Interface properties of the NiMnSb/InP and NiMnSb/GaAs contacts

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We study the electronic and magnetic properties of the interfaces between the half-metallic Heusler alloy NiMnSb and the binary semiconductors InP and GaAs using two different state-of-the-art full-potential ab-initio electronic structure methods. Although in the case of most NiMnSb/InP(001) contacts the half-metallicity is lost, it is possible to keep a high degree of spin-polarization when the interface is made up by Ni and P layers. In the case of the GaAs semiconductor the larger hybridization between the Ni-d and As-p orbitals with respect to the hybridization between the Ni-d and P-p orbitals destroys this polarization. The (111) interfaces present strong interface states but also in this case there are few interfaces presenting a high spin-polarization at the Fermi level which can reach values up to 74%.

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I. INTRODUCTION

The spin-injection from a metal into a semiconductor remains one of the main challenges in the field of magnetoelectronics. The use of half-metallic ferromagnets as electrodes was proposed to maximize the efficiency of spintronic devices. These compounds are ferromagnetic metals with a band gap at the Fermi level (EF) for the minority spin band leading to 100% spin-polarization at EF. But even in this case, interface states at the contact between the half-metal and the semiconductor can destroy the half-metallicity. Due to their orthogonality to all bulk states incident to the interface, in the ballistic limit these states should not affect the transport properties, but it is their interaction with other defect states which makes them conducting.

The first material predicted to be a half-metal is the Heusler alloy NiMnSb. There exist several ab-initio calculations on NiMnSb reproducing the initial results of de Groot and collaborators, and Galanakis et al. showed that the gap arises from the hybridization between the d orbitals of the Ni and Mn atoms. Experiments seem to well establish its half-metallicity in the case of single crystals, but in films the half-metallicity is lost. Theoretical calculations for the interfaces of these materials with the semiconductors are few and all results agree that in general the half-metallicity is lost at the interface between the Heusler alloy and the semiconductor. Wijs and de Groot have argued that in the case of the NiMnSb/CdS (111) contacts the Sb/S interface keeps the half-metallicity (or at least shows a very high degree of spin-polarization) of the bulk NiMnSb. Thus, even if half-metallicity is lost, it is possible that a high degree of spin-polarization stays at the interface and these structures remain attractive for realistic applications.

We should also mention that even in the absence of the interface states true half-metallicity can not really exist due to minority states induced by the spin-orbit coupling which couples the two spin-bands. But as it was shown for these systems in Refs. and this phenomenon is very weak and instead of a gap in the minority channel there is a region of still almost 100% spin-polarization. It was also found that the orbital moments are very small in these compounds. Thus, spin-orbit coupling can be assumed to be negligible with respect to the interface states.

In this communication we study the (001) interfaces of the half-metallic NiMnSb Heusler alloy with InP and GaAs and the (111) interface between the NiMnSb and InP compounds. We take into account all possible contacts and show that there are cases where a high degree of spin-polarization remains at the interface. In Section III we discuss the structure of the interfaces and the computational details and in Section IV we present and analyze our results for the (001) interfaces. In Section V we discuss the (111) interfaces and finally in Section VI we summarize and conclude.

II. COMPUTATIONAL METHOD AND STRUCTURE

In the calculations we employed different full-potential methods. Firstly we employed the full-potential version of the screened Korringa-Kohn-Rostoker (FSKKR) Green’s function method in conjunction with the local spin-density approximation (LDA) to the density functional theory to study the (001) interfaces between NiMnSb and the InP and GaAs semiconductors. The FSKKR method scales linearly with the number of atoms and, therefore, allows to study also very thick slabs of these materials. But it cannot give exactly the Fermi level of semiconductors due to problems arising from the orbital moments are very small with respect to the interface states.

