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First principles analysis of hydrogen chemisorption on Pd–Re alloyed overlayers and alloyed surfaces

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Gradient corrected periodic density functional theory (DFT–GGA) slab calculations were used to examine the chemisorption of atomic hydrogen on various Pd–Re alloyed overlayers and uniformly alloyed surfaces. Adsorption was examined at 33% surface coverage, where atomic hydrogen preferred the three-fold fcc sites. The binding energy of atomic hydrogen is observed to vary by as much as 0.7 eV due to Pd–Re interactions. The computed adsorption energies were found to be between $-2.35$ eV [for monolayer Pd-on-Re, i.e., $Pd_{33}/Re(0001)$] and $-3.05$ eV [for $Pd_{33}/Re_{66}/Pd(111)$]. A $d$-band weighting scheme was developed to extend the Hammer–Nørskov surface reactivity model [Surf. Sci. 343, 211 (1995)] to the analysis of bimetallic Pd–Re alloyed systems. The hydrogen chemisorption energies are correlated linearly to the surface $d$-band center, which is weighted appropriately by the $d$-band coupling matrix elements for Pd and Re. The farther the weighted $d$-band center is shifted below the Fermi energy, the weaker is the interaction of atomic hydrogen with the alloyed Pd–Re surface. © 2000 American Institute of Physics.

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I. INTRODUCTION

Much of the petrochemical industry practices hydrogenation of unsaturated hydrocarbon feeds over supported bimetallic particles.\(^1\) The synergy between the two metals often leads to catalytic reactivity that is significantly different as compared to each of the monometallic components. An improved understanding of how bimetallic metal–metal interaction influences surface reactivity will likely provide insights into designing surface alloys with specific catalytic properties. Over the past decade, first-principles density functional theory (DFT) calculations\(^2\)–\(^5\) and UHV surface-science studies\(^6\)–\(^13\) have helped to elucidate the chemical reactivity of well-defined pseudomorphic overlayers and uniformly alloyed surfaces. Simplified models have been proposed to correlate changes in chemical reactivity to electronic perturbations in these idealized surfaces. Goodman and co-workers, for instance, showed how changes in the chemisorption energy of CO on pseudomorphic overlayers, correlate with measured XPS shifts in the core-level shifts measured at the $d$-band center.\(^14\)–\(^15\) Using frontier orbital theory\(^16\),\(^17\) and the Newns–Anderson chemisorption model,\(^18\) Hammer and Nørskov developed a simplified chemisorption and reactivity model,\(^3\)–\(^5\),\(^20\)–\(^23\) which demonstrated that the metal–adsorbate bond strength is closely related to the surface metal $d$-band structure. The core-level shifts measured by Goodman and co-workers were shown to be commensurate with changes in the valence $d$-band center.\(^3\) Recent results have verified the effectiveness of this simple model in the analysis of chemisorption of adsorbates such as CO,\(^3\)\(^,\)\(^22\) H\(_2\),\(^21\)\(^,\)\(^23\)–\(^25\) NO,\(^26\)\(^,\)\(^27\) ethylene\(^28\) and maleic anhydride\(^5\) on monometallic and pseudomorphic overlay surfaces. The simple two-level interaction model,\(^3\)\(^,\)\(^5\)\(^,\)\(^20\)–\(^23\)\(^,\)\(^24\) which treats orbital overlap and Pauli repulsion, was also found to provide excellent correlation between activation barriers, for bond-breaking and coupling reactions on pseudomorphic overlayer surfaces, and shifts in the position of the surface $d$-band center.\(^2\)\(^,\)\(^28\)

The original model proposed by Hammer and Nørskov\(^19\) has been predominantly used to examine the interaction of adsorbates, such as CO and atomic hydrogen, with a single type of metal atom on monometallic or pseudomorphic overlayer surfaces. The changes in the reactivity were found to correlate well with the $d$-band center of the metal to which the adsorbate was bound. For a surface alloy, the situation may conceivably be more complicated, where adsorption in a multifold site may result in the adsorbate forming bonds with more than one type of metal atom. For example, species such as atomic carbon or hydrogen are most strongly bound to three-fold or four-fold hollow sites on a metal surface. On an alloyed surface, it is likely that the atomic adsorbate may form bonds concurrently with two or more types of metal atoms. In this paper, we demonstrate that the $d$-band center model of Ref. 21 may also be extended, in some cases, to describe such adsorption scenarios. To illustrate this point we consider the chemisorption of atomic hydrogen in the threefold fcc sites on various Pd–Re alloyed surfaces.

