Structure and stability of borohydride on Au(111) and Au3M(111) (M = Cr, Mn, Fe, Co, Ni) surfaces

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Structure and stability of borohydride on Au(111) and Au₃M(111) (M = Cr, Mn, Fe, Co, Ni) surfaces

Ryan Lacdao Arevalo, Mary Clare Sison Escaño, Andrew Yu-Sheng Wang and Hideaki Kasai

We study the adsorption of borohydride on Au and Au-based alloys (Au₃M with M = Cr, Mn, Fe, Co, and Ni) using first-principles calculations based on spin-polarized density functional theory. Favorable molecular adsorption and greater adsorption stability compared to pure Au are achieved on Au₃M alloys. For these alloys, there is an emergence of unoccupied states in the surface d band around the Fermi level with respect to the fully occupied d band of pure Au. Thus, the derived antibonding state of the sp-d interaction is upshifted and becomes unoccupied compared to pure Au. The B–H bond elongation of the adsorbed borohydride on these alloy surfaces points to the role of surface-parallell (dₓ₂-ᵧ₂ and dₓᵧ) states components of the d-band of the alloying metal M, most pronouncedly in the cases of M = Co or Ni. On the alloy surfaces, B binds directly with the alloying metal, unlike in the case of pure Au where the surface bonding is through the H atoms. These results pose relevant insights into the design of Au-based anode catalysts for the direct borohydride fuel cell.

1. Introduction

Hydrogen has been foreseen as a viable source of energy through hydrogen fuel cells. However, difficulties arise, especially for small-scale power applications, due to issues on hydrogen storage and gas handling safety concerns. Improvements were obtained through the “direct feeding” of liquid hydrogen-rich fuels such as borohydride to the anode. This direct borohydride fuel cell (DBFC) is a very promising power source for the future because of its high power densities for portable power applications. However, its efficiency and power density are limited in part by the lack of an effective catalyst for borohydride oxidation. Numerous experimental studies have been carried out on the oxidation of borohydride on pure (Pt, Pd, Au, Ag, Rh, Ir) and alloy (Pt–M, M = Au, Ag, Ru, Ni, Ir, Bi) catalysts using voltammetric, amperometric, rotating electrode, and in situ mass and FTIR methods. Among the possible candidates for use as DBFC anodes, Au is uniquely capable of producing almost 100% coulombic efficiency. However, high overpotential is necessary to achieve an appreciable rate of borohydride oxidation on the Au surface. Interestingly, recent experiments have shown that alloying Au with 3d transition metals can effectively reduce the overpotential for the oxidation of the borohydride anion and such Au–3d alloys have higher catalytic activity than Au. However, the electronic effect of 3d transition metals as alloying components for these alloy systems on the oxidation of borohydride is not understood. Such fundamental understanding of the interaction of borohydride with surfaces is important to efficiently utilize the high hydrogen density of the borohydride molecule.

First principles calculations based on density functional theory (DFT) were previously shown to corroborate the experimental findings on borohydride oxidation. For the case of Pt(111), borohydride was found to dissociate to BH_ads and 3H_ads fragments (ads denotes surface-bound species). Surface diffusion of H_ads and associative desorption to H₂ are very likely which explains the experimental observation of high H₂ evolution on Pt. For the case of Au, the low adsorption energy of borohydride corresponds to the experimental finding that the oxidation of borohydride requires high overpotential. Thus, DFT studies on the structure and energetics of borohydride on different catalysts can be used for screening desirable catalysts for borohydride oxidation.

In this paper, we provide for the first time, theoretical insights into the electronic effect of alloying Au with 3d transition metals on the adsorption of borohydride and show the importance of an unoccupied 3d orbital on the stability of borohydride on the surface and the orbital-specific effect on borohydride adsorption structure. These results are...
indispensable in the design of Au-based anode catalysts for the direct borohydride fuel cell.

2. Computational model

The adsorption of the borohydride anion (BH$_4^-$) has been experimentally discussed on both noble and non-noble metal catalysts and was found to be accompanied by the simultaneous transfer of an electron to the electrode, as shown by eqn (1).

\[
\text{BH}_4^- \rightarrow \text{BH}_4\text{ads} + e^-
\]

The adsorption of borohydride on metal surfaces can be modeled in an overall neutral unit cell. This model has also been utilized in other related DFT studies for the simultaneous transfer of an electron to the electrode, as shown by eqn (1).

The adsorption of borohydride on Au surfaces can be modeled in an overall neutral unit cell. This model has also been utilized in other related DFT studies for the simultaneous transfer of an electron to the electrode, as shown by eqn (1). The adsorption of borohydride on metal surfaces can be modeled in an overall neutral unit cell. This model has also been utilized in other related DFT studies for the simultaneous transfer of an electron to the electrode, as shown by eqn (1).

