An approximation to density functional theory for an accurate calculation of band-gaps of semiconductors

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Abstract

The local-density approximation (LDA), together with the half-occupation (transition state) is notoriously successful in the calculation of atomic ionization potentials. When it comes to extended systems, such as a semiconductor infinite system, it has been very difficult to find a way to half- ionize because the hole tends to be infinitely extended (a Bloch wave). The answer to this problem lies in the LDA formalism itself. One proves that the half-occupation is equivalent to introducing the hole self-energy (electrostatic and exchange-correlation) into the Schröedinger equation. The argument then becomes simple: the eigenvalue minus the self-energy has to be minimized because the atom has a minimal energy. Then one simply proves that the hole is localized, not infinitely extended, because it must have maximal self-energy. Then one also arrives at an equation similar to the SIC equation, but corrected for the removal of just 1/2 electron. Applied to the calculation of band gaps and effective masses, we use the self-energy calculated in atoms and attain a precision similar to that of GW, but with the great advantage that it requires no more computational effort than standard LDA.

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I. INTRODUCTION

The well known *Density-Functional Theory* (DFT) is an approach to the theory of electronic structure in which the electron density distribution, rather than the many electron wavefunction, plays a central role. The practical applications of DFT are based on approximations for the so-called exchange-correlation potential which describes the effects of the Pauli principle on the many-electron system. If we had the exact exchange-correlation potential we could solve the many-body problem exactly for the ground state. Although the potential is unknown, approximations are made. The most common is the so-called *Local-Density Approximation* (LDA) which locally uses the exchange-correlation energy density of a homogeneous system. The LDA to the Kohn and Sham DFT is still one of the most reliable methods for condensed matter calculations, having successfully predicted and explained a wide range of ground state properties in solid state physics and chemistry. Lately, but very slowly, it is being progressively abandoned in favor of the many GGA (Generalized Gradient Approximations). However, while LDA and GGA have predicted many ground state properties with good accuracy, the electronic properties such as band gaps are significantly smaller than experiment. These discrepancies are caused by the lack of the discontinuity of the exchange-correlation potential in going from the valence to the conduction band. Several methods for overcoming these limitations have been proposed. One of them is the GW approximation, in which one considers the energies of quasiparticles and calculate the electron self-energy in terms of perturbation theory. This procedure has been quite successful, achieving good accuracy, but it goes beyond the DFT. Other procedures were also proposed, among them we can mainly cite the Self Interaction Correction (SIC), the atomic SIC applied to solids (ASIC), screened exchange (SX-LDA), the so-called exact-exchange approach, the well known LDA+U, the work of Liberman, and others. Most of these approaches are computationally very demanding, which prohibits their application to large systems of atoms.

Slater half-occupation scheme was very successful for valence states. One example is that one could obtain energies which were comparable to the experimental ionization energies, though, at that time, good spin-polarized exchange-correlation approximations, as those based on Ceperley and Alder, did not exist. In order to illustrate to the reader
the quality of the results that can be obtained, we present in Table II the first and second ionization potentials of 12 atoms, measured and calculated with LDA with 1/2 occupation.

TABLE I: First and second Ionization Potentials (IP) for some atoms (eV). These results were obtained with spin-polarization but assuming spherical charge densities for ions and atoms. We used a code originally written by S. Froyen, modified by N. Troullier and J. L. Martins, and modified and maintained by A. Garcia.

| Atom | First IP   | Second IP   |
|------|------------|-------------|
|      | Calculation| Experiment  | Calculation| Experiment  |
| C    | 11.60      | 11.26       | 24.58      | 24.38       |
| N    | 14.81      | 14.53       | 30.01      | 29.60       |
| O    | 13.89      | 13.62       | 35.38      | 35.12       |
| Al   | 5.94       | 5.99        | 18.97      | 18.83       |
| Si   | 8.19       | 8.15        | 16.30      | 16.35       |
| P    | 10.44      | 10.49       | 19.80      | 19.73       |
| S    | 10.57      | 10.36       | 23.25      | 23.33       |
| Zn   | 9.70       | 9.39        | 18.65      | 17.96       |
| Ga   | 6.00       | 6.00        | 20.83      | 20.51       |
| Ge   | 7.99       | 7.90        | 15.88      | 15.93       |
| As   | 9.90       | 9.81        | 18.63      | 18.63       |
| In   | 5.73       | 5.78        | 18.56      | 18.97       |

Though the precision of the calculated results shown in the Table is much better than the precision one reaches in the calculation of band gaps, either by LDA or GGA, it has been difficult to find a way to make the ionization of 1/2 electron in extended systems as crystals. Of course the problem is that a crystal is described by means of Bloch waves and removing the population of just one Bloch state is of no consequence. In this paper we present a solution to this problem. We are specially concerned with the calculation of band gaps in semiconductors, for which we obtain calculated results that compare very favorably with experiment, and are not computationally demanding. We report the results for fourteen semiconductors, including the group II-VI, III-V and IV. Our method is inspired in the LDA and in the half ionization but, at some point, it has to be postulated. The quality of
II. LDA AND HALF-IONIZATION IN SOLIDS

