value of the first-layer relaxation reported in the experimental literature is anomalously small (the first interlayer spacing is practically equal to its bulk value within error bars). Recently, Cho and Sch鋟ler [9] pointed out that a proper account of the vibrational contribution to the surface free energy may result in a reduction of the inward relaxation of the first layer, thus bringing theoretical predictions in better agreement with experiments. No attempts of estimating these vibrational effects have been done in the present work. Our calculated values for the surface energy $s$ and work function $w$ are $s \approx 1.4 \text{ eV/atom}$ and $w \approx 5.5 \text{ eV}$ (see Table 1).

Special care must be taken when estimating the latter, for the XC potential goes to zero rather slowly in the vacuum region. As the electrostatic (Hartree) potential converges much more rapidly than the XC potential to its vacuum value, it is convenient to evaluate the vacuum level by simply neglecting the XC contributions. A preliminary study of oxygen adsorption, we performed a couple of simple ab-initio molecular dynamics simulations of an O$_2$ molecule impinging onto the surface. We tried two possible initial conditions for the molecule, in both of which the axis of the molecule is parallel to the surface. In the first case, the projection of the center of the molecule on the surface falls on top of a surface atom, whereas in the other, it falls on a bridge site. The initial velocity of the molecule is orthogonal to the surface, and its modulus corresponds to a temperature of 300 K, whereas the surface is initially assumed to be at zero temperature. The initial distance of the molecule from the surface is $\approx 6 \text{ a.u.}$, which is essentially in the vacuum. When the molecule arrives at the surface, the molecular bond breaks and the surface heats up. In order to find the ground-state atomic configuration, we slowly cooled the system, ending with the structure displayed in Fig. 2: the oxygen atoms adsorb in every second hollow site of the surface, resulting in a $c(2 \times 2)$ structure, where the squares whose centers

### Table 1

<table>
<thead>
<tr>
<th>Unit</th>
<th>$d_0/%$</th>
<th>$d_{11}$ (Å)</th>
<th>$d_{o1}/d_{11}$ (Å)</th>
<th>$s$ (eV)</th>
<th>$E-E_{\text{hollow}}$ (eV/atom)</th>
<th>$\delta s$ (eV/atom)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean</td>
<td>$-3.8%$</td>
<td>$5.5$</td>
<td>$1.36$</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Experiment</td>
<td>$-1.2 \pm 1.6%$</td>
<td>$5.0$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Top</td>
<td>$1.81$</td>
<td>$+3$</td>
<td>$7.5$</td>
<td>$+1.5$</td>
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<td></td>
</tr>
<tr>
<td>Bridge</td>
<td>$1.33$</td>
<td>$+0.5$</td>
<td>$6.8$</td>
<td>$+0.3$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hollow</td>
<td>$1.02$</td>
<td>$+0.5$</td>
<td>$6.2$</td>
<td>$+0.0$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S. clock</td>
<td>$-4$</td>
<td>$1.02$</td>
<td>$-6.2$</td>
<td>$-0.003$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>A. clock</td>
<td>$-11$</td>
<td>$0.98-1.06$</td>
<td>$-6.1$</td>
<td>$-0.030$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The coverage is $\theta = 1 \times 2$. $d_0$ is the bulk lattice spacing; $d_{11}$ is the distance between the oxygen atoms and the first rhodium layer; and $d_{o1}$ is the distance between the first and second layers. For the asymmetric clock reconstruction, the two numbers given refer to the two inequivalent first-layer rhodium atoms (see Fig. 2). $s$ is the amplitude of the movement of the first-layer rhodium atoms upon distortion (see Fig. 3b); $w$ is the work function; and $s$ is the surface energy.

a from Ref. [1].
b from Ref. [24].
c from Ref. [25].

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a from Ref. [1].
b from Ref. [24].
c from Ref. [25].
other adsorption sites (see Fig. 3), which are, however, much larger than the energy differences between the unreconstructed hollow structure and the corresponding clock reconstructions.

The work function is larger at a coverage $\theta = 0.5$ than for the clean surface (see Table 1), thus indicating that electrons tend to transfer from the Rh substrate to the oxygen layer, so as to increase the surface dipole. In Fig. 4, we display the adsorption energy $E_{\text{ad}}$ as a function of coverage defined as $E_{\text{ad}}(\theta) = s(\theta) - s(0) - (\theta / 2)E_{O_2}$, where $s(\theta)$ is the surface energy of the oxygen-covered system, and $E_{O_2}$ is the energy of the isolated $O_2$ molecule. Coverage $\theta = 1$ is realized by filling all the hollow sites; for $\theta = 0.75$ and $\theta = 0.25$, three and one of the four possible hollow sites in a $(2 \times 2)$ substrate supercell have been occupied. The minimum energy corresponds to a coverage in the range $0.5 \leq \theta \leq 1$. In this range, however, the curve is very flat, and the variation of the adsorption energy ($\sim 100$ meV) is smaller than the expected accuracy of the terms which enter its definition.

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The value of the most stable coverage is determined by a trade-off between the adsorption energy of an isolated oxygen molecule, which tends to favor a high coverage, and the oxygen-oxygen electrostatic repulsion due to charge transfer, which becomes more effective when the average O–O distance becomes smaller than some typical screening length, and which tends instead to favor a low coverage.

4. Discussion

From Fig. 2, it is evident that the oxygen sublattice forms a zigzag arrangement which is not observed in the experiments [10]. However, there are two equivalent three-fold sites for each cell occupied by an oxygen adatom, one on each side of the bridge. Neglecting the interactions between different adsorption sites, each of them is therefore two-fold degenerate. We postulate that the system undergoes an order-disorder transition at some critical temperature which depends on the magnitude of the adsorbate-adsorbate interactions, and that this temperature is lower than that at which...
Fig. 3. Sketch of the various O adsorption sites of the Rh(001) surfaces considered in this work. Small dark circles: O atoms. Large lighter circles: first-layer Rh atoms. Large darker circles: second-layer Rh atoms.