The stable configuration of the adsorbed borohydride (BH$_4^{ads}$) was determined by exhausting the possible orientations on the surface of a four-layer slab. This includes the H-up and H-down orientations (Fig. 1) of the tetrahedral borohydride with B at the high symmetry sites on the surface (top, bridge, hcp hollow and fcc hollow sites), and in-plane rotation. The adsorbate and the top two layers of the slab were fully relaxed in all directions while the bottom two layers were held fixed at their bulk structure. Each slab is separated by \( \sim 15.0 \text{ Å} \) of vacuum, which is large enough to avoid the surface atom interaction along the \( z \) axis with neighboring unit cells. The electric dipole correction layer in the vacuum area was used to cut the dipole interactions between the repeated image layer systems. The adsorption energy on each metal was computed by taking the difference between the total energy of the relaxed clean surface and the gas-phase borohydride.

Spin polarized density functional theory calculations were implemented via the Vienna \textit{ab initio} simulation package (VASP). The interaction between ions and electrons was described using the projector augmented wave (PAW) method. Plane wave basis sets were employed with an energy cut-off of 400 eV. The exchange-correlation term was described using generalized gradient approximation (GGA) based on the Perdew–Burke–Ernzerhof (PBE) functional. The surface Brillouin zone integrations were performed on a grid of \((4 \times 4 \times 1)\) Monkhorst–Pack \( k \)-points using a Methfessel–Paxton smearing of \( \sigma = 0.2 \text{ eV} \). A conjugate-gradient algorithm was used to relax the ions into their ground state. The convergence of numerical results with respect to the slab thickness, the kinetic energy cut-off and the \( k \)-point was established.

3. Results and discussion

3.1 Borohydride on Au(111)

We present the interaction of borohydride on a pure Au(111) surface to make relevant comparisons with Au alloys. The calculated lattice constant for fcc Au is 4.18 Å, in excellent agreement with other DFT studies. The stable adsorption configuration of borohydride on Au(111) is shown in the inset of Fig. 2. Borohydride adsorbs on the surface with B at the hollow site and H atoms at the top sites in the H-up orientation. The B–H bond length is 1.26 Å, which is in very close agreement with other DFT studies. This lies within the B–H bond length (1.26–1.49 Å) previously reported for the case of molecularly adsorbed borohydride. The calculated adsorption energies for \( \sim 1/9 \) and \( \sim 1/16 \) monolayer adsorbate coverages are \(-1.69 \text{ eV} \) and \(-1.83 \text{ eV} \) respectively. This shows the enhanced repulsive interaction between the adsorbates as the coverage is increased. The adsorption of borohydride on Au is the weakest among all the transition metals previously studied. This shows the property of Au being the most noble of all metals.
To understand the mechanism of adsorption of borohydride on Au, we analyzed the density of states (DOS) projected on surface Au atoms and borohydride for clean and adsorbed systems. We found that the shifting and broadening of states are prominent only for the d$_{zz}$ state of Au which hybridizes with the H-s state of borohydride. Thus, the molecular structure of borohydride on Au can be described by the s–d$_{zz}$ hybridization of adsorbate–metal states. This is illustrated by the partial charge density distribution projected using a 0.01 electron/\(a_0^3\) isosurface value at an identified bonding state shown in the inset of Fig. 2. Here, the mixing between charge density of the H atoms and the metal is very evident. This hybridization is promoted when borohydride is in the H-up orientation, which explains its preferential stability over the H-down orientation.

3.2 Adsorption energetics on Au alloys

Au is known to form stable alloys such as Au$_3$M (M = Cr, Mn, Fe, Co, Ni) with the L1$_2$ structure. These alloys have been successfully synthesized in the laboratory and examined using a variety of experimental techniques, such as XRD and TEM.$^{55-59}$ The (111) facet of these metals was used to rule out the structural differences between different surfaces and to extract meaningful trends in the properties as a function of the substrate identity. The calculated lattice constants are reported in Table 1, which are in excellent agreement with experiments.$^{55}$

Fig. 3 shows the adsorption configuration of borohydride on these surfaces in a 2 x 2 supercell (4 M and 12 Au atoms per layer). Borohydride tends to “seek” the alloying metal by adsorbing on the surface with B on top of the alloying metal M, and H atoms on the hollow sites. B bonds directly with the alloying metal M unlike in the case of pure Au wherein the H atoms bond directly with Au atoms. These alloying metals are known to be easily oxidized$^{60}$ by transferring electron charge to an electronegative atom. This oxidation can be facilitated by the direct bonding of B with M.