Accepting the LDA as a valid approximation to the density functional (DF), the Janak theorem follows
\[
\frac{\partial E}{\partial f_\alpha} = e_\alpha(f_\alpha) \tag{1}
\]
where \(E\) is the total energy of the system, function of the occupation \(f_\alpha\) of the one-particle Kohn and Sham state \(\alpha\). It is a well known fact that the eigenvalue \(e_\alpha(f_\alpha)\) is almost precisely linear with the occupation \(f_\alpha\). \(^{17}\)

Then integrating
\[
\int_{-1}^{0} df_\alpha
\]
between the ground state \((f_\alpha = 0)\) and the ion \((f_\alpha = -1)\) obtain
\[
E(0) - E(-1) = e_\alpha(-1/2) = - \text{ Ionization Potential}. \tag{2}
\]

Taking another derivative
\[
\frac{\partial e_\alpha}{\partial f_\alpha} = 2S_\alpha, \tag{3}
\]
where
\[
S_\alpha = \int \int d^3r d^3r' \frac{n_\alpha(\vec{r}) n_\alpha(\vec{r}')}{|\vec{r} - \vec{r}'|} + \frac{1}{2} \int \int d^3r d^3r' n_\alpha(\vec{r}) \frac{\delta^2 E_{xc}}{\delta n(\vec{r}) \delta n(\vec{r}')} n_\alpha(\vec{r}') \tag{4}
\]

is named “self-energy”, because of the first term in the right. In LDA the functional derivatives become common derivatives times delta-functions. We maintain the functional derivative notation because the final formulae can have extended use. Because of the linearity of \(e_\alpha(f_\alpha)\) \(^{17}\) we may write
\[
e_\alpha(-1/2) = e_\alpha(0) - S_\alpha \tag{5}
\]
and
\[
E(0) = E(-1) + e_\alpha(0) - S_\alpha \tag{6}
\]

Eq. (6) is quite surprising in its simplicity. The Eq. is telling us that, to restore the ground state, with total energy \(E(0)\), from an ion with a hole at state \(\alpha\) we add an electron whose
energy is the eigenvalue $e_\alpha(0)$ minus the hole self-energy. The self-energy is large when the function is much localized as an atomic wavefunction, and is small and zero when it is much spread as a Bloch function. Since the energy of the restored ground state must be a minimum the hole self-energy must be a maximum. Thus the hole should be representable by a very localized wavefunction. \textit{So far, this is the first time we see a demonstration of the hole localization}, though this proof is based on an approximation (LDA) to the DF theory and on the linearity assumption. Of course we cannot say that the localized hole state is truly stationary, specially if its energy is inside the band continuum of the Bloch states, into which the localized hole would be scattered.

The self-energy may be thought as the Quantum Mechanical average of a \textit{“self-energy potential”} $V_s(\vec{r})$ such that

$$S_\alpha = \int d^3 r n_\alpha(\vec{r}) V_s(\vec{r})$$

where $n_\alpha = \psi_\alpha^* \psi_\alpha$.

$$V_s(\vec{r}) = \int d^3 r' \frac{n_\alpha(\vec{r}')}{|\vec{r} - \vec{r}'|} + \frac{1}{2} \int d^3 r' \frac{\delta^2 E_{xc}}{\delta n(\vec{r}) \delta n(\vec{r}')^2} n_\alpha(\vec{r}')$$

$$+ \int d^3 r' \frac{\sum_\beta f_\beta \frac{\partial n_\beta}{\partial f_\alpha}}{|\vec{r} - \vec{r}'|} + \frac{1}{2} \int d^3 r' \frac{\delta^2 E_{xc}}{\delta n(\vec{r}) \delta n(\vec{r}')^2} \sum_\beta f_\beta \frac{\partial n_\beta(\vec{r}')}{\partial f_\alpha}$$

that depends on the state $\alpha$. From now on, in Eqs. such as (8) we will not write the last two terms, those depending on the derivative of the wavefunctions with respect to the occupation $f_\alpha$.

To derive Eq. (6) we assumed linearity, aside from the Janak theorem. The linearity results when the Kohn and Sham eigenfunctions of the ground state are equal to those of the ion, which is correct to a large extent.\footnote{Coherently we may neglect the last two terms in Eqs. such as (8) and make the difference $E(0) - E(-1)$, of two minima, an extremum. Then we minimize (extremize) $e_\alpha(0) - S_\alpha$ as suggested by Eq. (6). To do so we must write a variational expression that, upon minimization (extremization), leads to a differential Eq. for the hole wavefunction $\psi_\alpha(\vec{r})$. \textit{At this point we must set clearly what we imply by the term “hole”. From Eq. (6) we see that we are adding an electron to a hole state of the ion. The hole state might be in a valence or conduction band, the only requirement being that the state is empty in the ion. Thus by a “hole” we mean “an electron filling an empty state” or “a particle excitation”. This particle excitation may be in the valence or in the conduction band.}}
The variational expression must be such that upon calculating it with the solution \( \psi_\alpha(\vec{r}) \) of the differential Eq. it returns back the average \( e_\alpha(0) - S_\alpha \). Then we write it as

\[
e_\alpha - S_\alpha = \left\langle \psi_\alpha \left| \nabla^2 - 2 \sum_I \frac{Z_I}{|\vec{r} - \vec{r}_I|} + 2 \int \frac{n_0(\vec{r}')}{|\vec{r} - \vec{r}'|} d^3 r' + \frac{\delta E_{xc}}{\delta n_0(\vec{r})} - V_S \right| \psi_\alpha \right\rangle
\]  

