First-principles investigation of the stability of 3d monolayer/Fe(001) against bilayer formation

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Based on the full-potential linearized augmented plane-wave method combined with the generalized gradient approximation, we determine the ground-state spin configurations and the total energies of 3d transition-metal monolayer and bilayer films on Fe(001) within the c(2×2) unit cell. We find by energy analysis that V, Cr, and Mn layers prefer the layered antiferromagnetic coupling, and Fe, Co, and Ni layers favor the ferromagnetic coupling to Fe(001). One exception is the Mn monolayer, which favors the c(2×2) ferrimagnetic superstructure. We discuss the stability of the 3d transition-metal monolayer films on Fe(001) against the bilayer formation and find that, with the exception of Cr, all 3d monolayers on Fe(001) are stable against bilayer formation. We have confirmed that the interlayer relaxations do not change the overall features of the present results.

I. INTRODUCTION

3d metal monolayers (ML) on Fe(001) are prototypical systems where in-plane magnetic interactions compete with strong magnetic interactions between the ML and the Fe substrate. In particular the magnetism of ultrathin films of Cr and Mn on Fe(001) are under intense scrutiny for the understanding of the interfacial magnetism. The experimental results are still partly in contradiction and the complexity of those systems have been gradually revealed. For a Cr ML the measured magnetic moment was found to be at most 1 \( \mu_B \), which is less than half of the theoretical prediction. It was also reported that a layer-by-layer growth leads to a strong intermixing with the substrate Fe layers. Mn overlayers seem to be even much more involved. There is a general experimental consensus that for Mn around 1 ML coverage the signals related to the magnetization disappear. The microscopic origin for this observation is under strong debates. The difficulty in controlling and characterizing the morphology of the interface seems to be intimately related to the difficulties in the understanding of the interfacial magnetism of those systems.

This article is dedicated to shed some light on the aspect of the stability of the 3d metal MLs on Fe(001) with respect to the bilayer (BL) formation. So far no ab initio calculations on the 3d overlayers on Fe(001) have focused on the stability arguments of these systems. We calculated the magnetic moments, the magnetic structures, and the total energies for two types of 3d metal systems: (i) the 3d MLs on Fe(001) and (ii) the 3d BLs on Fe(001), and in addition the plain Fe(001) without 3d overlayers. Each type of systems includes the 3d metals V, Cr, Mn, Fe, Co, and Ni. By comparing the total energy of MLs on Fe(001) with the one of BLs covering 50% of the Fe surface and leaving 50% uncovered, we extract the BL formation energy as a measure of the stability of the 3d MLs on Fe(001) against BL formation during growth or after annealing. Since what occurs in real growth would be strongly governed by the energy barriers of elementary processes, we expect that the examination of such total energy aspects will give a good estimation of how systems evolve for systems where the energy barriers are sufficiently low. Therefore, we may also expect that comparisons of the present examinations with experimental growth behaviors will give some indications of such barrier heights.

II. METHOD

The calculations were carried out with the full-potential linearized augmented plane-wave (FLAPW) program FLEUR in film geometry combined with the generalized gradient approximation (GGA) proposed by Perdew et al. Two film geometries were considered, the first consisted of seven bcc Fe(001) layers and one 3d metal ML on each side of the Fe(001) surface, and the second consisted of seven Fe(001) layers with two 3d metal MLs on each side. We searched for the ground-state spin configurations within a c(2×2) unit cell excluding noncollinear magnetic structures. Prior to the ML and BL calculations we determined the theoretical (GGA–FLAPW) equilibrium lattice constant of FM bcc Fe as \( a_0 = 5.33 \text{ a.u.} \), which is smaller by 1.5% as compared with the experimental value of 5.41 a.u. Thus, the calculations are carried out with the in-plane lattice constant of 5.33 a.u. and results of structurally unrelaxed calculations use the interlayer distance of 2.665 a.u. Converged total energies were obtained by use of roughly 70 symmetrized augmented plane waves per atom as a variational basis set and 36 and 21 special \( k \) points for the \( p(1\times1) \) and \( c(2\times2) \) unit cells, respectively, to integrate over the irreducible wedge of the two-dimensional Brillouin zone. To determine the magnetic ground-state configurations of each system we have examined all conceivable spin configurations. The interlayer dis-

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For the BL systems there are eight magnetic configurations being a transient element between an AF and a FM coupling reduced making a distinguished dip. This is a result of Mn Mn MLs the magnetic moment of the interface Fe atom is flipped to the Fe substrate, emphasized by open filled symbols. Shown are three spin configurations, $p(1 \times 1)$ FM (solid line), $p(1 \times 1)$ LAF (dotted line), and the $c(2 \times 2)$ FI (dashed lines) for positive and negative moments. Included are the moments of interface Fe (chained line) for the ground-state spin configurations.

