Density Functional Theory Study of the Adsorption of Oxygen and Hydrogen on 3d Transition Metal Surfaces with Varying Magnetic Ordering

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ABSTRACT

We have employed density functional theory (DFT) calculations to investigate the adsorption of molecular oxygen and hydrogen on 3d transition metal (TM) surfaces with varying ordered magnetic structures in the bulk, namely ferromagnetic Fe(110), Co(0001), Ni(111) and diamagnetic Cu(111). The trend observed in the energies of adsorption was compared with the magnetic moment of the cell using the d-band centre model of chemisorption and the Stoner model of magnetic energy. As the gap between the d-band centre and the Fermi level of the TM decreases, more antibonding orbitals are present above the Fermi level and thus unoccupied, leading to stronger binding. Correspondingly, the shift in the d-band centre decreases the density of states (DOS) at the Fermi level giving rise to the ordered magnetic structure.

KEYWORDS

D-Band centre, chemisorption, Hedvall effect, magnetism.

1. Introduction

The interactions between gas molecules and transition metal (TM) surfaces are vital to understanding the underlying mechanics of heterogeneous catalysis. Thus, optimizing adsorption sites on TM surfaces has driven a variety of theoretical and experimental studies.1 It is well-understood that the interaction between an adsorbate and TM surface largely depends on the d-electrons of the metal surface. The most widely used descriptor of this interaction is the d-band centre model pioneered by Hammer and Nørskov.2 The d-band centre model, in principle, states that changes to adsorption energies over TM surfaces are related to the shifting of the metal’s d-band centre with respect to its Fermi level. As shown in Fig. 1, the higher the shift of the d-band centre (i.e. smaller ΔE – E dbc), the lower the occupancy of the anti-bonding states, corresponding to a stronger TM-adsorbate binding energy. Bhattacharjee3 has shown that for magnetically polarized TM surfaces, distinct d-band centres should be considered for spin-up and spin-down states. These two d-band centres are shifted in opposite directions relative to the unpolarized d-band centre, resulting in competing contributions (i.e. attractive and repulsive) to the TM-adsorbate orbitals. The extent of this competition depends solely on the magnitude of the TM magnetic polarization. For the purposes of this study, the conventional d-band centre model was used, since the trends observed for the TM of interest would be similar, as shown by Bhattacharjee. The work function (WF) of the metal is another important descriptor to consider during chemisorption. The WF is defined as the difference in energy between an electron in the surface and that of an electron removed to the vacuum4 and thus it can be associated with the minimum work required to facilitate the electron transfer between the metal and the adsorbate. The lowest WF for transition metals are generally found on the least densely packed surface.5

Another important property of TM, which also largely depends on the d-electrons, is their magnetic ordering (i.e. paramagnetic, ferromagnetic, diamagnetic). The role of the ordered magnetic structure, temperature-dependent magnetic-order transitions and bulk magnetic domain formation during the chemisorption of adsorbates is less well-understood. This is further complicated by the variation in magnetic ordering of the TM bulk structure compared to the surface where chemisorption occurs.7

The magnetic moment of 3d TM can be described by quantum formulations of electron spin and orbital contributions of...
individual atoms that comprise the bulk structure. Interestingly, spin-orbital coupling has been shown to make a negligible contribution to the atomic magnetic moments. This description, however, is difficult to relate to the chemisorption of adsorbates since the adsorption energy, as mentioned above, is easily described in terms of the d-band energy. A more useful descriptor to relate trends in chemisorption with magnetic ordering is the energy band theory of magnetism (or Stoner Mean-Field theory). In the Stoner model, the magnetically ordered structure is described in terms of the DOS at the Fermi level, $D(E_F)$, and an exchange integral parameter (or Stoner parameter, $I$) which in turn depends on the number of d-electrons. The Stoner criterion, described as $D(E_F)I > 1$, is a measure by which the ferromagnetic ordering of a TM can be determined. As shown in Fig. 2, the $D(E_F)$ for the 3d TM increases from Mn to Ni; however, only Fe, Co and Ni exceed the Stoner criterion.

![Figure 2](image)

The role of TM magnetic ordering in the kinetics and thermodynamics of chemisorption is crucial to the understanding of the magneto-catalytic effect. The magneto-catalytic (or Hedvall) effect is a contested theory based on early experimental and theoretical studies, which suggests that the magnetic ordering of a metal can influence the catalytic properties of a reaction. The initial experimental studies were conducted by Hedvall, showing that the activity of nickel catalysts for the decomposition of $\text{N}_2\text{O}$ could be manipulated by varying the temperature-dependent magnetic ordering of the bulk structure. Early theoretical work conducted by Iliszca and Selwood focused on the application of magnetic fields for ortho-para hydrogen conversions. A handful of theoretical works have surfaced in recent years on the role of magnetism in catalysis, which provide further evidence of Hedvall’s theory.

Melander et al. computed the adsorption of $\text{H}_2$ and CO on face centred cubic (fcc) Fe(111) surfaces with different magnetic configurations, namely ferromagnetic and antiferromagnetic structures. The hypothetical variation of the magnetic ordering results in the orbital occupancy of the surface differing due to the variation of the spin structures. The molecular adsorption study indicates that there is charge transfer between the iron surface and either adsorbate, which is greater for the bilayer system with the antiferromagnetic order than with the ferromagnetic configuration. In addition, the dissociation of $\text{H}_2$ is easier on the bilayer with the antiferromagnetic order, whereas CO dissociation is more difficult on the ferromagnetic surface.