(9)

where \( n_0(\vec{r}) \) does not include the hole wavefunction \( \psi_\alpha \) and it is the number-density of the ground state. In this case, the wavefunction \( \psi_\alpha \) is to be interpreted as the hole state to be filled with an electron to restore the ground state.

Performing the extremization and then inserting Eq. 8 we obtain the equation below with the top entries of the brackets \{\}

\[
\left[ -\nabla^2 - 2 \sum_I \frac{Z_I}{|\vec{r} - \vec{r}_I|} + 2 \int \frac{n_0(\vec{r}')}{|\vec{r} - \vec{r}'|} d^3 r' + \frac{\delta E_{xc}}{\delta n_0(\vec{r})} \right]
\left\{ \frac{1}{2} \int \frac{\psi_\alpha(\vec{r}')^* \psi_\alpha(\vec{r}')}{|\vec{r} - \vec{r}'|} d^3 r' \right\}
\]

(10)

If we insert Eq. 8 before extremization we obtain the Eq. 10 with the entries at the bottom of \{\}. In this latter case, the next to last term of the operator is exactly the term in the SIC equation. Its effect is to exclude the electron being considered from the Hartree interaction. The last term in the operator, the exchange-correlation term, is very different from the corresponding SIC term, since it depends on the whole density of the system and not only on the density of the \( \alpha \) state. The SIC equation, Eq. 10 with the bottom entries in \{\} is not what we want because the eigenvalue \( \lambda_\alpha = e_\alpha - 2S_\alpha \) and not \( \lambda_\alpha = e_\alpha - S_\alpha \) as the half-ionization requires. It is worth mentioning that, in the calculation of band gaps, SIC over corrects and halving it seems to be a better procedure.

Except for atoms, solving Eq. 10 is very difficult, either with the top or bottom entries in \{\}. One important problem is that the solutions of Eq. 10 are not orthogonal. The SIC solution for atoms is used in the ASIC method, which is excellently reviewed in ref. 9. In our case we proceed differently. We introduce a parametrized self-energy potential and use a variational expression that is an extremum for variations in the parameter(s). The first question to be answered is whether it is possible to define a unique self-energy potential that is state-independent. In Table II we show a study of how the atomic self-energy of many states vary with the assumed self-energy potential. One sees that for \( s \) and \( p \) orbitals, the self-energy does not vary much whether it is calculated with \( s, p \), and even \( d \) self-energy potentials. Of course the self-energy potential that we will use is the one corresponding to
the atomic orbital dominating the crystal energy bands around the gap. In extreme cases we can define a self-energy potential that is angular momentum dependent.

\[ V_S = \sum_l V_{S,l}(r) \sum_{m=-l}^l |l, m><l, m| \] (11)

This possibility was explored in the case of diamond, as shown in the discussion of our results.

TABLE II: Atomic self-energies (eV), Eq. 7, for some third-row atoms and different valence electrons, calculated with self-energy potentials (Eq. 18) derived from half-ionization of different states. The fact that the self-energy of a given state has no important dependence on the self-energy potential, that is the entries at each column for a given atom do not differ much, shows that, for practical purposes, one can neglect the state-dependence of the self-energy potential. In the table below, only the self-energies of d-states seem to be much dependent on the way the self-energy potential is derived.

|     | half-ionized valence state | Self-energy (eV) |   |   |   |
|-----|---------------------------|-----------------|---|---|---|
|     |                           | s               | p | d |
| Zn  | s                         | 3.63            | 4.62 |
|     | d                         | 4.62            | 7.43 |
| Ga  | s                         | 3.15            | 3.67 | 4.67 |
|     | p                         | 3.44            | 3.02 | 3.91 |
|     | d                         | 4.75            | 4.02 | 8.12 |
| Ge  | s                         | 4.13            | 3.71 | 5.03 |
|     | p                         | 3.70            | 3.35 | 4.22 |
|     | d                         | 5.11            | 4.30 | 9.00 |
| As  | s                         | 4.11            | 3.73 | 5.26 |
|     | p                         | 4.26            | 3.93 | 4.73 |
|     | d                         | 5.33            | 4.57 | 9.79 |

