

**A FIRST-PRINCIPLES STUDY OF DEFECTS
AND ADATOMS ON SILICON CARBIDE
HONEYCOMB STRUCTURES**

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By

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August, 2009

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ABSTRACT

A FIRST-PRINCIPLES STUDY OF DEFECTS AND ADATOMS ON SILICON CARBIDE HONEYCOMB STRUCTURES

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M.S. in Materials Science and Nanotechnology

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In this thesis a study of electronic and magnetic properties of two dimensional (2D), single layer of silicon carbide (SiC) in hexagonal structure and its quasi 1D armchair nanoribbons are presented by using first-principles plane wave method. In order to reveal dimensionality effects, a brief study of 3D bulk and 1D atomic chain of SiC are also included. The stability analysis based on the calculation of phonon mode frequencies are carried out for different dimensionalities. It is found that 2D single layer SiC in honeycomb structure and its bare and hydrogen passivated nanoribbons are ionic, non magnetic, wide band gap semiconductors. The band gap further increases upon self-energy corrections. Upon passivation of Si and C atoms at the edges of nanoribbon with hydrogen atoms, the edge states are discarded and the band gap increases. The effect of various vacancy defects, adatoms and substitutional impurities on electronic and magnetic properties in 2D single layer SiC and in its armchair nanoribbons are also investigated. Some of these vacancy defects and impurities, which are found to influence physical properties and attain magnetic moments, can be used to functionalize SiC honeycomb structures for novel applications.

Keywords: ab initio, first principles, silicon carbide, density functional theory, adsorption, substitution, binding, vacancy.

ÖZET

BALPETEĞİ YAPISINDA SiC ÜZERİNDEKİ HATALARIN VE İLAVE ATOMLARIN ETKİSİNİN İLK PRENSİPLERDEN İNCELENMESİ

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Çalışmamda kısmi bir (nanoşerit) ve iki boyutlu bal peteği SiC malzemesini ilk prensiplerden başlayarak inceledim. Boyutsallık etkilerini takip için üç boyutlu SiC allotroplarından bazılarını ve gerçek bir boyutlu SiC zincirlerini çalışmama dahil ettim. Değişik boyutlardaki SiC malzemesinin kararlılığı, fonon hesaplarıyla tahlil edilmiştir. İki boyutlu bal peteği şeklindeki SiC ve nanoşeritleri iyonik, manyetik olmayan, geniş bant aralıklı yarı-iletkenlerdir. Öz enerji (GW_0) hesaplarıyla bu bant aralığı artmıştır. Hidrojen atomları ile doyurulduğunda, “armchair” SiC nanoşeritlerin yanlarındaki atomlardan oluşan bantlar kaybolmakta ve bant aralığı artmaktadır. Atom boşluğu, söğürülen yabancı atomlar ve Si ya da C atomlarının yerini alan yabancı atomların iki boyutlu ve nanoşerit SiC üzerindeki etkileri araştırılmıştır. Bu araştırılan etkilerden bazılarının tek katmanlı SiC malzemesini işlevlendirebileceği saptanmıştır.

Anahtar sözcükler: ab initio, temel prensipler, SiC, durum fonksiyonu teorisi, söğürülme, bağlanma, kristal hataları.

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Chapter 1

Introduction

Owing to its exceptional thermal and physical properties [1], silicon carbide (SiC) is a material, which is convenient for high temperature and high power device applications. Because of its wide band gap, SiC bulk structure has been a subject of active studies in optical and optoelectronic research. Unlike the polymorphs of carbon, SiC is a polar material. In spite of the fact that both constituents are Group IV elements, charge is transferred from Si to C, due to higher electronegativity of C atom relative to Si atom.