II. COMPUTATIONAL METHODS

DFT–GGA periodic slab calculations were performed to examine the chemisorption of atomic hydrogen for a 33% surface coverage of the fcc hollow sites. The $\gamma_3 \times \gamma_3 R30^\circ$ unit cell was used to model the periodic adsorp-
tion of hydrogen for 1/3 ML coverage of the surface. The fcc hollow site was determined to be the most favorable adsorption site for hydrogen chemisorption on the monometallic Pd(111) and Re(0001) surfaces and was therefore chosen for this study. The Kohn–Sham equations were solved self-consistently using the Perdew–Wang 91 (PW91) exchange–correlation functional. The inner-shell electrons of all the atoms were described by scalar-relativistically corrected, norm-conserving pseudopotentials. The maximum kinetic energy of the plane wave basis was 40 Rydberg.

III. RESULTS AND DISCUSSIONS

The bimetallic Pd–Re surfaces that were examined for hydrogen chemisorption are shown in Fig. 1. In a previous study, we examined the effect of the number of metal layers on the hydrogen chemisorption energy and found negligible changes in the adsorption energy by increasing the number of metal layers beyond three. Three metal layer slabs are therefore used throughout this study. Slabs (a), (b), (e), and (f) in Fig. 1 represent uniformly alloyed Pd–Re monolayers, pseudomorphically placed on a monometallic Pd(111) or Re(0001) substrate. The slab surfaces (c) and (d) are uniformly alloyed Pd–Re surfaces containing a 33:66 and 66:33 ratio of Re/Pd, respectively. Since the bulk crystal structure for Re is hcp, the alloyed surface containing more than 50% Re was modeled using an hcp pattern. Analogously, an fcc pattern was used for the Pd-rich alloyed surface. At the initial structure, the surfaces were chosen to have the lattice parameter of the metal with the larger atomic concentration in the bulk. Since the optimized interatomic distances in monometallic Pd(111) (2.75 Å) and Re(0001) (2.76 Å) are very similar, the choice of the lattice parameter at the starting geometry should not introduce significant strain in the lattice. During the geometry optimization scheme, the adsorbate and the first two metal layers of the slab were completely relaxed. Throughout the following discussion, overlayer or surface layer refers to the top metal layer to which hydrogen is bound. Substrate refers to the second and third metal layers.

Table I shows the DFT–GGA optimized metal–hydrogen distance for hydrogen chemisorption on monometallic Pd(111), Re(0001), and the various alloyed Pd–Re surfaces. It is observed that when hydrogen is bonded to both Pd and Re, the Pd–H bond distances are significantly longer than that on Pd(111) (1.8 Å). The Re–H distances in these cases are slightly shorter than they are on Re(0001) (1.91 Å), suggesting that atomic hydrogen forms stronger bonds with the surface Re atoms as compared to the surface Pd atoms. This is especially evident for the surfaces containing 66% Re, where the Pd–H distance (2.0 Å) is about 0.2 Å longer than that on Pd(111), and hydrogen is almost bridge bound over the Re atoms. The surface structure in this case can to some extent be considered as Re dimers embedded in a Pd(111) surface, with atomic hydrogen almost bridge bound to the dimer.