Thus, assuming a self-energy potential that is state-independent, we use the following variational expression.\textsuperscript{20,21}

\[ E[n, v, \rho] = K[n] - \int V[p]\rho + \frac{1}{2} \int V[\rho]\rho - \int V_S\rho \]
\[ + \int v(n - \rho) + E_{xc}[\rho] \] (12)

where \( p \) is the proton number-density, \( n = \sum_\beta f_\beta \psi_\beta \psi_\beta^* \) is the electron number-density made out of the squares of the wavefunctions, \( f_\beta \) being the occupation numbers, \( v \) is the Kohn-Sham potential, \( \rho \) is the model number-density, \( V_S \) is the given parametrized self-energy potential, \( K \) and \( E_{xc} \) have their usual meaning of kinetic and exchange-correlation of the Kohn-Sham DFT. The functional \( V[\rho] \) is defined as

\[ V[\rho(\vec{r})] = 2 \int \frac{\rho(\vec{r}')}{|\vec{r} - \vec{r}'|} d^3r' \] (13)

The functional \( E \) is an extremum for variations in any of the three functions \( n, v, \) and \( \rho \):

1. \( \delta E/\delta v = 0 \) leads to \( \rho = n \)

2. \( \delta E/\delta \rho = 0 \) leads to

\[ v = -V[p] + V[\rho] - V_S + \frac{\delta E_{xc}/\delta \rho}{\delta \rho \int \rho \delta E_{xc}/\delta \rho} \] (14)

3. \( \delta E/\delta n = 0 \) leads to Schroedinger Equations with potential \( v \) and eigenvalues \( e_\alpha \), rewritten as.

\[
\left[ -\nabla^2 - 2 \sum_I \frac{Z_I}{|\vec{r} - \vec{r}_I|} + 2 \int \frac{\rho(\vec{r}')}{|\vec{r} - \vec{r}'|} d^3r' + \frac{\delta E_{xc}/\delta \rho(\vec{r})}{\delta \rho \int \rho \delta E_{xc}/\delta \rho} - V_S(\vec{r}) \right] \psi_\alpha(\vec{r}) = e_\alpha \psi_\alpha(\vec{r})
\] (15)

Using Eq. 14 to determine \( \rho \) for given \( v \) and \( V_S \) and solving the Schroedinger equations, find

\[ E = \sum_\beta f_\beta e_\beta - \frac{1}{2} \int V[\rho] \rho + E_{xc}[\rho] - \int \rho \frac{\delta E_{xc}/\delta \rho}{\delta \rho \int \rho \delta E_{xc}/\delta \rho} \] (16)

It must be understood that both \( n \) and \( \rho \) are number densities of \( N \) electrons, not \( N - 1/2 \), but the eigenvalues correspond to a situation where 1/2 electron is removed, if \( V_S \) is well chosen.

We want to find band gaps by taking the difference of total energies due to different occupations \( f_\alpha \). Maintaining the Kohn and Sham potential \( v \) and the model number-density \( \rho \), solution of Eq. 14, the band gap becomes a difference between eigenvalues \( e_\alpha \). Now, because the total energy is a variational functional, that is an an extremum for variations in \( v \), resulting from variations of the self-energy potential \( V_S \), one should look for extreme eigenvalue differences.

\[ \frac{\delta (e_\alpha - e_\beta)}{\delta V_S} = 0 \] (17)
III. THE LDA-1/2 METHOD

Consider the case of an atom. We first prove that the self-energy potential is given by

$$V_S \simeq -V(-1/2, r) + V(0, r)$$

(18)

namely the difference between the all-electron potentials of the atom and of the half-ion.

We begin by writing the potential difference as

$$V(-1/2, r) - V(0, r) = \int_{0}^{-1/2} df_{\alpha} \frac{\partial}{\partial f_{\alpha}} \left\{ -2Z \frac{1}{r} + 2 \int d^{3}r' \frac{n\left(\vec{r}'\right)}{\left|\vec{r} - \vec{r}'\right|} + \frac{\delta E_{xc}}{\delta n(\vec{r})} \right\}$$

$$= \int_{0}^{-1/2} df_{\alpha} \left\{ 2 \int d^{3}r' \frac{n_{\alpha}(\vec{r}')}{\left|\vec{r} - \vec{r}'\right|} + \int d^{3}r' \frac{\delta^{2}E_{xc}}{\delta n(\vec{r})\delta n(\vec{r}')} n_{\alpha}(\vec{r}') \right\}$$

$$+ \int_{0}^{-1/2} df_{\alpha} \left\{ \frac{1}{2} \int d^{3}r' \sum_{\beta} \frac{f_{\beta} \frac{\partial n_{\beta}(\vec{r}')}{\partial f_{\alpha}}}{\left|\vec{r} - \vec{r}'\right|} + \frac{1}{2} \int d^{3}r' \frac{\delta^{2}E_{xc}}{\delta n(\vec{r})\delta n(\vec{r}')} \sum_{\beta} f_{\beta} \frac{\partial n_{\beta}(\vec{r}')}{\partial f_{\alpha}} \right\}$$

