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METAL-SEMICONDUCTOR INTERFACE FORMATION

D. BOLMONT

Laboratoire de Physique des Solides, associé au CNRS

Université P. et M. Curie, 75230 - Paris Cedex 05, France

The more and more detailed study of an increasing number of metal-semiconductor (MS) interfaces tends to demonstrate the prevailing part played by the different local bonds associated with the compositional change at the interface. In the case of the abrupt interfaces both geometrical properties and an electronic equilibrium are settled at a coverage below one monolayer. A few cases are discussed here which show the variety of physical phenomena involved in the barrier height fixation in contradiction with all attempts to a unified model.

INTRODUCTION

The historical evolution of the MS (metal-semiconductor) interface concept has followed the progress in the diverse experimental methods and theoretical approaches. It appears as the refinement of an initially simple model, associated with a better perception of the different structural parameters. At first the electronic properties of the MS contact have been thought to be governed either by the metal M (Schottky (1)) or the semiconductor S (Bardeen (2)). Then after swinging between the two (3-6), the physical properties of the system appear progressively as governed by the atomic interaction at the interface between the atoms of both metal and semiconductor, the local bonds which are formed being dominant. The multiplicity of the systems studied has led to a gradation of the reactions at the interface in which the limit cases are :

- a cohabitation metal-semiconductor without any visible modification of the substrate properties and with a geometric extension of the transitional zone limited to the first atomic planes in contact.
- a form of intermixture more or less spread along the

growing direction with a transition zone between semiconductor and metal which might, according to the cases, go from a two-dimensional buffer layer, a few $\text{\AA}$ thick, to a three dimensional alloy several hundred $\text{\AA}$ thick.

This first view of the MS interface sets the problem of the adequacy between the system under study and the experimental approach. With regard to our work and discussions in this article we are chiefly interested by interfaces with no or little intermixture in their phase of formation : it makes the use of surface physics methods specially suited to study the early stages of the formation under vacuum of such interfaces. Although the studies on these systems started a few decades ago, a good appreciation of interfacial physics has been reached only within the last few years. Nowadays it looks like a game between the semiconductor and metal characteristics and likewise between the interactions of both. In order to separate more easily the determinant part of each component, the study of the less reactive systems is the best adapted to a comprehensive approach of the MS interfaces. During the interface formation this study is deployed round the three following aspects :

- the evolution of the semiconductor surface intrinsic properties
- the development of the interfacial structures
- the formation of the metal layer.

The success of this approach is subordinated to the possibility to carry out the following operations :

- the preparation of the semiconductor substrate surface, clean, well characterized (reproductiveness) both geometrically (orientation, defects, surface restructuration) and electronically (electronic states). As a matter of fact, for many years, in spite of the lack of many present experimental skills (7) the influence of the surface quality had been already observed. Until now our experimental work has been concerned with the Si (III) face, both 2×1 and 7×7 reconstructed

and GaAs (IIIO) surfaces cleaved under ultra-high vacuum. The experimental methods were : LEED (low energy electron diffraction), AES (Auger electron spectroscopy), ELS (electron loss spectroscopy) and PYS (photoemission yield spectroscopy) - the deposition of an accurately calibrated amount θ of pure metal on the clean surface. The coexistence of preparation and measurements in the same ultra-high vacuum system requires the use of the MBE technic (molecular beam epitaxy) where the effusion cells can provide a deposition rate as low as 10^{-4} ML/s. The smallest coverage able to provoke a measurable modification of the surface is used as a start. If 1 monolayer (ML) is 1 deposited atom per semiconductor surface atom, less than 0.1 ML on Si (III) give observable changes in the interfacial characteristics while 10^{-3} ML already modifies the surface properties of GaAs (IIIO).

The development of the MS interface which comes with the substrate evolution and with the metallic deposition may be easily described through two sets of phenomena well established in surface physics. The first concerns geometrical properties where the following effects can appear

- surface substrate relaxation induced by the deposited atoms
- two-dimensional structuration of the deposition ($\theta < 1$ ML) the ordering of which depends as much on the substrate surface as the metallic species
- epitaxy of the metal on the semiconductor ($\theta > 1$ ML).

