One-dimensional half-metallic interfaces of two-dimensional honeycomb insulators

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We study zigzag interfaces between insulating compounds that are isostructural to graphene, specifically II-VI, III-V and IV-IV two-dimensional (2D) honeycomb insulators. We show that these one-dimensional interfaces are polar, with a net density of excess charge that can be simply determined by using the ideal (integer) formal valence charges, regardless of the predominant covalent character of the bonding in these materials. We justify this finding on fundamental physical grounds, by analyzing the topology of the formal polarization lattice in the parent bulk materials. First-principles calculations elucidate an electronic compensation mechanism not dissimilar to oxide interfaces, which is triggered by a Zener-like charge transfer between interfaces of opposite polarity. In particular, we predict the emergence of one dimensional electron and hole gases (1DEG), which in some cases are ferromagnetic half-metallic.

Since the first two-dimensional material (graphene) was successfully synthesized by mechanical exfoliation [1], a growing number of studies has been devoted to other planar systems. These include BN monolayers [2], layered transition metal oxides [3], dichalcogenides [4, 5], or topological insulators such as Bi2Te3 or Bi2Se3 [6]. Of particular note in this context are a number of II-VI, III-V and IV-IV compounds, whose stable bulk phase is wurtzite, but may adopt a planar graphitic phase in ultrathin films. These were first predicted theoretically several years ago [7, 8], and later grown in the lab [9].

Recent improvements in growth techniques, allow joining these nanosheets together. In addition to the many heterostructures studied so far involving graphene and BN, obtained by modified stacking [10] or segregated nanosheets [11, 12], there are and will be coplanar heterostructures [13, 14], made from different insulators on the same sheet many of them having a significant degree of ionic character. This brings up the important question of whether “polar discontinuities” [15] might play a role in these low-dimensional systems. Vertical stacking of nanosheets is not dissimilar to oxide interfaces [16], and polarity effects will resemble those of ultrathin films already discussed in the literature [7, 8]. However, to our knowledge, coplanar interfaces between 2D insulators have not been addressed from first principles, except for a qualitative discussion of the electrostatics [17].

Compared with the case of polar oxide interfaces, there are a few additional challenges that are specific to the 2D case. First, the electrostatics is somewhat more complicated: unlike a plane of charge (interface between 3D materials), which generates a uniform electric field in the whole space (essentially a 1D problem), a line of charge (interface between 2D materials) produces a logarithmically divergent potential with inhomogeneous stray fields in the vacuum region. Second, the symmetry group (with a main six-fold rotation axis but no center of symmetry) gives rise to different polarization classes than the ones found in centrosymmetric insulators [18]. This calls for special care in the application of the modern theory of polarization, and in particular of the interface theorem [19], for the determination of the excess “bound” charge at a given interface. Moreover, the lack of center of symmetry implies that these materials are piezoelectrically active.

Here we perform extensive density-functional theory calculations to show that (i) there indeed exists a net charge at the 1D zigzag interface between two chemically distinct polar insulating honeycomb domains; that (ii) the linear charge density is uniquely determined from bulk properties of the participating materials, and is related to the nontrivial topology of their respective polarization lattice; that (iii) the divergent electrostatic potential produced by the lines of charge triggers a Zener-like breakdown mechanism in the limit of increasingly wide domains; and that (iv) such mechanism gives rise to spin-polarized one-dimensional electron/hole gases (1DEG/1DHG). In what follows, we proceed addressing points (i) to (iv) in the same sequential order.

According to the modern theory of polarization [19], and as further elucidated in Refs. [20,22], the net charge at the interface between two compounds can be exactly given by invoking only bulk properties of the parent materials. In particular, one needs to consider the lattice (Fig. 1) of allowed values for the so-called formal polarization, P, of either crystalline constituent. Then, based on the interface orientation and termination [22], one can readily deduce the interface net charge. An intuitive way to do this is by representing the electronic structure of the
material in terms of localized Wannier functions [23, 24]. For polarization purposes, these effectively map the continuous charge density distribution of the periodic crystal into a discrete set of classical point charges, located at the Wannier centers [39] shown in Fig. 1(a). Then the formal polarization of the crystal is simply given by the total dipole moment of the ensemble of these charges and the ion cores divided by the unit cell volume. Of course, there are infinite possible choices for the primitive basis of atoms and Wannier orbitals, hence the multivaluedness of the formal $P$ [see Fig. 1(b)].