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cell is that of a fcc lattice with four atoms as basis at A=(0 0 0), B=(1 1 1), C=(2 2 2) and D=(3 3 3) in Wyckoff coordinates. In the case of NiMnSb the A site is occupied by Ni, the B site by Mn and the D site by Sb, while the C site is unoccupied. The C1b structure is similar to the L21 structure adopted by the full Heusler alloys, like Ni2MnSb where also the C site is occupied by a Ni atom. The zincblende structure, adopted by a large number of semiconductors like GaAs and InP, can also be considered as consisting of four fcc sublattices. In the case of GaAs the A site is occupied by a Ga atom, the B site by an As atom, while the C and D sites are empty. Depending on the electronic structure method used to perform the calculations one either uses empty spheres or empty polyhedra to account for the vacant sites (as it is done in the FSKKR) or the vacant sites just make part of the interstitial region (as in FLAPW). Within 1% accuracy NiMnSb (5.91Å) has the same experimental lattice constant as InP (5.87Å) and epitaxial growth of NiMnSb on top of InP has been already achieved experimentally by molecular beam epitaxy. On the other hand, the lattice constant of GaAs (5.65Å) is almost 4% smaller. The dominant effect at the interface is the expansion or the contraction of the lattice of the half metal along the growth axis to account for the in-plane change of its lattice parameter. Since in the case of the NiMnSb/InP interface both compounds have similar lattice parameters, in the calculations perfect epitaxy can be assumed.

Within the FSKKK the space is divided into non-overlapping Wigner-Seitz polyhedra and thus empty ones are needed to describe accurately the vacant sites (similarly to the use of empty spheres in the early electronic structure methods). To simulate the (001) interface within the FSKKR calculations we used a multilayer consisting of 15 layers of the half-metal and 9 semiconductor layers. This thickness is enough so that the layers in the middle of both the half-metallic part and the semiconductor one exhibit bulk properties. There are several combinations at the interface, e.g. at the NiMnSb/InP contact the interface can be either a Ni/In one, Ni/P, MnSb/In or MnSb/P (see Fig. 1). We will keep this definition through out the paper to denote different interfaces. We should also mention that since the multilayer contains 15 half-metal and 9 semiconductor layers, there are two equivalent surfaces at both sides of the half-metallic spacer. Finally, for our FSKKR calculations we used a 20×20×4 grid in the k-space and we took into account wavefunctions up to ℓmax=3 and thus the potential and the charge density were expanded on lattice harmonics up to ℓmax=6. All FSKKR calculations have been performed at the experimental lattice constant of NiMnSb (5.91Å).

In the FLAPW method the space is divided into non-overlapping muffin-tin spheres around each atom and an interstitial region, that is described in terms of plane waves. To perform the calculations for the (001) interfaces we employed a repeated slab made up of 8 layers of NiMnSb and 8 layers of the semiconductor. Thus if the one contact is Ni/P the other one is MnSb/In. As will be shown in subsection III A the smaller number of layers (as compared to the FSKKR calculations) does not influence the properties near the Fermi level. For the (111) interfaces the supercells consisted of 16 layers of NiMnSb and 12 layers of InP. The FLAPW calculations were performed using density functional theory in the generalized gradient approximation (GGA) as given by Perdew et al. For the calculations, a planewave cutoff Kmax=3.45 a.u. was used. Lattice harmonics with angular momentum ℓ ≤ 8 were used to expand the charge density and the wavefunctions within the muffin-tin spheres, having a radius of 2.4 a.u. for Sb and 2.34 a.u. for all the other atoms. The Brillouin-zone (BZ) was sampled with 128 special k-points in the irreducible wedge (1/8 of the whole BZ) for (001) interfaces, and 90 k-points in the irreducible wedge (1/12 of the whole BZ) for the (111) interfaces. All FLAPW calculations were performed at the experimental lattice constant of InP (5.87 Å).