The second concerns the change in the electron state densities which may reveal

- the evolution (conservation, displacement or removal) of the electron surface states of the clean substrate surface. This evolution is the double mark of both deposition reactivity and the part played by the semiconductor in the Fermi level pinning and consequently of the Schottky barrier.
- the creation of new states at the interface related to the MS interactions and also to the first MM interactions

when the deposition becomes sufficient or if some metallic aggregates are formed

Where the MS interface is established

The major point which appears in the MS interfaces formed on Si (III) and GaAs (IIIO) by Ag, Cu, Au, Ga, In or Al with or without annealing, is the extreme sensitivity of the electronical properties of the system to the number of deposited atoms. It governs the fixation of the Fermi level position with respect to the bulk bands of the semiconductor (barrier height) at metal coverages lower or much lower than one monolayer for all the studied cases. The MS contact no longer appeared as a metal-semiconductor surface and a two-dimensional layer of foreign atoms which on the whole determines the future contact physics : once this two-dimensional layer completed, further metal deposition do not modify its character. The adsorbed atoms may be considered as any impurities on the surface and in some cases (GaAs) the metal effect in the first deposition stage is similar to the interaction with a gas such as oxygen. Let us note also that this intermediate area between semiconductor and metal is often a two-dimensional structured network where the metal atom arrangement is peculiar to this zone and does not correspond to the one projected from the bulk.

The MS interfacial geometry

θ_{ml}	Ag.Si(111) 2x1	Ag.Si(111) 7x7	Ag.Si(111) 7x7 annealed	Ga.Si(111) 2x1	In.Si(111) 2x1	Al.Si(111) 2x1
0.05						
0.3					$\sqrt{3} \times \sqrt{3} - R30^\circ$	
0.7	$\sqrt{7} \times \sqrt{7} - R19^\circ$			$\sqrt{3} \times \sqrt{3} - R30^\circ$	2x2	$\sqrt{3} \times \sqrt{3} - R30^\circ$
1			$\sqrt{3} \times \sqrt{3} - R30^\circ$			
2	epitaxy 1x1			1x1	epitaxy	
4		epitaxy 1x1				epitaxy

Table I - LEED structures of some M-Si (III) systems.

Certain surfaces such as Si (111) have a geometry which promote the building of structured interfacial planes (table n°I). The variety of the encountered structures in the M-Si (III) compositions is remarkable and points out the natural tendency of the (III) face's three-fold symmetry to accept different parallel structures provoked by the metal arrangement in the surface network. The ad-atom position varies from one metal to another either along the surface plane or within the last atomic layer. For instance let us quote two characteristics examples. First

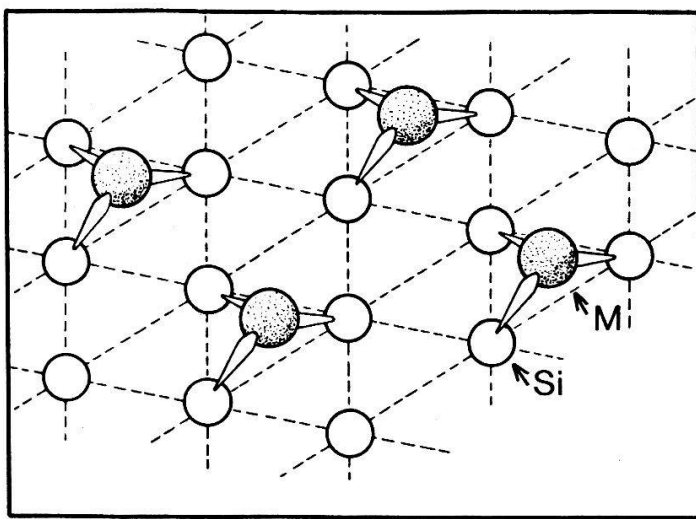


Fig.1 Metal (like Ga, In, Al) bound on Si(111). The dangling bonds are used. The deposition is $\sqrt{3} \times \sqrt{3}$ -R30° structured.

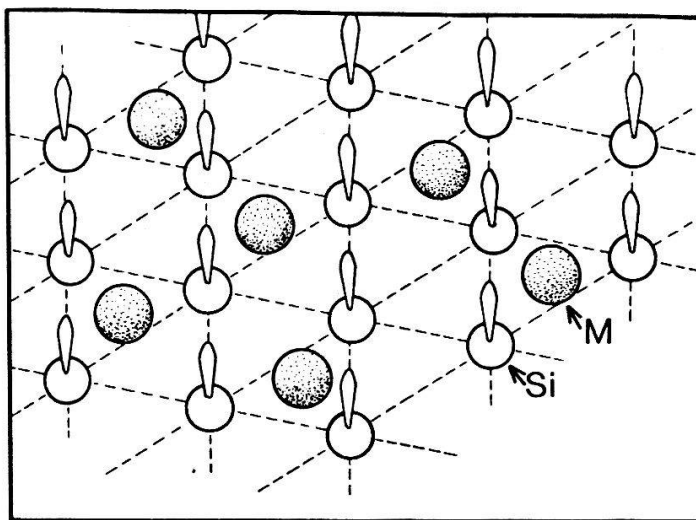


Fig.2 Metal (like Ag) in interstitial position on Si(111). The dangling bounds are conserved.