Table I summarizes the DFT–GGA computed binding energies (BE) for atomic hydrogen on the monometallic Pd(111), Re(0001), and bimetallic Pd–Re surfaces examined in this study. Note that the BE for (\(\sqrt{3}\times\sqrt{3}\)) adsorption of atomic hydrogen reported here are about 0.1–0.15 eV stronger than the BE reported earlier for 100% surface coverage. The strengthening of the adsorption energy with decrease in surface coverage is primarily due to the reduction in through-space repulsive interactions and is discussed in detail in Ref. 25. The resulting DFT-computed adsorption energies for hydrogen on monometallic Pd(111) (−2.77 eV) and Re(0001) (−2.98 eV) are in reasonably good agreement with previously reported theoretical calculations and experimental measurements. Calculations suggest that the interaction of atomic hydrogen with the PdML/Re(0001) and PdRe3/Re(0001) surfaces are relatively weak with binding energies of −2.35 eV and −2.48 eV, respectively. These

![FIG. 1. Pd–Re alloyed surfaces examined in this paper for the chemisorption of atomic hydrogen. (a) 66:33 Pd-Re alloyed monolayer on Pd(111); (b) 33:66 Pd-Re alloyed monolayer on Pd(111); (c) 66:33 Pd-Re alloy (fcc); (d) 33:66 Pd-Re alloy (hcp); (e) 66:33 Pd-Re alloyed monolayer on Re(0001); and (f) 33:66 Pd-Re alloyed monolayer on Re(0001).](image)
binding energies are significantly weaker than those for hydrogen on the monometallic Pd(111) (−2.77 eV) and Re(0001) (−2.98 eV) surfaces. This indicates that the Re(0001) substrate significantly weakens the interaction of atomic hydrogen with the surface metal atoms. Although there is no reported experimental estimate for the binding energy of hydrogen on the Pd$_{\text{ML}}$/Re(0001) surface, it is known experimentally that adsorbates such as CO bind weaker to this pseudomorphic overlayer as compared to monometallic Pd

Table I indicates that the hydrogen-surface bond strength for most of the bimetallic Pd–Re surfaces are between −2.73 eV and −2.94 eV, which is intermediate to metal–hydrogen surface bond strengths for monometallic Pd(111) and Re(0001) surfaces. The binding of atomic hydrogen on the Pd$_{33}$Re$_{66}$/Pd(111) (−3.05 eV) surface, however, is slightly stronger than on the Pd–Re alloyed and monometallic Pd(111), Re(0001) surfaces. An important observation from this study is that the variation in the hydrogen binding energy over the various Pd–Re alloyed surfaces examined here is about 0.7 eV. This implies that the energy for dissociative adsorption of hydrogen can vary by as much as 1.4 eV, depending on the bimetallic Pd–Re ensemble to which hydrogen is bound. The second important conclusion from Table I is that bimetallic Pd–Re surfaces can exhibit metal–hydrogen bond strengths that are significantly weaker, or sometimes even stronger, than each of the monometallic surfaces.

It is apparent that bimetallic surfaces can have markedly different reactivity than monometallic surfaces, and an appropriate treatment of bimetallic catalysts will require the analysis of bimetallic alloyed systems. Since an exhaustive treatment of adsorbate interaction with all possible bimetallic ensembles is practically impossible, there is considerable value in determining the important electronic parameters of the surface that control surface reactivity and evaluating how bimetallic interactions affect them.