III. NiMnSb/InP AND NiMnSb/GaAs (001) INTERFACES

Compared to simple surfaces, interfaces are more complex systems due to the hybridization between the orbitals of the atoms of the metallic alloy and the semiconductor at the interface. Thus, results obtained for the surfaces (as the ones in Refs. 11 and 12) cannot be easily generalized for interfaces since for different semiconductors different phenomena can occur. In both (001) and (111) surfaces of NiMnSb, the appearance of surface states destroys the half-metallicity. In Sections III A and III B we present the FSKKR results for the NiMnSb/InP(001) and NiMnSb/GaAs(001) contacts, respectively, and in Section III C we give the valence band offsets calculated with the FLAPW method for the NiMnSb/InP(001) interfaces and compare the results obtained with the two different methods.

FIG. 1: Schematic representation of the (001) interface between NiMnSb and InP. There are several different combinations at the interface which can be either Ni/In, Ni/P, MnSb/In (shown in the figure) or MnSb/P.
TABLE I: FSKKR-calculated atomic spin moments given in $\mu_B$ for the interface between the MnSb-terminated (001) NiMnSb and the In or the P terminated InP. We do not present the spin moments at the vacant sites. Last columns are the moments for the MnSb-terminated NiMnSb(001) surface and the bulk NiMnSb. “I” denotes the interface layers and $\pm$ means one layer deeper in the half-metal or the semiconductor.

|         | MnSb/In | MnSb/P | MnSb surf. | bulk     |
|---------|---------|--------|------------|----------|
| I-3     | Ni: 0.270 | Ni: 0.275 | Ni: 0.269  | Ni: 0.264 |
| I-2     | Mn: 3.704 | Mn: 3.734 | Mn: 3.674  | Mn: 3.705 |
| I-2     | Sb: -0.057 | Sb: -0.044 | Sb: -0.066 | Sb: -0.062 |
| I-1     | Ni: 0.289 | Ni: 0.316 | Ni: 0.223  | Ni: 0.264 |
| I      | Mn: 3.405 | Mn: 3.718 | Mn: 4.018  | Mn: 3.705 |
| I-2     | Sb: -0.037 | Sb: -0.045 | Sb: -0.096 | Sb: -0.062 |
| I+1     | In: -0.053 | P: 0.015 |
| I+2     | In: -0.017 | P: 0.011 |
| I+3     | P: -0.012 | In: 0.002 |

A. NiMnSb/InP contacts

The first case which we will study are the interfaces between NiMnSb and InP. In Table I we have gathered the FSKKR spin moments for the case of the MnSb/In and MnSb/P interfaces. “I” stands for the interface layers, +1 means moving one layer deeper in the semiconductor and $-1$ one layer deeper in the half-metallic spacer. In the case of the MnSb terminated half-metallic film there is a difference depending on the semiconductor termination. In the case of the In termination the Mn spin moment decreases considerably and is now 3.4 $\mu_B$ compared to this surface, for the interfaces between MnSb-terminated NiMnSb and InP the situation is completely different. The hybridization between the half-metallic film, the spin moments regain theirbulklike behavior while, if we move deeper in the semiconductor film, the induced spin moments quickly vanish.

On a MnSb-terminated (001) surface the spin polarization at the Fermi level, $E_F$, was found to be as high as 38% (the spin-polarization is defined with respect to the density of states $n(E)$: $P = \frac{n(\uparrow(E_F)) - n(\downarrow(E_F))}{n(\uparrow(E_F)) + n(\downarrow(E_F))}$ where $\uparrow$ stands for the majority electrons and $\downarrow$ for the minority electrons). In Ref. 43 two surface states at $E_F$ were reported to destroy the half-metallicity, but still the population of the majority electrons at the Fermi level was twice as large as the one of the minority states. Compared to this surface, for the interfaces between MnSb-terminated NiMnSb and InP the situation is completely different. The hybridization between the half-metallic film and the semiconductor is strongly reduced and the density of states at the Fermi level is almost zero. This is clearly seen in Fig. 2, where we present with the dashed line the spin and atom resolved density of states (DOS) of the MnSb-P contact. There is a minority interface state pinned at the Fermi level which destroys the half-metallicity. In the Mn local DOS, this state overlaps with the occupied minority Mn states and it is not easily distinguished but its existence is obvious if one examines the Ni and Sb DOS. The situation is similar also for the MnSb/In contact not shown here.