A particularly convenient approach is that of combining the valence Wannier functions with their nearest neighbor ion core, obtaining a negatively charged object that we call “Wannier anion” [22]. This way, we obtain polarization lattices that include only a subset of the original points [larger symbols in Fig. 1(b)]. More specifically, each of these restricted points simply corresponds to the dipole moment of a system of two integer point charges $Q = \pm Ne$, placed at two arbitrarily chosen lattice sites, where $N$ is the “formal valence” of the cation in each compound. Note that, due to the three-fold symmetry of these structures, both the Wannier anion and the cation carry zero dipole moment; hence, the formal polarization only contains the point-charge contribution [22]. This has some analogies with the LaAlO$_3$/SrTiO$_3$ case, where it was shown that covalency effects are also irrelevant to charge-counting purposes [22, 23].

Each arbitrary choice of atomic basis, in turn, yields a well-defined interface termination once the unit cell is periodically repeated to generate a semi-infinite domain. For the zigzag interfaces considered in this work, the appropriate choice of the bulk unit cell (i.e., one of the many choices that tile the whole system without leaving any unpaired ion at the interface) is that shown by shaded regions in Fig. 1(c). The dipole moment along $y$ of such a unit cell is given by $d = -Qa/(2\sqrt{3})$, yielding a formal polarization of $P_y = -Q/3a$. The interface theorem relates differences in formal polarization to the net bound charge at the interface, and we have

$$\sigma_{\text{bound}} = P_y^{(b)} - P_y^{(a)} = -\frac{\Delta Q}{3a}. \quad (1)$$

This implicitly assumes that both parent compounds are hexagonal and have the same equilibrium lattice parameter. In practice there is always a small lattice mismatch, and to realize a pseudomorphic interface one or both constituents need to be strained. Then, Eq. (1) must be corrected by piezoelectric contributions on $P_y$.

The electrostatic potential produced by this interfacial net charge diverges in the limit of increasingly wide domains and needs to be somehow neutralized. To investigate possible compensation mechanism, we use explicit first-principles calculations of 2D superlattices, which are constructed by periodically repeating two compositionally distinct nanoribbons along the stacking direction ($y$), so that the interface direction ($x$) follows the zigzag edge of the hexagonal lattice [see Fig. 1(c)], consistent with the above discussion of the bulk polarity. Having a relatively low ($<7\%$) lattice mismatch, we chose SiC, AlN and ZnO as representative compounds for groups IV-IV, III-V and II-VI respectively. The periodicity of the superlattices is controlled by varying the nanoribbon width, $m$ (given by the number of AB zigzag chains). Only stoichiometric compounds are considered, which means that two chemically distinct interfaces are present in our superlattices: one of type $A_aB_zA_yB_y$ ($A_{a\gamma}$: cation of material $a$, $B_{\gamma}$: anion of material $b$ etc.) and one of type $A_yB_yA_yB_y$. Along the normal direction to the sheet we include a vacuum layer of 20Å, sufficiently thick to obtain converged results (no significant differences are observed with a vacuum layer of up to 200Å). The calculations are performed using the spin-polarized Perdew-Burke-Ernzerhof (PBE) exchange correlation functional [20] as implemented in the SIESTA code [27, 28]. Norm conserving pseudopotentials [29] and a double-$\zeta$ polarized basis set are used to represent the valence electrons [30]. Atomic forces are relaxed to less than 20 meV/Å.