Table II summarizes the electronic properties of the bare surfaces of monometallic Pd(111), Re(0001) and the alloyed Pd–Re surfaces. The $d$-band filling for the surface metal atoms, reported in Table II, were determined by calculating the fractional area under the curve of the density-of-states (DOS) projected to the $d$ band, which is below the Fermi energy. The $d$-band center for the surface metal atoms were calculated by taking the first moment of the normalized projected DOS about the Fermi level. From Table II, it appears that bimetallic Pd–Re interaction in the alloyed surfaces has nominal effect on the $d$-band filling for Pd and Re. The Pd $d$-band filling is 0.95–0.96 and the Re $d$-band filling is between 0.68 and 0.73, in all cases. It therefore appears that the significant changes in the hydrogen adsorption energy cannot be attributed to charge-transfer effects alone. In Fig. 2, we have plotted the projected DOS for the alloyed Pd–Re overlayers and uniformly alloyed surfaces. The solid lines depict the DOS projected to the surface Pd $d$ band. The dotted lines
correspond to the projection onto the d band for the surface Re atoms. It is evident that there are significant differences in the shape of the d band (see Fig. 2) due to bimetallic alloying. Integration under the curve up to the Fermi level indicates that the d-band filling remains approximately the same. The changes in the shape of the d band primarily affect the location of the d-band center, i.e., the first moment of the normalized projected DOS. This effect is more predominant in the Pd d-band structure as compared to that for Re. In examining Fig. 2, the plots from top to bottom [(a)–to–(c)–to–(e) and (b)–to–(d)–to–(f)], show the effect on the surface metal d band due to increasing the amount of Re in the substrate. It appears that increasing the amount of Re in the substrate shifts the weight of the Pd d band away from the Fermi energy. Looking from left-to-right in Fig. 2 [(a) and (b); (e) and (f)], we see the effect of increasing the amount of Re in the surface layer. The d band for the surface Pd atoms below the Fermi level is observed to shift further away from the Fermi energy as the number of Re neighbors is increased.

From Table II, the d band center for Pd is observed to vary anywhere between ~ −0.44 eV and ~ −1.18 eV from the d-band center for Pd(111) (~ −1.98 eV). The changes in the Re d band due to the presence of Pd neighbors, on the surface and in the bulk, are relatively subtle. There are smaller shifts in the d-band position due to bimetallic Pd–Re interaction. The Re d-band center for the bimetallic surfaces, which are tabulated in Table II, vary between ~ −0.18 eV and ~ −0.53 eV relative to Re(0001) (~ −1.16 eV). Oelhafen and co-workers have experimentally demonstrated similar shifts in the valence d-band of metals for amorphous alloys such as Pd–Zr, Cu–Zr, and Au–Sn, using ultraviolet photoelectron spectroscopy. They find that increasing the Zr content in a Pd–Zr alloy shifts the average Pd d-band position to higher binding energies by as much as 2 eV. The shift in the Zr d band is relatively small due to forming the bimetallic amorphous alloy. Our results seem to be consistent with the experimental observation.

In Ref. 21, a simple model was proposed that correlated changes in the metal–adsorbate bond energy to changes in the metal d-band center for the bare-metal surface. In a previous paper, we demonstrated that the hydrogen binding energies on monometallic Pd(111), Re(0001), and pseudomorph Pd$_{ML}$/Re(0001) and Re$_{ML}$/Pd(111) overlayers correlate with the d-band center for the surface metal atom. In this paper, we extend the model to correlate the hydrogen chemisorption energy on alloyed Pd–Re overlayers and surfaces by using a weighted d-band center as the key parameter to represent the surface electronic structure.

For adsorption of hydrogen on an alloyed Pd–Re surface, the d-band positions for the Pd and Re atoms on the surface are markedly different. The extent of overlap between hydrogen and the d bands of Pd and Re are also different. This is seen as a significantly elongated Pd–H and contracted Re–H bond distances on the alloyed Pd–Re surfaces. It is therefore necessary to use an appropriate weighting scheme to account for the differences in the interaction of Pd and Re with hydrogen. Since the coupling of the hydrogen 1s state with the metal d band is proportional to the d-band coupling matrix element, it seems to be an appropri-
hydrogen chemisorption on alloyed Pd–Re surfaces. It is demonstrated that the $d$-band center, weighted by the $d$-band coupling matrix element, provides an appropriate parameter by which to correlate the surface reactivity of bimetallic coupling matrix element, provides an appropriate parameter to which the number of Re neighbors around Pd shifts the $d$-band center for Pd away from the Fermi energy. The same is true for the surface Re atoms, but the shift in the $d$-band center due to increasing the number of Pd neighbors is relatively small.

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