In the case of the Ni terminated NiMnSb films, DOS at $E_F$ is more bulklike than the case of the MnSb films. Already Ni interface atom has a spin moment of 0.29 $\mu_B$ in the case of an interface with In and 0.36 $\mu_B$ for an interface with P compared to the bulk value of 0.26 $\mu_B$. In
the bulk case Ni has 4 Mn and 4 Sb atoms as first neighbors. On the Ni-terminated (001) surface the Ni atom loses half of its first neighbors. But if an interface with P is formed, the nickels two lost Sb neighbors are replaced by two isovalent P atoms and – with the exception of the Mn neighbors – the situation is very similar to the bulk. Now the Sb $p$ bands at lower energy are not destroyed since P has a behavior similar to Sb and still they accommodate three transition metal $d$ electrons. Thus the only change in the DOS comes from the missing two Mn neighboring atoms. The DOS in Fig. 2 for the Ni/P case is clearly very close to the bulk case and in Fig. 3 we have gathered the DOS for the Ni and the void at the interface and the Mn and Sb atoms at the subinterface layer for both Ni/In (dashed line) and Ni/P (solid line) contacts and we compare them with the bulk results from Ref. 10. In the case of the Ni/In interface there is an interface state pinned at the Fermi level which completely suppresses the spin polarization, $P$ (if we take into account the first two interface layers is $P \approx 0$). In the case of the Ni/P interface the intensity of these interface states is strongly reduced and now the spin-polarization for the first two interface layers is 39%, i.e. about 70% of the electrons at the Fermi level are of majority spin character.

B. NiMnSb/GaAs contacts

In the previous section it was shown that in the case of the Ni/P interfaces the spin-polarization was as high as 39%. In order to investigate whether this is a general result for all semiconductors or specifically for this interface we also performed calculations for the case of the NiMnSb/GaAs(001) contacts using the same lattice parameter as for the previous ones; thus the lattice constant of GaAs was expanded by approximately 4%.

In the top panel of Fig. 4 the atom-resolved DOS for the Mn at the interface layer is shown for the case of the MnSb/In semiconductor interfaces. The hybridization between the $d$-orbitals of the Mn atom at the interface and the $p$-orbitals of the $sp$ atoms of the semiconductor is larger in the case of the InP spacer. This leads to an about 0.1-0.2 $\mu_B$ smaller Mn spin moments at the interface and the exchange splitting between the occupied Mn majority and the unoccupied Mn minority $d$-states is smaller. Thus the large minority peak above the Fermi level moves lower in energy and now strongly overlaps with the occupied minority peak below the Fermi level increasing the minority DOS at the Fermi level. The final spin-polarization at
Employing the FLAPW method, we calculated the (minority states) valence-band offset which is the energy difference between the maximum of the valence band (VBM) of the semiconductor and the maximum of the minority valence band of the Heusler alloy. To calculate it we referenced the binding energies the core states in the interface calculation to their corresponding bulk values as described in Ref. 44. We found that the VBM of the semiconductor is 0.83 eV lower than the one of the half-metal for the In/MnSb contact. For the other interfaces the valence band offsets are: 0.69 eV for the In/Ni, 0.69 for the P/Ni and 0.80 eV for the P/MnSb contact. In the bulk InP semiconductor the experimental gap is 1.6 eV, thus the Fermi level, which is 0.07 eV above the maximum of the minority NiMnSb valence band falls in the middle of the semiconductor bulk bandgap. This is similar to what is happening also in the case of the Co2MnGe/GaAs (001) interfaces and these junctions can be used to inject spin-polarized electrons in the semiconductor.