the metal atom can take and adsoption site governed by the dangling bond geometry of the semiconductor surface (Fig.1). This is the case of Ga, In, Al on Si (111) (8) where $1/3$ ML brings out a $\sqrt{3} \times \sqrt{3}$ -R 30° structure which is interpreted as the binding of the metal atom to three Si surface atoms the dangling bond of which is replaced by a covalent bond. In the second case it may be possible that the metal atom ignores these preferential sites and adsorbs in some interstitial positions (Fig.2) where the nature of the bonds is less pronounced and where a two-dimen-

sional coordination is allowed. This is the case for instance of Ag on Si (III) 2×1 (9) where a new $\sqrt{7}\times\sqrt{7}$ - R 19° structure (Fig.2) has been observed. In these two kinds of adsorption the substrate surface geometry is only very slightly modified as shown by the different LEED analyses. Another class of geometrical interface interaction appears with the (II0) face of GaAs. In this case the additional structures induced by the first metal plane are uncommon, but the electronical structure analyses show, as we shall see later, that the surface atoms of the substrate undergo a restructuration perpendicular to the surface which tends to bring the surface geometry toward its unreconstructed (unrelaxed) structure. The lack of well defined adsorption sites along the (II0) face explains the absence of metal induced superstructures. Let us note however that Ag exhibits a peculiar behavior where an atom alignment occurs at low coverage in the $[\bar{1}\bar{1}0]$ direction, which initiates the metal epitaxy in an unexpected direction of growth (10). The first transition layer being achieved, the other atoms of metal arriving see a two-dimensional system with both M and S atoms. It may be possible that the new arrivals find an adsorption position in this system which keeps the initial coordination but with an increased number of atoms per elementary cell : this is the case of Ga on Si (III). On the contrary the new ad-atoms can break the existing coordination and produce some transitory structures as in In on Si (III). The surface atom mobility along the forming MS system is often sufficient to provoke atom clustering which gives a voluminal metal. Typical examples of epitaxy are found at room temperature with Ag and In on Si (III) 2×1 as well as Ag on GaAs (II0) where the surface substrate cristalline direction induces an epitaxial direction of the growing metallic layer. At coverage higher than 10 ML the layer thickness is often inhomogeneous with islands more or less spread along the surface. The shape of these islands can be changed by thermal processing as in the case of Ag on Si (III) 7×7 (II).

MS interfacial electronic structures

From the clean semiconductor surface with its geometrical defects (steps, etc..) and with its intrinsic electronic state density (valence, dangling bonds, defect of the real surface), the adsorbed metal atom can be thought of as a perturbation center from which the effect on the surface's electronic equilibrium can be interpreted either in term of defect (of which the nature might be specified) or in that of surface reactivity. Altogether microscopic interactions give some structural evolution macroscopically marked by some Fermi level E_F pinning position in the semiconductor gap and perceptible in the Schottky barrier measurement. As a first exemple let us note the Ag-Si