(19)

or, for a certain value of $f_{\alpha}$ in $[-1/2, 0]$

$$-V(-1/2, r) + V(0, r) = \int d^{3}r' \frac{n_{\alpha}(\vec{r}')}{\left|\vec{r} - \vec{r}'\right|} + \frac{1}{2} \int d^{3}r' \frac{\delta^{2}E_{xc}}{\delta n(\vec{r})\delta n(\vec{r}')} n_{\alpha}(\vec{r}')$$

$$+ \int d^{3}r' \frac{\sum_{\beta} f_{\beta} \frac{\partial n_{\beta}(\vec{r}')}{\partial f_{\alpha}}}{\left|\vec{r} - \vec{r}'\right|} + \frac{\delta^{2}E_{xc}}{\delta n(\vec{r})\delta n(\vec{r}')} \sum_{\beta} f_{\beta} \frac{\partial n_{\beta}(\vec{r}')}{\partial f_{\alpha}}$$

(20)

Fig. 1 depicts $r$-times the self-energy potential for the nitrogen atom, a typical case, for degrees of ionization $I$ ranging from 0.5 to $-0.2$. Observe that the ratio

$$\frac{V(f_{\alpha}, r) - V(0, r)}{f_{\alpha}}$$

(21)

has a very poor dependence on $f_{\alpha}$ meaning that in Eq. 20 we can take the $xc$ functionals at the full occupation $f_{\alpha} = 0$. Then, comparing Eqs. 8 and 20 our proof is completed.

We will leave to another paper a discussion on the lone hole solution we can get out of Eq. 10. Here we are interested in calculating band gaps of semiconductors. For that purpose we will repeat the atomic self-energy potential (Eq. 20) in the whole lattice and calculate eigenvalues for “hole bands”. But observe that the first term on the right of Eq. 20 when repeated in the whole crystal, diverges because it is Coulomb-like. On the other hand, as Fig. 1 shows, the Coulomb tail of the atomic $V_S$ has no importance because the wave function never goes far. Then, in using the self-energy potential $V_S$ defined in the atoms we first trim the potential with a function as

$$\Theta(r) = \begin{cases} 
1 - \left(\frac{r}{\text{CUT}}\right)^{3} & r \leq \text{CUT} \\
0 & r > \text{CUT} 
\end{cases}$$

(22)
FIG. 1: (Color online) Self-energy potential \( rV_S \) calculated for the N atom at different ionizations \( I \) ranging from 0.5 to \(-0.2\). The lines bunch around that of \( I = 0.5 \), which is made thicker. The wavefunction for the ionized state \( (2p) \) is also shown. Also shown is the potential after the cut-off by \( \Theta(r) \).

The idea is that the atomic self-energy potential is only meaningful where the atomic wave function is not negligible. Of course the trimmed self-energy potential is being repeated throughout the infinite crystal, so that we are actually calculating “filled hole bands”. Due to the trimming, the Coulomb tail (of \(-1/2\) electrons) of the atomic \( V_S \) does not penetrate into the neighboring atoms. With the trimming, the eigenvalues \( e_\alpha \) in Eq. 15 become dependent on the trimming parameter \( CUT \). However, Eq. 17 sets a recipe to choose the value of \( CUT \): one should make the energy gaps extreme.

The function of Eq. 22 has some important properties: 1 - its derivative is also zero at \( r = CUT \), so that its electric field is zero at that point and the cut-off does not add to the total charge of the atom; 2 - the trimmed self-energy potential \( \Theta(r)V_S(r) \) is wholly contained inside a sphere of radius \( CUT \), which facilitates its use in band-calculation methods such as
SIESTA and APW-like. The power \( n \) should be large so that the cut-off is sharp. In actual practice we tried \( n = 8 \) and \( n = 50 \) with similarly good results, thus we adopted \( n = 8 \), which is less abrupt, and does not introduce numerical problems into the programs. Fig. 2 shows a typical behavior of a band gap as function of the parameters defining the cut-off function. The first increase of the band gap with \( CUT \) only means that we are getting more of the valence band self-energy as the cut-off is made at larger radii. In principle the larger the \( CUT \) the more we get of the valence self-energy. After reaching a peak, the gap decreases because (i) the potential \( V_s \) is penetrating into neighboring atoms, tending to a uniform negative potential everywhere in space, shifting downwards all bands, valence and conduction alike, and (ii) the self-energy potential perturbation, being broad, diffuses the excitation wavefunction thus making it to loose locality and self-energy. In other words, the cut-off function should be broad enough so as to include most of the excitation wavefunction and thin enough so as not spread it. Thus, the procedure to determine \( CUT \) is based in Fig. 2 namely we look for the extreme band gap according to Eq. 17.

For a given atom and bonding type, the value of \( CUT \) depends little on the chemical environment. Because we are using \( CUT \) values that make the gaps extreme, small deviations from the optimal values produce only second-order deviations in the gaps. In the Fig. 3 we present the values of anion p state \( CUT \) optimized for arsenides, phosphides and nitrides. The anion \( CUT \) value has a small dependence on the chemical environment, which is approximately linear with the compound bond length. However, the relative variation produced in the energy gap is very small. This can be easily verified in the Fig. 2. Around the energy gap maximum, the range of \( CUT \) found in the optimization for all nitrides leads to a change of only 0.05 eV in the energy gap value. This behavior was also verified for all other calculated compounds. Therefore, we conclude that is very reasonable to consider the same \( CUT \) value for the anion potentials. In the Table 111 we show the optimal values of the parameter \( CUT \) of the trimming function \( \Theta \) of Eq. 22. The values for \( CUT \) in the Table reminds a table of ionic, covalent, or atomic radii, but are not equal to either.