The influence of magnetic ordering in surfaces on the molecular adsorption behaviour is also observed when the adsorbate has a net magnetic moment, as is the case for oxygen. During the oxygen evolution reaction, the spin conversion from singlet-state oxygen in water to triplet state molecular oxygen is considered the rate-limiting step for water splitting. This is a spin-forbidden isogyric reaction, where a heterogeneous catalyst is needed to allow non-conserving spin reactions. For example, Lim et al. showed that the antiferromagnetic ordering of perovskite catalysts was important for facilitating this conversion. Similarly, Torun et al. showed that the presence of magnetic moments in the RuO2 (110) surface facilitates the selection rule for the spin conversion and evolution of the triplet state molecular oxygen. This spin conversion can be further amplified by weak magnetic fields (<0.4 T) applied to the magnetic anodes of alkaline water electrolysis.

In this study the adsorption properties of selected 3d-TM surfaces with varying magnetic ordering is compared using molecular oxygen and hydrogen. Density functional theory (DFT) calculations are used to model the magnetic structures (i.e. non-paramagnetic ordering) and molecular adsorptions to the surfaces. Iron, cobalt and nickel are ferromagnetic materials, while copper is diamagnetic. The bcc-Fe(110), hcp-Co(0001), fcc-Ni(111) and Cu(111) surfaces are investigated since these planes are essentially the same or (as in the case of Fe) similar, thus minimizing the contribution of varying lattice.

2. Methodology

Spin-polarized DFT calculations were performed using the Perdew–Burke–Ernzerhof (PBE) exchange correlation functional within the generalized gradient approximation (GGA). The Vienna Ab Initio Simulation Package (VASP) was employed for the optimizations of the atomic positions and lattice parameters of the metal bulk phases and the position of the exposed atoms in the surfaces. An electronic step convergence of 10$^{-5}$ eV was used, along with a tight atomic force threshold of 0.01 eV/A. The projector augmented wave pseudopotentials were used to treat the core electrons and their effect on the valence electrons. A plane-wave basis set has been used with a kinetic energy limit of 400 eV and a Monkhorst Pack k-points mesh of $7 \times 7 \times 1$ to sample the Brillouin zone. Dipole corrections were applied in the direction perpendicular to the surface. The formalism of Grimme with the Becke-Johnson damping (D3-BJ) was applied to account for the long-range dispersion corrections.

The Methfessel-Paxton method order of 1 was used to determine the electronic partial occupancies with the width of the smearing set at 0.02 eV. To simulate the Fe(110), Co(0001), Ni(111) and Cu(111) surfaces, a set of slab models were constructed using $(1 \times 1)$ supercells keeping a vacuum region of 10 Å normal to the surface. The energy of adsorption $(E_{\text{ads}})$ of a single oxygen molecule in the triplet spin state or hydrogen molecule in equilibrium with a reservoir of oxygen or hydrogen, respectively, was calculated according to:

$$E_{\text{ads}} = E_{\text{tot}} - E_{\text{M}} - \frac{1}{2}E_{\text{O}_2}$$

where $E_{\text{tot}}$ is the total energy of the TM-adsorbate system, $E_{\text{M}}$ is the optimized energy of the TM surface and $E_{\text{O}_2}$ is the energy of a single O or H atom in the diatomic $\text{H}_2$ or $\text{O}_2$ molecules, respectively, accounting for over-binding in the case of $\text{O}_2$. The magnetic moment of the cell was also obtained and correlated with the energy of adsorption for each TM adsorption.
3. Discussion

Both hydrogen and oxygen adsorb dissociatively to the metal surfaces. The adsorption energy of oxygen (see Fig. 3 bottom panel) is lower than hydrogen across most surfaces with the largest difference observed for Co(001). The trend across the metals agree previous literature reports, generally growing more positive from iron to copper. The magnetic moment of the cell for each metal-adsorbate is inversely proportional to the adsorption energies, decreasing from iron to copper.

The trends observed in the adsorption energies and magnetic moments can both be explored as functions of the DOS at the Fermi level. As the number of d-electrons increases from iron to copper, the d-band centre shifts further from the Fermi level. As the adsorption energies, respectively (Fig. 4). As discussed above, using the Stoner Mean Field theory, the DOS at the Fermi level determines the magnetic ordering of the TM. As the adsorption energy decreases the local DOS at the Fermi level, a reduction in the local magnetic moment is also observed.

This study has sought to explore the trends observed in chemisorption and magnetic ordering from a common fundamental property of TM. An inverse relationship between chemisorption and magnetic ordering was observed for the 3d TM, which can both be rationalized using theories founded on the DOS at the Fermi level. This is a useful basis from which to understand experimental theories of magneto-catalysis currently under investigation.

Figure 3 Trends in (top) the work function$^{14}$ and (bottom) magnetic moment per atom and adsorption energy of oxygen and hydrogen.$^{22}$

Figure 4 DOS for fcc-Cu(111) and hcp-Co(0001), dashed line indicates the Fermi level.

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