Bulk SiC has six commonly used stacking configurations denoted as 3C (zincblende), 2H (wurtzite), 4H, 6H, 15R and 21R. Lubinsky *et.al.* [2] reported optical data, indirect transitions, dielectric function and reflectivity of 3C SiC using first-principles Hartree-Fock-Slater method. A more comprehensive study [3] of SiC comprises lattice constants, Si-C bond distances, band structures along high symmetry points, dielectric function and hence optical properties for all six stacking configurations of bulk SiC. It uses first-principles OLCAO (orthogonalized linear combination of atomic orbitals). In addition, computational modelling for optical characteristics of SiC and their utilities in actual devices was also carried out. Breakdown luminescence spectra, distribution functions with and without interband transition, hole initiated impact ionization coefficients [7] of 4H SiC were obtained using a Monte Carlo simulation model. Experimentally, a second order optical process (i.e two photon absorption) has been observed

in SiC. Furthermore, a SiC photodiode fabricated this way was able to detect small (up to 90 fs) laser pulses.[8] Again with 4H SiC, a Schottky UV photodiode [9], which uses the pinch-off surface effect was constructed. It has an internal efficiency of 78% and 1.8 times the responsivity of the traditional planar metal-semiconductor-metal junction.

As SiC in lower dimensionality, SiO₂ coated SiC nanowires [4] were synthesized and showed favorable photocatalytic behavior. A theoretical work on hydrogen passivated SiC nanowires[5] provided the energy bands both using local density approximation within Density Functional Theory (LDA-DFT) and sp^3s^* LCAO tight binding (TB) methods. SiC-ZnS core-shell structures were also fabricated [6]. 3C SiC (zincblende) nanoparticles were synthesized by carbothermal reduction method [11]. Band gap of zincblende nanoparticles were estimated to be around 3 eV from photoluminescence measurement (Blueshift is present on the order of 0.6 eV due to quantum confinement). With a similar carbothermal method, microribbons [44] with widths in the range of 500 nm - 5 μ m and thickness of 50-500 nm were synthesized. SiC is frequently used as a substrate to grow other materials [14, 15]. Characteristics of those materials (exciton localization, photoluminescence spectra etc.) are then analyzed. Few layers of graphene was also grown on SiC [16]. Not only regular SiC crystals but also SiC clusters (Si_nC_n, n=1-10) were investigated [13] using DFT and generalized gradient approximation (GGA). With the aim of developing a material for future nanoelectronic applications, binding energy, HOMO-LUMO gap, Mulliken charge, vibrational spectrum and ionization potential of Si_nC_n clusters are revealed.

Carbon/carbon (C/C) composites are widely used at high temperature, but they are vulnerable to oxidation. SiC, as a coating material has been proven to prevent oxidation [17].

For transistors used in microcomputers that we use today, silicon is the perfect choice. However, when it comes to high voltages and high currents, SiC is better due to its fundamental characteristics [18].

The appeal of SiC is that it appears in more than 200 structural configurations. Mostly used types, 4H and 6H, have large indirect band gap, large breakdown

electric field as well as high electron mobility and thermal conductivity. Hence, SiC power switch is a more useful alternative to Si [18].

Using metal induced characterization technique, SiC thin films for p-n junction devices were fabricated. Current-voltage (I-V) measurements confirm rectification ability of the junction from -2V to 2V [19].

Readily, 4H SiC DiMOSFET has been fabricated and its electrical properties are given. When compared to Si power MOSFET, SiC DiMOSFET has a five times higher voltage rating without an increase in the specific on-resistance [20].

SiC is a promising candidate for minimally invasive monitoring applications. Superiority of SiC to Si in terms of mechanical response and electrical properties has been confirmed. This fact enables the way to SiC based probes in biomedical applications [21].

To be used in harsh environments and high temperatures, all-SiC capacitive pressure sensor has been developed. The prototype has been tested at pressures up to 700 psi and at temperatures up to 574 degrees. The stability and performance reproducibility of the sensor after tests are promising [22].

1.1 Summary of Projects including Silicon Carbide

- Controlled Drug Delivery
- Micro-electronics / semiconductors
- Schottky Barriers
- p-n junctions
- Magnetized surfaces
- Surface Defects

- Pressure sensors
- Biomedical devices
- High temperature devices
- Field Effect transistors
- Nano electronics
- Doping
- Data storage
- Nanotubes
- Molecular Quantum Wires
- Nano-Interconnects
- Solar cells
- p-i-n diodes
- Solar Storage
- Anti-oxidation Coating
- Spintronics

1.2 Motivation

Because of interesting electronic, mechanical and thermal properties it possesses, bulk SiC drew attention of both scientists and device engineers for past 20 years. During this period of time a new interdisciplinary field - nanotechnology emerged. Nanotechnology not only aims to reduce well known technology to nanoscale but also tries to explore the new application areas, taking advantages of the quantum world. While the two-dimensional heterostructures of semiconductors have a subject of active study leading to the fabrication of exotic devices and quantum

structures, analogous results are expected to occur in 2D honeycomb SiC. In this thesis, we question the existence of strictly two dimensional honeycomb SiC . The main aim of the thesis is to reveal the fundamental properties of SiC nanoribbons and explore ways to functionalize the planar honeycomb SiC.