**IV. RESULTS AND DISCUSSION**

We calculated, within the LDA-1/2 approach, the electronic structure for several semiconductors. By comparing the LDA and LDA-1/2 calculation procedures, the LDA-1/2
FIG. 2: (Color online) Band gap of GaN as function of the parameter $CUT$ of the cut-off function $\Theta(r)$ applied to the self-energy potential of $N\,2p$. The band gap extreme values depend to some extent on the exponent $n$ being used ($n=8$ and 50 are shown).

calculations lead to no more computational effort than standard LDA, because the values of $CUT$ depend only on the atoms, not on their environment, and are calculated just once. This is a great advantage of our method. Most of the calculations were made with the code VASP “Vienna Ab-initio Simulation Package” using the ultra-soft pseudopotential$^{22,23}$ to which we added the trimmed self-energy potential. In some instances we repeated the calculations with the SIESTA code$^{24}$ and the results differed by no more than 0.1eV. The two codes are so different, for they use different basis functions and pseudopotentials, that the agreement of their results runs in favor of the reliability of our LDA-1/2 procedure. The $k$-space integrals were approximated by sums over a $9\times9\times9$ special-point of the Monkhorst-Pack type within the irreducible part of the Brillouin zone.$^{25}$ The number of plane waves for the expansion of wave functions was optimized for each system and it is basically the same value obtained for optimization of the equivalent standard LDA calculation. The lattice parameters used were the experimental ones.
The present LDA-1/2 proposal is assuming that in promoting an electron from the valence band to the conduction band, the hole thus created is similar to the hole created in the atomic photo-ionization. In other words, the hole has the extent of an atomic hole. If the hole in the extended system overlapped \( N \) equal atoms its self-energy would be \( 1/N^2 \) that of the atomic hole, and the self-energy potential \( V_s \) would be \( 1/N \) that of the atom, or it would have to be calculated with an ion with \( 1/2N \) electrons removed. The results for the semiconductors III-V and II-VI, to be presented shortly, definitely point to \( N = 1 \), meaning that the hole in the solid resembles much the hole in the atom. In fact, the valence band of these semiconductors is known to be made of the anion wavefunctions. On the other hand, for the IV elements Ge and Si (and also for diamond), the results point to \( N = 2 \) meaning that the hole covers the two atoms with covalent bond.

**TABLE III:** Values of \( CUT \) that make the band gaps extreme, that is, when the self-energy potential is defined by Eq. [18] and trimmed by Eq. [22]. The optimal value of \( CUT \), as is the case of an ionic or covalent radius, is typical of each atom, and the orbital that was half-ionized. In most cases only the anion matters.

| Atom | orbital | \( CUT \) (a.u.) | Atom | orbital | \( CUT \) (a.u.) |
|------|---------|----------------|------|---------|----------------|
| Si   | p       | 3.67           | Ge   | p       | 3.46           |
| N    | p       | 2.90           | P    | p       | 3.86           |
| As   | p       | 3.81           | Zn   | d       | 1.665          |
| O    | p       | 2.67           | S    | p       | 3.39           |
| Ga   | d       | 1.23           | In   | d       | 2.126          |

The band gaps calculated with LDA-1/2 are presented in Table IV. Here we must remind that LDA-1/2 is still a scheme to calculate excitations, not the total energy and the equilibrium lattice parameter. Whereas the LDA results exhibit the well known underestimation of the energy gap, LDA-1/2 results present an excellent agreement with experiment. In general, by comparing the theoretical LDA and LDA-1/2 band structures we observe that, as in LDA-1/2 the self-energy is removed, the valence states are now more localized and are pulled down in energy in comparison with the LDA, which results in a larger energy gap.

The LDA-1/2 entries in Table IV require a further explanation. In the cases marked
with two asterisks ** we are adding the trimmed self-energy potential derived from the half-ionized anion p-state, and the trimmed self-energy potential derived from the half-ionized cation d-state. The questions then are why adding the p-correction to the anion and not the s-correction, and why d and not s to the cation. The case of C, Si and Ge, when we used -1/4 and not -1/2 ionization, has been discussed above. Thus there seems to be a certain degree of arbitrariness in a LDA-1/2 scheme. But that is not so because, from what is known from the chemical bonding of these compounds, we could not proceed differently. Further we are always keeping in mind the criterion of an extreme band gap (Eq. [17]). The case of diamond (C) is even more puzzling because we are adding trimmed self-energy s and p potentials to a single atom. In this case we are defining the self-energy potential as in Eq. [11] and approaching the method of Filippetti and Spalding.8 Again, Eq. [17] is our guide.