We can now also compare the results obtained with the FLAPW with the results from the FSKKR calculations. In the left panel of Fig. 4 we present the DOS of the Mn atom at the MnSb/In and MnSb/P interfaces together with the bulk FLAPW calculations while in the right panel of the same figure we present the DOS for the Ni atom at the Ni/P and Ni/In interfaces. We can directly compare these results with the FSKKR results on the same systems shown in the left-top and left-down panels of Fig. 4. Except for very small details both methods give a similar density of states. In the case of the MnSb/In interface the Fermi level falls within a local minority minimum while for the MnSb/P interface, due to the smaller exchange splitting Mn unoccupied minority states move lower in energy crossing the Fermi level.

The same effect occurring for the MnSb interface can also be seen at the Ni interfaces, as shown in the bottom panels of Fig. 4. The stronger hybridization of the Ni atom with either the Ga or As atoms at the interface with respect to the InP semiconductor provokes a movement of the Ni unoccupied minority d states towards lower energies while the occupied majority ones are moving higher in energy. If one looks in detail at the Ni/In and Ni/Ga contacts, one observes that the minority peak at the Fermi level present in the Ni/In contact is now smeared out in the case of the Ni/Ga contact due to the unoccupied minority states which move lower in energy. Similarly the unoccupied Ni minority d-states have a larger bandwidth in the case of the Ni/As contact than in the case of the Ni/P one inducing a high minority Ni DOS at the Fermi level. The high spin-polarization at the Fermi level presented in the case of the Ni/P interfaces is completely destroyed in the case of the Ni/As contact due to the larger hybridization between the Ni-d and As-p orbitals with respect to the hybridization between the Ni-d and P-p orbitals. Thus, the properties of the interface depend also in a large extent on the choice of the semiconductor.

C. Band offsets and partial DOS for NiMnSb/InP contacts

FIG. 5: Left panel: Atom- and spin-resolved DOS for the case of MnSb/In(001) (dashed line) and MnSb/P(001) (solid line) contacts for the Mn atom at the interface layer as calculated with the FLAPW method. Right panel: Results for the Ni atom at the Ni/In and Ni/P interface for the Ni-terminated half-metallic spacers. The thin solid line indicates the local DOS of bulk NiMnSb.
axis perpendicular to the interface). Similarly, the bottom graph contains the results for the MnSb/P and Ni/In interfaces. The layers at the middle of the semiconductor spacer show a small DOS due to both the induced states from the half-metal and bulk NiMnSb states which decay slowly outside the half-metallic spacer and travel throughout the semiconductor. It is clearly seen that none of the interfaces is in reality half-metallic. For the MnSb/In interface the Mn atom at the interface shows an almost zero net spin-polarization while the Mn atom at the MnSb/P interface shows a quite large minority DOS as we have already discussed. In the case of the Ni/In interface shown in the bottom panel of Fig. 6 the net spin-polarization is also almost zero as was the case for the FSKKR results. The Ni/P interface shows a spin-polarization $P$ around 40% due to the high polarization of the Ni atom at the interface which polarizes the P atom at the interface also presenting a high majority DOS at the Fermi level. This value is almost identical to the 39% calculated within the FSKKR method as discussed in Section III A.

Although different types of slabs are used to describe the interfaces and different approximations to the exchange-correlation potential are employed both FSKKR and FLAPW calculations lead to very similar results. The latter fact confirms the results in Ref. 9, where it was shown that for the same lattice constant both LDA and GGA reproduce the same electronic and magnetic properties for the Heusler alloys.