1.3 Organization of the Thesis

The thesis is organized as follows: Chapter 2 summarizes the basic properties of two dimensional (2D) honeycomb structures including SiC 2D honeycomb, Chapter 3 focuses on the theoretical background and approximation methods. In Chapter 4, our studies and results are presented. Finally , a brief conclusion summarizes the result of our studies.

Chapter 2

Graphene and 2D Honeycomb SiC

Graphene, 2D honeycomb structure of carbon has been the source for the inspiration of all other monolayer honeycomb materials. Advances in materials growth and control techniques have made the synthesis of the isolated graphene [23] and its ribbons [24] in different orientations possible. Recent studies on the quasi one dimensional graphene ribbons revealed interesting size and geometry dependent electronic and magnetic properties [25].

What makes carbon atoms stay planar in honeycomb form is the strong coupling of p_z orbitals. Silicon having a larger radius than carbon makes honeycomb structure by getting slightly buckled (puckered) [27]. This slight buckling enables bonding with the next nearest neighbor of each silicon atom and grants stability.

2D SiC honeycomb consists of one Si and one C in its unit cell, both of which make the honeycomb structures by themselves.

Firstly, a brief introduction to graphene and its nanoribbons are given. In the second part, we will present a brief comparison of 2D SiC honeycomb lattice to graphene and give some experimental results from literature. Detailed quantitative results of 2D SiC honeycomb material will be given in Results section.

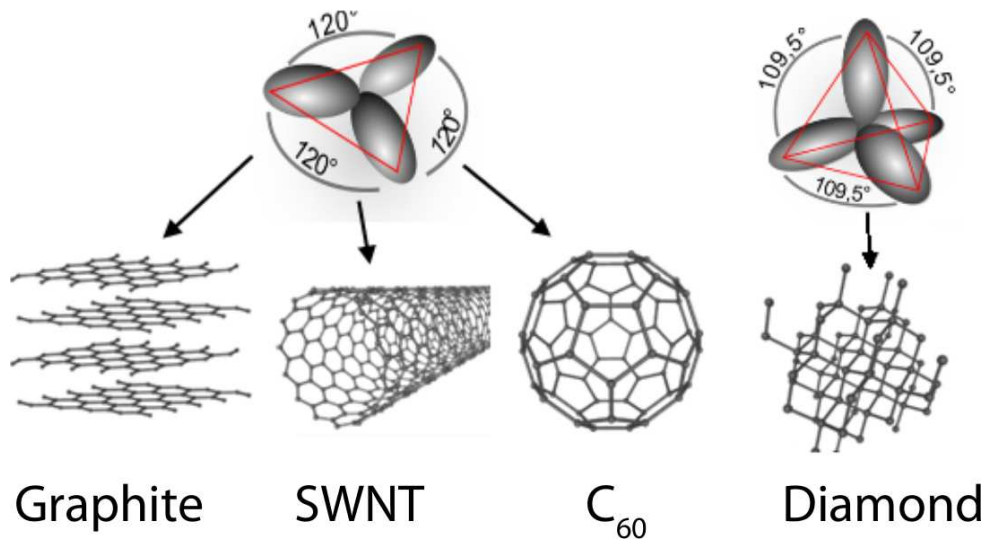


Figure 2.1: (Reproduced from Ref.[26]) Graphene, graphite, single-walled carbon nanotube (SWNT) and C_{60} structures make sp^2 type bonding, whereas diamond makes sp^3 type bonding. Graphite can be viewed as a stack of graphene layers. Carbon nanotubes are rolled up cylinders of graphene and fullerenes are the molecules consisting of wrapped graphene by the introduction of pentagons on the hexagonal structure. The diamond is a transparent crystal of tetrahedrally bonded carbon atoms and crystallizes into the face centered cubic lattice structure.