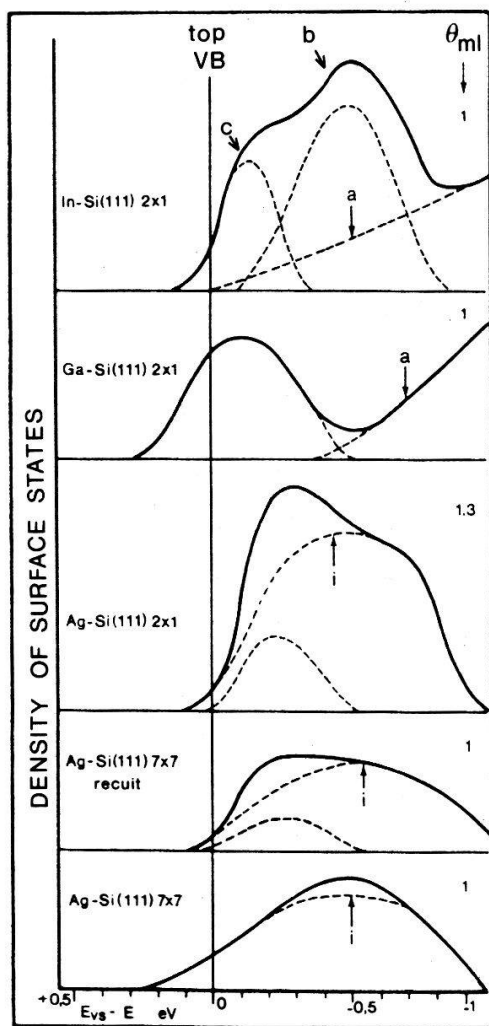


Fig.3 Density of electronic surface states M -Si i -Si(111)2x1-7x7 clean
a,b,c-metal induced.

(III) 2x1 (9) case where the Si dangling bonds (i in Fig.3) appear as maintained, E_F being determined by the substrate's intrinsic surface states, is not modified by the very weak density of states induced by the metal. Only a work function shift towards the metal one, induces an overall shift of the energy levels without any band bending modification. This behavior shows the determinant effect of the semiconductor surface on the Schottky barrier determination. Such a behavior is specific to the Ag-Si (III) 2x1 system but it cannot be generalized to any system on Si. For example on Si (III) 7x7 the effect of Ag is more important and with Ga and In the deposited atoms control the electronic properties of the interface. For the latest systems in accordance

with the adsorption model given above (Fig.1) the dangling bond states of the semiconductor disappear and are rapidly replaced by bonding electron states (a, b, c in Fig.3) near the valence band edge. They spread into the gap and stabilize the Fermi level in a position, which appears almost independent of the initial surface structure and of the metal species, close to $\frac{2}{3} E_g$

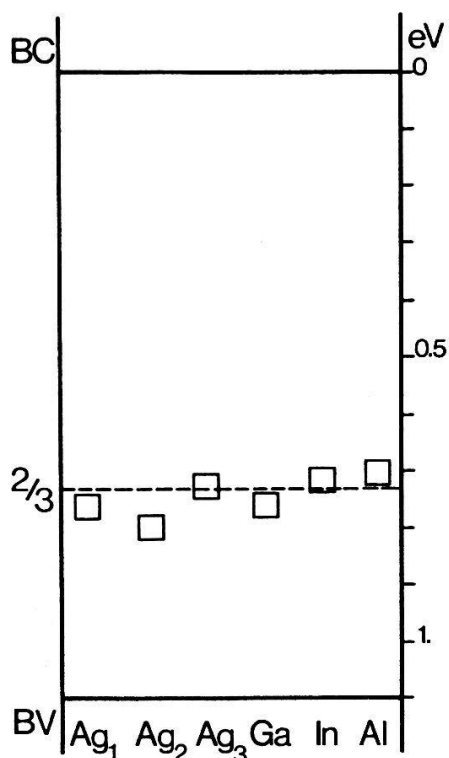


Fig.4 Fermi level M-Si
 $Ag_1 = Ag-Si(111)$, $Ag_2 = Ag-Si(111)7 \times 7$, $Ag_3 = Ag^2-Si(111)$ annealed.

below the conduction band edge (fig.4). The shape of the interface state distribution gives to this two-dimensional region a character which is more covalent than metallic, the MS bonds on the silicon sites showing a strong spatial localization. With such systems the metal plays a determinant role for the electrical characteristics of the MS contact. Another class of interactions at the interface is found with the deposition of some metals (here Ag, Ga, In, Al) on cleaved GaAs (110). The analysis of the density of states near the threshold shows a great sensitivity of the electronical equilibrium to the metal coverage. More precisely a few 10^{-2} ML of metal are