Figs. 4, 5 and 6 depict the corresponding band structures (BS) along the main symmetry directions of the Brillouin Zone (BZ) for Si, ZnO, and InN comparing the LDA-1/2 with LDA. The zero of energy was placed at the top of the valence band. We chose to show the BS for these semiconductors for two reasons: firstly, silicon is one of the most important semiconductors, secondly, we would like to show the results for cases where the LDA fails completely, as InN, for which LDA gives a semi-metal instead of semiconductor, and ZnO, which became a very interesting material with large band gap, and the LDA predicts an energy gap much smaller than the experimental value. Moreover, for ZnO it is difficult to obtain the correct BS, even performing quasiparticle calculations using GW if the starting point is the standard LDA wavefunctions.26 From our results, we observe that for Si, the LDA-1/2 dispersion relations are similar to the LDA but with the correct band gap energy. For the InN and ZnO, the same behavior as Si occurs, but with some differences concerning the cation d-states. In both InN and ZnO, the semicore cation d states play an important role. In the nitride, the states derived from the atomic 4d (In)-orbital lie close to the bottom of the valence band 2s (N)-like and hybridize with it.27 On the other hand, in ZnO, the cation d states lie approximately in the middle of the valence band. Moreover, recently it was shown that in both cases the d-states interact and hybridize with the top of valence band, and in DFT-LDA there is an underestimation of the binding energies of these semicore d states and consequently an overestimation of their hybridization with the anion. The enhanced p-d coupling then pushes up the valence-band maximum and the energy gap becomes smaller.28,29,30,31 By taking these facts into account, for InN and ZnO, LDA-1/2
corrects not only the top of valence but also the cation d-states. Thus, by comparing the LDA with LDA-1/2 band structures, we observe that the latter has the cation d-orbitals deeper in energy. This effect is more pronounced in InN. In both cases, ZnO and InN, LDA-1/2 is remarkable, leading to values very near the experimental ones. In order to study deeply the influence of the cation d-state in both InN and ZnO, in the Table [V] we present the results for the valence band width (VBW) and energy gap at different levels of the LDA-1/2 procedure. We observe that in the case of ZnO, the O p-state correction increases the VBW and the Zn d-state correction decreases the VBW. The combination of (O-p)+(Zn-d) corrections presents a smaller VBW than the pure LDA calculation. However, both corrections increase the value of the energy gap. The combination of O-p and Zn-d corrections results in energy gap in very good agreement with experiment, which again states the importance of taking into account the cation d-state. It is worth to point out that we obtain this good result, in spite of the fact that the position of Zn d-state (∼5 eV below the top of valence band) is higher in energy than the experimental data (∼7.8 eV).24 In the case of InN, as the d state is deeper than in the case of ZnO, the In d-state correction is more important for the VBW value, while the N-p correction is more important for the energy gap value. Particularly, with (N-p)+(In-d) correction we obtain a value for the energy gap which is in good agreement with experiment. Moreover, our full N-p + In-d LDA-1/2 calculation is in precise agreement with the measured value obtained from x-ray photoemission spectroscopy experiments,35 from which the d state of In atom is found to lie 16.0 eV below the valence band maximum.

In order to analyze the band dispersion in more detail, we also performed calculations to obtain the conduction band effective masses. Thus, now we focus our attention on the electronic structure around the conduction-band minima. We fit a parabola to the curves of energy versus $\mathbf{k}$ around the conduction-band minimum up to 1.0% along the main symmetry directions of the Brillouin Zone. Considering the degeneracies and making weighted averages we obtain the electron effective masses. Table [VI] summarizes the effective conduction band masses for several semiconductors. Since a negative value for the LDA-InN band gap was obtained, it was not possible to calculate an effective mass in that case and only the LDA-1/2 value is shown. We note from the Table that the LDA-1/2 method systematically gives larger electron effective masses than LDA. This is due to the fact that with the LDA underestimation of band gap energy, the $\mathbf{k} \cdot \mathbf{p}$ interaction between valence (VB) and conduc-
tion (CB) bands is stronger leading to smaller effective masses. Therefore, in the cases where
the correction of the energy gap is more pronounced, the difference between the LDA and
LDA-1/2 electron effective masses is larger. This is the case as, e.g., the GaAs and ZnO,
for which the LDA values agree rather poorly with experimental data, and the LDA-1/2
gives excellent agreement with experiment. Moreover, if we take a look at the whole table
we observe that the LDA-1/2 effective masses are generally in very good agreement with
experimental data.

Therefore, the LDA-1/2 not only improves the band gaps as a “scissor operator” approach,
but also provides reliable important band structure-derived properties, such as the effective
masses.

V. SUMMARY

The very important problem concerning the calculation of excitations in solids is ad-
dressed and a method to overcome this problem is developed. The method is inspired in
the simple half-ionization method. The localization of the hole created by promoting an
electron from the valence band to the conduction band follows naturally from the method.
The hole is shown to be representable by a square integrable wavefunction, instead of the
usual Bloch wave hole of band structure calculations.

The major success of this method is its reliable description of excited states in solids,
giving band gap energies, effective masses and band structures in very good agreement with
experiment, even in the cases for which the LDA markedly fails, e.g., ZnO and InN. The
method is not more computationally demanding than the LDA calculations. Moreover, the
method is general and can be applied to a broad class of DFT self-consistent methods,
all-electron and pseudopotential based.