2.1 Graphene

Graphene, graphite, carbon nanotubes and fullerenes are categorized in carbon-based π electron systems in honeycomb network, which are distinguished from sp^3 -based nanocarbon systems having a tetrahedral network such as diamond. With the sp^2 hybridization of one s-orbital and two p-orbitals results in a triangular planar structure with a formation of a σ -bond between carbon atoms which are separated by 1.42 Å.

The perfect 2D graphene is an infinite network of hexagonal lattice, in contrast to ideal graphene which is a nanosized flat hexagon network with the presence of open actual edges around its periphery. The open edges become important for nanoribbons.

2.1.1 Structure of Graphene

The structure of graphene layers have been explored by using the high resolution microscopy techniques such as Raman [29] and Rayleigh [30]. The graphene structures based on the hexagonal lattice of carbon atoms have been confirmed [31]. Once identified, graphene layers can be processed into nanoribbons by lithography techniques [32].

One carbon atom in honeycomb structure bound to three neighbour through strong, covalent bonds. This configuration gives exceptional structural rigidity within its layers.

The structure is not a Bravais lattice but it can be seen as a triangular lattice with a basis of two atoms per unit cell. The lattice vectors can be written as :

$$a_1 = \frac{a}{2}(3, \sqrt{3}, 0), \quad a_2 = \frac{a}{2}(3, -\sqrt{3}, 0) \quad (2.1)$$

where $a \approx 1.42 \text{ \AA}$ is the C-C distance. The reciprocal lattice vectors are given by :

$$b_1 = \frac{2\pi}{3a}(1, \sqrt{3}, 0), \quad b_2 = \frac{2\pi}{3a}(3, -\sqrt{3}, 0) \quad (2.2)$$

The two points at the corners of graphene's Brillouin zone (BZ) is of special importance. They are named Dirac points. Their positions are given by:

$$K = \left(\frac{2\pi}{3a}, \frac{2\pi}{3\sqrt{3}a}, 0 \right), \quad K' = \left(\frac{2\pi}{3a}, -\frac{2\pi}{3\sqrt{3}a}, 0 \right) \quad (2.3)$$

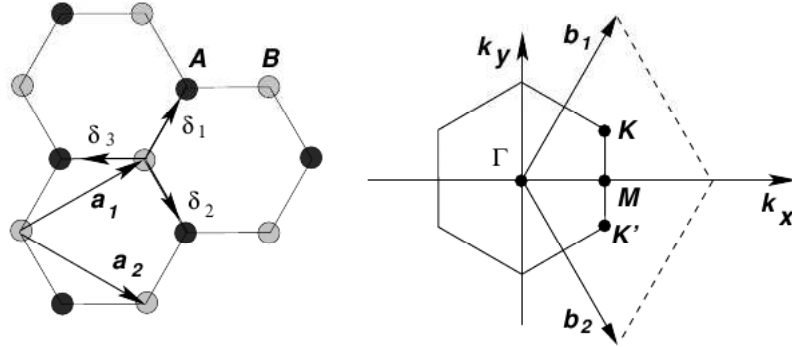


Figure 2.2: (Reproduced from Ref.[26]) Left: Lattice structure of graphene made of two interpenetrating hexagonal lattices (a_1 and a_2 are lattice unit vectors, and δ_i , $i=1,2,3$ are the nearest neighbor vectors); Right: corresponding Brillouin zone. The Dirac corners sit at the K and K' points.

2.1.2 Synthesis

Graphene sheets (a single sheet or a few layer sheet) can mainly be prepared by micromechanical cleaving of graphite crystals according to recent experiments [23, 33, 34] or by epitaxial growth on silicon carbide (SiC) [35, 36]. The first method can be used to obtain high quality of graphene sheets which are comparable to that in graphite, but it is restricted by small sample dimensions and low visibility. On the other hand, the second one is more suitable for large area fabrication and is more compatible with current Si processing techniques for future applications. Nevertheless, the epitaxial graphene was shown to interact with SiC by first principles calculations [37, 38] and experiments [39, 40].