IV. NiMnSb(111)/InP(111) INTERFACES

In the last section we will discuss our FLAPW results for the NiMnSb/InP (111) interfaces. As mentioned in Section II for these calculations 16 layers of NiMnSb and 12 layers of InP have been used. Along the [111] direction the semiconductor is composed by pure alternating In and P layers and, thus, our semiconducting spacer is ending in P on the one side and In on the other side. In the case of the half-metallic alloy the structure could be understood easier if we also assume that there is a vacant site in the bulk structure (these “voids” have been explicitly included in in the FSKKR calculations as described in Section II and Ref. 42). For the Mn termination, as we proceed from the interface deeper into the half-metallic spacer, the succession of the layers can be either Mn-NiSb-Void-Mn-... or Mn-Void-Sb-Ni-Mn-... . We denote the two different terminations here as Mn-Ni-Sb-Mn-... or Mn-Sb-Ni-Mn-... . Similarly for the Sb terminated interface we can have either Mn or Ni as subinterface layer and for the Ni termination we can have either Sb or Mn at the subinterface layer. Since we have 16 layers of the half-metallic alloy, we will have from both sides the same layer at the interface, e.g. Mn, but with different subinterface layers, e.g. Sb from one side and Ni from the other.

In Fig. 7 the layer-resolved partial DOS for the Ni-terminated interfaces are shown. In the top panel are the ...-In-P/Ni-Sb-Mn-... and ...-P-In/Ni-Mn-Sb-... contacts and in the bottom panel the ...-In-P/Ni-Mn-Sb-... and ...-P-In/Ni-Sb-Mn-... ones. As it was shown in Ref. 42, in the case of the Ni and Mn terminated (111) surfaces there are strong surface states pinned at the Fermi level which also penetrate deeply into the subsurface layers. These surface states are present also in the case of the interfaces studied here, although their intensity decreases slightly due to the hybridization with the sp atoms of the semiconductor. In all cases the net spin-polarization of the Ni atom at the interface is very small with the exception of the ...-In-P/Ni-Mn-Sb-... interface (middle of the bottom panel). For this case the simultaneous presence of the P atom from the one side and of the Mn atoms at the subinterface layer create an atomic-like environment for Ni similarly to what happened in the case of the Ni/P(001) contact and the spin-polarization, taking into account the two semiconductor layers at the interface and three first NiMnSb layers, is as high as ~53% and thus more than 76% of the electrons at the Fermi level are of majority character. In the case of the Mn-
FIG. 7: Layer-resolved DOS at the Fermi level for the Ni-terminated NiMnSb/InP(111) contacts calculated using the FLAPW method. In the middle of the figures a Ni/P interface is shown with Sb (top) or Mn (bottom) in the subinterface layer, while at the borders of the figures the layers of a Ni/In interface can be seen with Mn (top) or Sb (bottom) in the subinterface layer.

In Fig. 7 we also show the two different P/Sb-terminated interfaces: In the top panel the one with Mn as subinterface layer is of particular interest since the Mn atom shows a practically zero net spin-polarization decreasing considerably the overall spin-polarization at the interface. On the other hand, when the subinterface layer is Ni as in the middle of the bottom panel, all atoms at the interface show a very high majority DOS at the Fermi level and the resulting spin-polarization, $P$, is $\sim 74\%$ and thus $\sim 86\%$ of the electrons at the Fermi level are of majority character. We should also mention that, although the induced majority DOS at the Fermi level for the P atom at the interface seems very large (it is of the same order of magnitude with the Ni one), when we move away from the Fermi level it becomes very small compared to the majority DOS of the transition-metal terminated NiMnSb-films (not shown here) the interface states are even stronger than for the Ni-terminated spacers and the spin-polarization at the interface vanishes.

In the last part of our study we will concentrate on the Sb-terminated (111) interfaces. In their paper Wijs and de Groot predicted that the interfaces between the Sb-terminated NiMnSb(111) film and a S-terminated CdS(111) film should keep the half-metallicity or at least show an almost 100% spin-polarization at the Fermi level. Thus it is of particular interest to study the interfaces between the Sb atom and P, although P has one electron less than S. Firstly, we should note that contrary to the Mn and Ni terminated, in the case of the Sb-terminated NiMnSb(111) surfaces the interface state was not pinned exactly at the Fermi level but slightly below it and the spin-polarization in the case of the Sb surfaces was still high. In the case of the interfaces between In and Sb half-metallicity is completely destroyed and the spin-polarization is even negative; there are more minority-spin electrons at the Fermi level than majority ones as can be seen from the DOS at the boundaries of the pictures in Fig. 8.