FIG. 1: (Color online) (a) The position of the Wannier centers (small circles) in the primitive unit cell of ZnO, AlN and SiC. The “cation” (Zn, Al, Si) is represented by squares, and the “anion” (O, N, C) by circles. “a” is the cell parameter (3.316 Å, 3.121 Å, 3.110 Å for ZnO, AlN and SiC respectively), and $\lambda$ is the distance of the $sp^3$ Wannier centers from the anion ($\lambda=0.431$ Å, 0.491 Å, 0.677 Å respectively), which increases with the bond covalency. (b) The triangular lattice of allowed polarization values for the II-VI (red circles), III-V (black triangles) and IV-IV (blue squares) 2D honeycomb compounds. Each symbol represents a polarization vector, indicated by $P$. The box highlights the values obtained with “Wannier anions” (see text). (c) Sketch of the zigzag interface between two compounds ($A_aB_z$ in red and $A_yB_y$ in blue).
eventually becomes higher in energy than the conduction band minimum (CBM) at the $n$-type (Si-N) interface, and a charge transfer becomes favorable in energy. The resulting accumulation of free carriers along the interfaces originate the aforementioned 1DEG/1DHG whose spatial localization can be appreciated from their calculated planar-averaged densities, plotted in Fig. 2(b). The 1DEG/1DHG, in turn, partially neutralize the net charge at either interface, reducing the potential offset. This feedback mechanism results in an effective pinning of the $p$-VBM and the $n$-CBM, which remain close in energy regardless of the value of $m$ after breakdown occurs. Thus, many aspects of the present electronic mechanism resemble those already reported for oxide superlattices \[24\]. However, we stress that the electrostatics of this system is different, as evidenced by the macroscopic potential profile, $V(y)$, followed by the DOS, which is analogous to that of an infinite array of alternating positive and negative linear charge densities [sketched in Fig. 2(a)].

At smaller ribbon widths (not shown) the associated potential drop is smaller than the electronic band gap of either participating compound, and the system can remain insulating (i.e. no “Zener breakdown” occurs). In the large $m$ limit, on the other hand, the 1DEG/1DHG densities tend to the “ideal” value, which corresponds to perfectly neutralizing the discontinuity in the normal component of the bulk $P$, according to Eq. 1. This dependence of the charge transfer with ribbon width is shown in Fig. 2(c) for both AlN-SiC (III-V / IV-IV) and ZnO-SiC (II-VI / IV-IV) interfaces. This nicely shows a cross-over scenario from a short-period uncompensated regime, to a longer-period regime where a Zener-like breakdown mechanism occurs, similar to the case of a LaAlO$_3$-SrTiO$_3$ superlattice. The precise details of the cross-over depends on the band gaps and alignments between the two materials, which might not be properly given by DFT calculations, although the physical mechanism remains valid. Note that the asymptotic limits (dashed lines in Fig. 2(c)), in fact, do not correspond exactly to the ideal rational values (1/3 and 2/3) of Eq. 1. This is due to piezoelectric effects, which again are accounted for by macroscopic bulk properties of the participating materials. In particular, the lattice mismatch between SiC and ZnO, forces a compression in the ZnO ribbon (and a minor expansion in SiC) resulting in a shift of its formal polarization lattice along the axis normal to the interface, hence decreasing $P_{SiC} - P_{ZnO}$.

We find that the 1D gases of carriers at the interfaces of these insulating materials are fully spin-polarized. To further investigate magnetic effects, we plot in Fig. 3 the band structure near the Fermi level, $E_f$, for a AlN-SiC superlattice with thickness that corresponds to the uncompensated regime (a), and a partially compensated superlattice with larger thickness (panels (b)-(d) in the figure). There are three dispersive $r$ bands close to $E_f$ that are involved in the compensation mechanism, one occu-

![FIG. 2: (Color online) (a) Spin-resolved layer-by-layer density of states of the AlN-SiC ($m=16$) superlattice. The top half corresponds to the AlN slab, and the bottom half to the SiC slab, with the interface terminations indicated. Dashed (red) line schematizes the electrostatic potential, $V(y)$, for an infinite array of alternating +/- cylindrical charges. (b) The spatial (plane averaged) charge distribution computed from the local density of states (LDOS) is shown in the lower panel (shaded curves are macroscopically averaged LDOS). (c) The density of free electrons (or holes) calculated as a function of slab width ($m$) for the two superlattice systems studied (AlN-SiC and ZnO-SiC). Theoretical values corrected by piezoelectric effects are marked by dashed lines.](image-url)
plied and mostly localized at the Al-C interface, and two empty and mostly localized at the Si-N interface. The flat bands at X for the non-magnetic solutions Fig. 3(a) and (b) very much resemble the edge states in graphene nanoribbons described with a simple tight-binding model for the \( \pi \) electrons. The corresponding high-density of electronic states at \( E_f \) suggests magnetic ordering due to a Stoner instability.