For all these alloys, the adsorption energy has increased by 0.33–0.58 eV compared to the case of pure Au. To explain this, we compared the DOS projected on the surface metal atoms for pure Au and clean Au$_3$Fe slabs as a representative alloy (red and black curves respectively in Fig. 4a). For the case of pure Au, the d band is fully occupied. When this Au–d band hybridizes with the sp state of borohydride, the derived sp–d bonding and antibonding states are both occupied as shown in Fig. 4c (red curve). We have confirmed that the occupied states near the Fermi level (insets 1 and 2 in Fig. 4c) have antibonding characteristics which contribute to a repulsive interaction. On the other hand, for the clean Au$_3$Fe surface (black curve in Fig. 4a), spin-down states around the Fermi level have appeared as indicated by the dashed ellipse. We note that these states are mainly Fe states as shown in Fig. 4b. The lower

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<tr>
<td>Au</td>
<td>4.18</td>
<td>—</td>
<td>1.26</td>
<td>−1.83</td>
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Fig. 4 (a) Surface d band projected on pure Au (red) and clean Au$_3$Fe (black), (b) atom projected surface d band of Au$_3$Fe, and (c) sp states of borohydride adsorbed on pure Au (red) and Au$_3$Fe (black). States near the Fermi level are shown by blue dashed ellipses in (a) and (b).
band in Fig. 4a for Au3Fe is mainly from Au. The derived antibonding state of the sp–d interaction (inset 3 in Fig. 4c) is upshifted and becomes unoccupied unlike on pure Au. The corresponding bonding state below the Fermi level (inset 4 in Fig. 4c) is occupied and is due to the B–Fe bonding interaction.

We note that for the d band projected on the surface alloying metal M of other clean Au3M slabs, there is also an emergence of states near the Fermi level in the same energy range as that for Au3Fe. The same bonding mechanism as for the case of Au3Fe explains the increase in the adsorption energy of borohydride on other Au3M surfaces compared to pure Au.

3.3 Adsorption structures on Au alloys

The B–H bond lengths of borohydride on all these alloys fall within the previously set B–H bond lengths for molecular adsorption of borohydride.33,39 For Au3Cr, Au3Mn and Au3Fe, the B–H bond lengths are almost the same as that for pure Au, while for Au3Co and Au3Ni, the B–H bond lengths are more elongated (Table 1). To understand this, we analyzed the DOS projected on the surface alloying atom M for the clean and adsorbed systems. In Fig. 5, we present the DOS of the Fe atom (left panel) on the surface of Au3Fe (representing Au3Cr, Au3Mn and Au3Fe alloys) and the surface Co atom (right panel) of Au3Co (representing Au3Co and Au3Ni alloys). For all these alloys, the components of the d band that protrude out of the surface (dzz, dx²−y², dz²) interact with borohydride. However, the shift and formation of peaks for “surface-parallel” components of the d band (dxz, dx'y², dyz) are noted only for Au3Co and Au3Ni. This happens because borohydride adsorbs at a closer distance to the surfaces of Au3Co and Au3Ni (B–M bond length in Table 1) compared to the other alloys studied. Thus, the interaction of borohydride with surface-parallel components of the d band leads to further elongation of the B–H bonds on Au3Co and Au3Ni alloys. The magnetic moment of Co has almost diminished upon borohydride adsorption (from 1.905 μB to 0.293 μB). However, this large decrease in the magnetic moment of Co is not observed for Ni (from 0.328 μB to 0.111 μB). Thus, the demagnetization of the substrate can be ruled out as the cause of the shifting of DOS peaks (Fig. 5) for dxz and dx'y² components of the d band of Co and Ni. Further discussions in the magnetic aspects of borohydride adsorption can be found in Ref. 39.

These DFT studies shed light on the vast selections of pure and alloy catalysts for BH4 oxidation. We have shown that molecular adsorption structure with greater adsorption energy compared to pure Au can be achieved on these Au-based alloys. Further studies that involve the calculation of free energies, influence of the solvent, and treatment of an externally applied electrode potential shall be done in the next steps of the research to address the electro-catalytic reactions on these surfaces.

4. Conclusion

The structure and stability of borohydride on Au(111) and Au3M(111) (M = Cr, Mn, Fe, Co and Ni) surfaces were studied using first-principles calculations based on spin-polarized density functional theory. Borohydride adsorbs on Au(111) molecularly at lower adsorption energy compared to other transition metals previously reported. This explains why high overpotential is necessary to achieve an appreciable rate of borohydride oxidation on Au. This molecular adsorption is characterized by the strong hybridization of the H-s state of borohydride with the dzz state of Au. This interaction is promoted when borohydride is adsorbed on the surface with B at the hollow site and H atoms at the top sites. Alloying Au with 3d transition metals results in greater stability of borohydride adsorption compared to pure Au. For these alloys, there is an emergence of unoccupied states in the surface d band around the Fermi level with respect to the fully occupied and low-lying d band of pure Au. Thus, the derived antibonding state of the sp–d interaction is upshifted and becomes unoccupied compared to pure Au. The difference in the B–H bond elongation of borohydride on these alloy surfaces points to the effect of the surface parallel (dxz, dx'y² states) components of the d-band of the alloying metal. On the alloy surfaces, B binds directly with the alloying metal unlike the case of pure Au wherein the surface bonding is through the H atoms. This happens in order to facilitate the oxidation of the alloying metal by transferring an electron to B. These results pose relevant insights into the design of Au-based anode catalysts for the direct borohydride fuel cell.

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Notes and references