Fig. 4. Oxygen adsorption energy $E_{ad}(\text{eV})$, as function of coverage $\theta$.

the STM images were taken. The fact that no disorder in the O adlayer appears from these images indicates that the frequency of oxygen barrier crossing is much larger than the scanning frequency of the STM. This is compatible with our estimate of the barrier height (~30 meV, as obtained by comparing the energies of the symmetric clock and the asymmetric clock structures) which is of the order of the room temperature.

The STM pictures reported in Ref. [10] are such that only Rh or O atoms are visible in turn in a same picture. Because of this, it is difficult to judge whether the squares which undergo a rhomboid distortion are those filled by an adatom, or they are instead empty. What appears to be rather evident is that the distorted squares display a depletion with respect to the undistorted ones. In analogy with a similar reconstruction which is observed to occur in N/Ni(001) and which is much better characterized [26], the authors of Ref. [10] conclude that O atoms are thrust down the first Rh layer in the middle of the undistorted squares. We have simulated STM pictures using the Tersoff–Hamann model [27], but we have not been able to eliminate the images of the O atoms from any of our simulated pictures. In all of them, O atoms appear as protrusions in the middle of the distorted squares, in qualitative agreement with the experimental observation that undistorted squares are depleted with respect to the distorted ones.

In Fig. 5, we display the surface-projected densities of states (SDOS) of the clean and of the oxygen-covered Rh(001) surface, along with their decompositions into various atomic-like contributions. The SDOS of the clean surface differs from that of the bulk FCC structure mainly because of its more pronounced narrowness due to the lower coordination of surface atoms [8,20], and it is almost entirely determined by its d-like component (see Fig. 5a). In the bulk FCC structure, the three $d_{xy}$, $d_{xz}$, and $d_{yz}$ orbitals are degenerate, and so are $d_{3z^2}$ and $d_{3z^2-r^2}$. This degeneracy is partially lifted at the surface because the surface atoms lose the neighbors in the direction orthogonal to the surface. For the (001) surface, the $d_{xy}$ and $d_{yz}$ orbitals are still equivalent by symmetry and, hence, degenerate. The position of the oxygen atomic p-level is around 3.1 eV below the Fermi energy of the oxygen-covered surface. Inspection of Fig. 5d shows that the O–O level gives rise to a bonding and an antibonding main peak, respectively, below (~5.8 V) and above (~1 eV) the Fermi level. Upon oxygen adsorption, first-layer Rh atoms become locally inequivalent according to whether the neighboring oxygen atoms are aligned along the x or y directions (see Fig. 5e). The projected densities of states (PDOS) plotted in Fig. 5d refer to those Rh atoms which have O neighbors aligned along the y direction (the PDOS of the other Rh atoms can be obtained by simply exchanging x with y). It is also easy to recognize...
that the same bonding and anti-bonding features occur in the $d_{xz}$ band whose atomic orbitals have lobes oriented towards the adsorbed oxygen, whereas the $d_{yz}$ band remains similar to that of the clean surface (see Fig. 5c) because in that direction, the surface sites are empty. The $O_p$ orbitals make bonds with the $d_{xz}$ orbitals of the Rh atoms along the $x$ direction, whereas the $O_p$ orbitals hybridize with the $d_{yz}$ orbitals of the Rh atoms along the $y$ direction. We find that the $c(2 \times 2)$ structure is not the most stable one, and that a $(2 \times 2)p4g$ reconstruction occurs which shortens and strengthens the O–Rh bonds along one direction, while lengthening them in the perpendicular direction, as illustrated in Fig. 5e. The amplitude of the distortion depends rather sensi-
tively on the lattice parameter: using our calculated lattice parameter, we estimate the distortion to be ~4%. If one uses a lattice parameter which is 1% larger, the amplitude of the distortion is also increased, reaching a value of ~6%. The opposite occurs if the lattice parameter is reduced. The tendency of the system to strengthen two of the four rhodium bonds at the expenses of the other two results in a net lowering of the surface energy of ~3 meV/atom, thus stabilizing the (2 × 2)p4g structure. In the asymmetric, more stable clock structure, the size of the distortion is ~3% (using our calculated lattice parameter), and the surface energy gain is ~30 meV/atom. This shows that the reconstruction occurs because the optimal O–Rh bond length is shorter than that realized in the ideal geometry. The chemical contribution to the energy reduction which determines the distortion is illustrated in Fig. 5f, which shows the differences between the Rh and O PDOS after and before the reconstruction. In both cases, we notice a push of states towards lower energies.

5. Conclusions

Our ab-initio results confirm the experimental evidence of the dissociative character of the oxygen adsorption on Rh(001) and that the favored adsorption site is the four-fold one. A (2 × 2)p4g reconstruction of the surface is also predicted, in agreement with SPA-LEED data. At variance with claims based on recent STM work, we find that this reconstruction is due to a rhomboid distortion of the squares formed by first-layer Rh atoms which have an O atom adsorbed in the middle, and that this structure is unstable with respect to a departure of the adatoms from the centers of the resulting rhombi. We argue that the experimental evidence upon which these claims are found is rather weak, and we suggest therefore that further experimental work is needed to fully characterize the reconstruction of this surface. Furthermore, we find evidence that an order–disorder transition should occur at some temperature below that at which the STM pictures have been taken. Our findings are substantiated by a simple chemical model of the mechanisms responsible for the reconstruction.

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