Indeed, two spin-polarized phases with ferromagnetic orderings along the interface are lower in energy than the paramagnetic solution, and correspond to the transfer of charge (electrons) with well defined spin from one interface (\( p \)-VBM) to the other (\( n \)-CBM); when the spin is preserved in the transfer, as in Fig. 3(c), the electronic magnetization at both interfaces is antiferromagnetically aligned (AF), whereas a spin-flip of the transferred charge gives the ferromagnetic (FM) solution (panel d). The two solutions, AF and FM between interfaces, are almost degenerate in energy, because the 1DEG and the 1DHG are decoupled by the wide insulating ribbon. Notice that the whole system is half-metallic for the AF solution, with full spin polarization of the conducting electrons.

It is generally agreed that a one-dimensional metal is unstable with respect to a symmetry lowering modulation of the charge. The coupling between lattice vibrations and electrons near the Fermi level drives Peierls distortions that open a gap. Such deformation could also be a way to resolve the instability due to the large density of states at the Fermi level shown in Fig. 3(b). To address the stability of the 1DEG against Peierls distortions, calculations using zone-folding techniques were performed for the AlN-SiC \((m=16)\) superlattice, including up to three or four unit cells along the \( x \) axis (in the limit of large \( m \) a supercell with three unit cells should be enough to accommodate symmetry-lowering deformations capable of opening a gap in the right region of the Brillouin zone). We did not detect any trace of bond-length alternations in our tests, all of which remained in a metallic state. This is in agreement with previous studies that showed that Peierls distortions can open a gap in similar edge bands only for narrow (<6Å) graphene and graphene nanoribbons.

In addition to competing structural/electronic ground states, another factor that could thwart experimental realization of the 1DEG/1DHG predicted here are defects. In particular, these polar structures may, in certain growth conditions, find more energetically favorable compensation mechanisms than electronic transfer. For example an anionic vacancy could donate electrons to the \( n \)-type interface, or combinations of anionic vacancies and cationic vacancies at each interface could provide the necessary charge compensation without free electrons. Similar mechanisms have been proposed at the \( \text{LaAlO}_3/\text{SrTiO}_3 \) interface and in ferroelectric thin films. Substitutional defects and charge transfers between edge states in isostructural C/BN superlattices have been also discussed. If the growth conditions can be controlled such that these point defects are minimized it may be possible to utilize the systems here for various applications such as spintronics, sensors and electronics.

In conclusion, we have shown that polar compensation mechanisms can give rise to fully spin-polarized one-dimensional electron and hole gases at the interface between two coplanar insulating bidimensional materials, in close analogy with the 3D case of \( \text{LaAlO}_3/\text{SrTiO}_3 \) (001) interface. The net charge at the interface can be determined solely from bulk properties (the polarization) of the parents materials, as required by the interface theorem. The polarization is given by a simple charge counting procedure, irrespective of covalency/ionicity, that results in three distinct sets of allowed polarization lattices, for the II-VI, III-V and IV-IV compounds, and to the formation of new electronic states (1DEGs) at the interface between two materials of different classes.

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**FIG. 3:** (Color online) The band structure of the AlN-SiC superlattices for \( m = 8 \) (a), and \( m = 16 \) with paramagnetic-NM (b), antiferromagnetic-AF (c) and ferromagnetic-FM (d) solutions. Solid (black) and dashed (red) lines correspond to bands with different spin components. The inset in (b) shows the Fermi level crossing of bands at the zone boundary.


[39] The Wannier calculation was performed with the Wannier90 [38] code and the density functional theory (DFT) code SIESTA [27] [28].