In Fig. 8 we also show the two different P/Sb-terminated interfaces: In the top panel the one with Mn as subinterface layer is not of particular interest since the Mn atom shows a practically zero net spin-polarization decreasing considerably the overall spin-polarization at the interface. On the other hand, when the subinterface layer is Ni as in the middle of the bottom panel, all atoms at the interface show a very high majority DOS at the Fermi level and the resulting spin-polarization, $P$, is $\sim 74\%$ and thus $\sim 86\%$ of the electrons at the Fermi level are of majority character. We should also mention that, although the induced majority DOS at the Fermi level for the P atom at the interface seems very large (it is of the same order of magnitude with the Ni one), when we move away from the Fermi level it becomes very small compared to the majority DOS of the transition-metal
the second case the Mn is deeper in the interface and as can be seen in Fig. 9, the contrary effect occurs. In occupied minority states would be higher in energy but, case the exchange splitting should be larger and the unoccupied minority states to move slightly lower in energy. Thus, the Fermi level does not fall in the local minimum but shifts into the peak of the unoccupied minority states and the net spin-polarization vanishes. The Ni states are strongly polarized by the Mn ones and also in the case of the Ni atom which is deeper than the Mn one, the Fermi level does not fall anymore within the local minimum.

Similarly to the (001) interfaces in Section III C, we also calculated the band-offset in the case of the (111) interfaces. The band-offset ranges from 0.36 eV in the case of the ...-In-P/Mn-Sb-Ni-... contact up to \( \approx 1 \) eV for the ...-In-P/Sb-Ni-Mn-... configuration. Thus the conclusions of Section III C are valid also for these interfaces.

V. SUMMARY AND CONCLUSIONS

In the first part of our study we investigated the electronic and magnetic properties of the (001) interfaces between the half-metal NiMnSb and the binary semiconductors InP and GaAs using two different full-potential \textit{ab-initio} techniques. Both methods gave similar results in the case of the NiMnSb/InP(001) contacts. In all cases the (001) interfaces lose the half-metallicity but in the case of the Ni/P contact the Ni has a bulk like behavior since the P atoms substitute the cut-off Sb isovalent neighbors and 70\% of the electrons are of majority-spin character at the Fermi level. But in the case of the Ni/As interface the large hybridization at the interface suppresses this high spin-polarization. MnSb-terminated interfaces, on the other hand, present very intense interface states which penetrate also into the deeper layers of the NiMnSb film.

In the second part of our study we investigated all the possible (111) interfaces between NiMnSb and InP. In all cases interfaces states destroy the half-metallicity but in two cases the interface presents high spin-polarization. Firstly, when the contact is the ...-In-P/Ni-Mn-Sb-..., the Ni atom at the interface has a bulklike environment and the spin-polarization at the Fermi level is as high as 53\%. In the case of the ...-In-P/Sb-Ni-Mn-... contact the spin-polarization is even higher reaching a value of 74\%.

Although half-metallicity at the interfaces is in general lost, there are few contacts in which a high spin-polarization remains, that makes them attractive for realistic applications. Interface states are important because the interaction with defects makes them conducting and lowers the efficiency of devices based on spin-injection. Thus, building up interfaces with the highest spin-polarization possible like the ones proposed here is a perquisite but not a guarantee to achieve highly efficient

![Fig. 9: Atom- and spin-resolved DOS for the ...-In-P/Sb-Mn-Ni-... interface (left panels) and the ...-In-P/Sb-Ni-Mn-... interface (right panels) calculated with the FLAPW method. The values in the figures are the spin moments of the atoms at the interface in \( \mu_B \). The thin solid line indicates the bulk results.](image-url)
spin-injection.

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