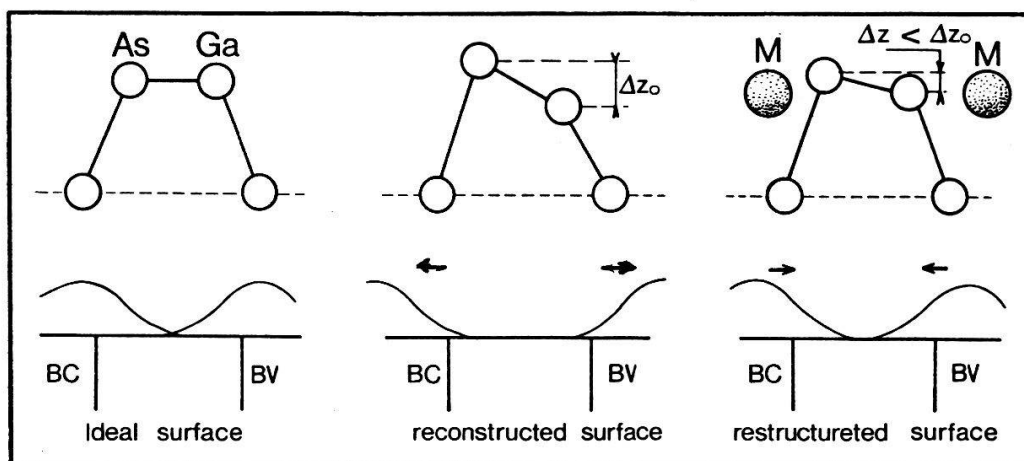


Fig.5 Metal-GaAs(110)restructured surface model

sufficient to stabilize the Fermi level in the gap at a position which is that of the future Schottky barrier, and which is almost independent of the deposited metal. One explanation for such sensitivity is the effect of substrate restructuring induced by the deposited atoms (Fig.5) which as been mentioned

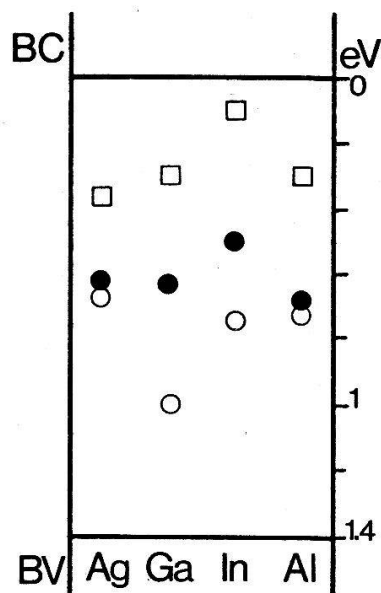


Fig.6 Fermi level M-GaAs (1110) - $\bullet$ n.type, $\circ$ p.type (1ML), $\square$ metal work function (0=4eV under vacuum level)

earlier. A return of As and Ga atoms from the semiconductor toward their ideal extrapolated bulk positions should bring back into the gap the electronic states initially removed by the substrate relaxation after cleavage. These intrinsic states lead to the Fermi level pinning at a position which is a function of the quantity of displaced states, then in relation with the restructuring importance. The other contributions come from the MS interaction states and are proportional to the number of deposited atoms below 1 ML and play a part in the 1 ML coverage range. Fig.6 gives the Fermi level position, in the gap, for the four M-GaAs (1110) interfaces studied in our experiments.

The Al has a different behaviour and seems to be more reactive than the other three metals : the formation of an interfacial compound is possible in this case.

CONCLUSION

The few results and observations which have been given here on MS systems with small intermixing show the double aspect of the interfacial physics which is at the origin of several generations of more or less unified explanatory models. It is the diversity of the physical processes which determines the MS contact characteristics in apparent contrast with certain behaviour constant (such as the barrier height) which would rather favorise a unified model. Nowadays in the present state of our

knowledge, if we want to put the accent on one of the most characteristic features of the MS interfaces, we should hold our attention on the extreme sensitivity of the electronic structure of the local bonds in the interfacial region. As a consequence the future MS contact parameters are defined in the early coverage stage, in the one monolayer range.

- I - W.SCHOTTKY ; Zeits.F.Physik II8,539 (I942)
- 2 - J.BARDEEN ; Phys.Rev. 7I,7I7 (I947)
- 3 - W.G.SPITZER, C.A.MEAD ; J.Appl.Phys. 34,306I (I963)
- 4 - V.HEINE ; Phys.Rev. I38,AI689 (I965)
- 5 - J.C.PHILLIPS ; Phys.Rev. BI,593 (I970)
- 6 - J.C.INKSON ; J.Phys.C : Solid St.Phys. 5,2599 (I972)
- 7 - F.G.ALLEN , A.B.FOWLER ; J.Phys.Chem.Solids 3,I07 (I957)
- 8 - D.BOLMONT ; Thèse d'Etat Paris (I982)
- 9 - D.BOLMONT, P.CHEN, C.SEBENNE, F.PROIX ; Phys.Rev.B24,4552 (I98I)
- IO- D.BOLMONT, P.CHEN, F.PROIX, C.SEBENNE ;
J.Phys.C : Solid State Phys. I5,3639 (I982)
- II- D.BOLMONT, P.CHEN, C.SEBENNE ;
J.Phys.C : Solid State Phys. I4,33I3 (I98I)