VI. ACKNOWLEDGMENT

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0 and 2006/61448-5) and CNPq.
TABLE IV: Band energy gaps (eV) for several semiconductors obtained with the LDA-1/2 at experimental lattice constant, by using the VASP code and SIESTA (S), compared with pure LDA, GW and experimental results of Ref. 32, except where noted. Direct energy gaps are noted as (d) and indirect as (i). The majority of the LDA-1/2 calculations were obtained using only the trimmed self-energy potential of p-anion, exceptions are noted.

|        | LDA-1/2 | LDA | Exp. | GW       |
|--------|---------|-----|------|----------|
| C (i)  | 5.25(S)*| 4.13| 5.47a| 5.48-5.775|
| C (d)  | 6.75(S)*| 5.54| 7.3a |          |
| Si (i) | 1.137, 1.21(S) 0.51| 1.17a| 1.3218, 0.95-1.105 |
| Si (d) | 2.9, 2.94(S) 2.54| 3.05, 3.40a |
| Ge (i) | 0.70    | 0.08| 0.66-0.74a | 0.66-0.835 |
| AlN (d)| 6.06    | 4.27| 6.23 | 5.83-6.245 |
| GaN (d)| 3.52**  | 1.95| 3.507| 3.15-3.475 |
| InN (d)| 0.95 ** | -0.29| 0.7-1.9 | 0.20-0.332 |
| AlP (i)| 2.79    | 1.47| 2.52 | 2.5918 |
| GaP (i)| 2.36(G-L)** | 1.49(G-X) 2.35 | 2.5518 |
| InP (d)| 1.12**  | 0.50| 1.42 | 1.4418 |
| AlAs (i)| 2.73   | 1.34| 2.24 | 2.1518 |
| GaAs (d)| 1.41   | 0.41| 1.519| 1.2218, 1.40-1.705 |
| InAs (d)| 0.75   | -0.34| 0.417| 0.3118 |
| ZnO (d)| 3.29**  | 0.83| 3.4a | 2.51-3.075 |
| ZnS (d)| 3.68**  | 2.02| 3.91a| 3.21-3.575 |

a Ref. 33
* -1/4p-1/4s
** -1/2p-anion-1/2d-cation

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TABLE V: InN and ZnO valence band width (VBW) and band gap energy values at different levels of the LDA-1/2 calculation procedure. The levels presented are: (i) standard LDA calculation, (ii) LDA-1/2 anion-p correction only, (iii) LDA-1/2 cation-d correction only, and (iv) full LDA-1/2 anion-p + cation-d corrections.

|        | correction | VBW (eV) | band gap (eV) |
|--------|------------|----------|--------------|
| ZnO    | none       | 17.72    | 0.83         |
|        | O-p        | 19.44    | 2.14         |
|        | Zn-d       | 17.01    | 1.48         |
|        | (O-p)+(Zn-d)| 17.28   | 3.29         |
| InN    | none       | 15.25    | -0.29        |
|        | N-p        | 14.49    | 1.16         |
|        | In-d       | 18.23    | -0.49        |
|        | (N-p)+(In-d)| 16.85   | 0.95         |

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TABLE VI: Effective masses (units of electron free mass, $m_e$), for several semiconductors obtained with the LDA-1/2 at experimental lattice constant, compared with pure LDA and experimental results. The calculations were made using the VASP code. The experimental data were extracted from Ref. 32, except where noted. The same trimmed self-energy potential used in Table IV were also used here.

| Electron Effective mass |
|-------------------------|
| LDA-1/2 | LDA | Exp. |
|---------|-----|------|
| AlN     | 0.38| 0.30 | -    |
| GaN     | 0.30| 0.17 | 0.18-0.29 |
| InN     | 0.12| -    | 0.11-0.23 |
| AlP     | 0.256| 0.18 | -    |
| GaP     | 0.17| 0.10 | 0.09-0.17 |
| InP     | 0.088| 0.04 | 0.077-0.081 |
| AlAs    | 0.064| 0.022| 0.06-0.15 |
| GaAs    | 0.064| 0.026| 0.065-0.07 |
| InAs    | 0.047| 0.033| 0.023-0.03 |
| ZnO     | 0.39| 0.14 | 0.3-0.36$^{36}$ |
| ZnS     | 0.26| 0.16 |      |

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FIG. 3: (Color online) The optimized values of the parameter $CUT$, in which we correct only the anion p-state for nitrides, phosphides and arsenides compounds.

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FIG. 4: Calculated band structures for Si (in eV). Dashed lines display LDA and solid lines represent LDA-1/2 results. The zero of energy was placed at the top of the valence band.

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FIG. 5: Calculated band structures for ZnO (in eV). Dashed lines display LDA and solid lines represent LDA-1/2 results. The zero of energy was placed at the top of the valence band.
FIG. 6: Calculated band structures for InN (in eV). Dashed lines display LDA and solid lines represent LDA-1/2 results. The zero of energy was placed at the top of the valence band.