III. RESULTS

We show in Fig. 2 the magnetic moments for FM, LAF, and FI spin structures of the unrelaxed 3$d$ MLs on Fe(001). The magnetic moments of the interface Fe atom for ground-state spin configurations are also shown. The magnetic ground-states are LAF for V and Cr, FI for Mn, FM for Fe, Co, and Ni, as has been reported in the literature.\textsuperscript{3,4,10} For Mn MLs the magnetic moment of the interface Fe atom is reduced making a distinguished dip. This is a result of Mn being a transient element between an AF and a FM coupling between early and late 3$d$ transition-metal elements, respectively, as discussed by Mirbt et al.\textsuperscript{3} Here the size of the dip at Mn is a little bit larger than the one calculated by Mirbt et al. owing to the present incorporation of the $c(2 \times 2)$ configuration. The total energy differences between FM and LAF configurations and between FM and FI show some differences to those of Handschuh and Blügel\textsuperscript{4} coming mostly from the different choice of the in-plane lattice constant. Extending the search for the magnetic ground-state of Mn to larger surface unit-cells will most likely lead to a more complicated spin-structure\textsuperscript{11–13} not discussed here any further.

For the BL systems we have examined all possible magnetic configurations within the $c(2 \times 2)$ unit cell. With the exception of Mn all stable magnetic solutions found have the $p(1 \times 1)$ structure. The magnetic ground-states of the 3$d$ BLs are the $p(1 \times 1)$ LAF one, ($\uparrow \downarrow \uparrow \downarrow$) for Fe, Co, and Ni, and the $p(1 \times 1)$ FM one for Fe, Co, and Ni, as seen in the signs of magnetic moments shown in Fig. 3. For the case of the LAF Mn BL we observe an almost vanishing magnetic moment of the substrate Mn atom. For Mn, in addition to the ground-state structure, we found another LAF configuration, ($\downarrow \uparrow \downarrow \uparrow$), about 9 meV/Mn higher in energy than the ground state, which is the one found by Wu and Freeman.\textsuperscript{13} Furthermore other than these two LAF structures we found a superstructure ($\uparrow \uparrow \uparrow \uparrow$) as a stable solution with the total energy higher than the ground state by 43.2 meV/Mn. Thus, there exist three states within an energy range corresponding to about 400 K, which seems to show already a part of difficulties involved in the calculation of thicker Mn overlayers on Fe(001).\textsuperscript{14}

In Fig. 4 we show the BL formation energy defined as the total energy difference of three systems, $E_{\text{ML/Fe}} - 1/2[E_{\text{BL/Fe}} + E_{\text{Fe(001)}}]$, whose negative value corresponds to the case where ML formation is preferred. The figure indicates that, with the exception of Cr, all 3$d$ transition metals favor the ML formation over the BL formation. Though this overall trend is already visible in the nonmagnetic results, the strong magnetic effect for Mn changes the sign of its BL formation energy. The results seem to meet the experimental observation on the Cr overlayer where its ML is unstable.\textsuperscript{5} For other 3$d$ overlayers ML stability assessment requires...
extension of the present investigations to, e.g., the effect of interdiffusion, which are now in progress.

IV. CONCLUSION

In summary, we have determined by ab initio total energy and force calculations the ground-state spin configurations and the total energies of 3d transition-metal ML and BL films on Fe(001). We found that for both MLs and BLs V, Cr, and Mn prefer the p(1×1) LAF configuration and Fe, Co, and Ni the p(1×1) FM with a single exception of the Mn ML which prefers a FI structure within the restricted choice of a c(2×2) unit cell. We found that all 3d elements favor the ML over the BL except for Cr. The present result of the Cr ML instability against BL formation may meet the experimentally found instability of the Cr ML.

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FIG. 4. Shown is the bilayer formation energy $\Delta E$ per 3d transition-metal atom for unrelaxed geometries. For $\Delta E<0$ ($\Delta E>0$), ML (BL) is energetically preferred. The solid (dashed) line denotes results derived from the (non)magnetic ground-state spin configuration within the $c(2\times2)$ unit cell. The interlayer relaxations do not change the overall features of this figure.