2.1.3 Electronic Properties of Graphene

The investigations of electronic properties of graphene trace back to 1946 when P. R. Wallace wrote the first scientific paper on the band structure of graphene as an approximation trying to understand the electronic properties of more complex, three dimensional (3D) graphite. He did not use the word graphene and referred

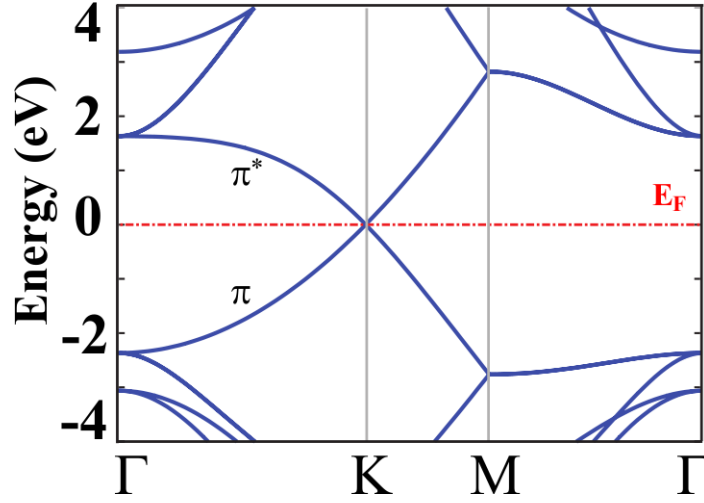


Figure 2.3: (Reproduced from Ref.[26]) Band structure of the bare graphene calculated for the 2×2 unitcell.

to "a single hexagonal layer " [28]. The electrical properties of graphene can be described by a conventional tight-binding model; in this model the energy of the electrons with wavenumber \mathbf{k} is

$$E = \pm \sqrt{\gamma_0^2 \left(1 + 4 \cos^2 \pi k_y a + 4 \cos \pi k_y a \cdot \cos \pi k_x \sqrt{3} a \right)} \quad (2.4)$$

[28].

with the nearest-neighbour-hopping energy $\gamma_0 \approx 2.8$ eV. $+$ and $-$ corresponds to the π^* and π energy bands, respectively. Figure 2.3 shows the band structure of 2D graphene. The energy dispersion around K is linear in momentum, $E = \hbar k v_f$, as if the relation for relativistic particles (like photons). In this case the role of the speed of light is given by the Fermi velocity $v_f \approx c/300$. Because of the linear spectrum, one can expect that particles in graphene behave differently from those in usual metals and semiconductors, where the energy spectrum can be approximated by a parabolic dispersion relation.

2.1.4 Graphene Nanoribbons

Graphene nanoribbons can be thought of as single wall CNTs cut along a line parallel to their axis and then unfolded into a planar geometry. There are two main shapes for graphene nanoribbon edges, namely armchair and zigzag edges. We can cut a graphene sheet in two different line with a difference of 30° in the axial direction between the two edge orientations to produce armchair and zigzag graphene nanoribbons (see Fig. 2.4). If a ribbon is restricted by one of these edges, it is defined either as an armchair GNR (AGNR) or as a zigzag GNR (ZGNR) (see Fig. 2.4 (a &b)).

The ribbon width can be defined by the number of carbon atoms in the primitive unit cell. Dashed rectangle in Fig. 2.4-a shows an armchair graphene nanoribbon containing 20 carbon atoms in its unitcell. This ribbon can be labelled as AGNR (20). On the other hand the ribbon in part (b) can be labelled as ZGNR (10) since it has zigzag shaped edges. Similar to the carbon nanotubes the width plays a crucial role on the electronic and magnetic properties of graphene nanoribbons.

2.2 Two Dimensional (2D) SiC

2D honeycomb SiC unit cell can be thought of a graphene unit cell with one carbon atom replaced by a silicon atom. As Si has a larger radius than C, the lattice gets extended. Lattice vectors for honeycomb SiC :

$$a_1 = d(\sqrt{3}, 0, 0) \quad a_2 = \frac{d}{2}(\sqrt{3}, 3, 0) \quad (2.5)$$

where $d \approx 1.786 \text{ \AA}$ is the Si-C distance. Apparently, it is larger than C-C distance. The reciprocal lattice vectors are given by :

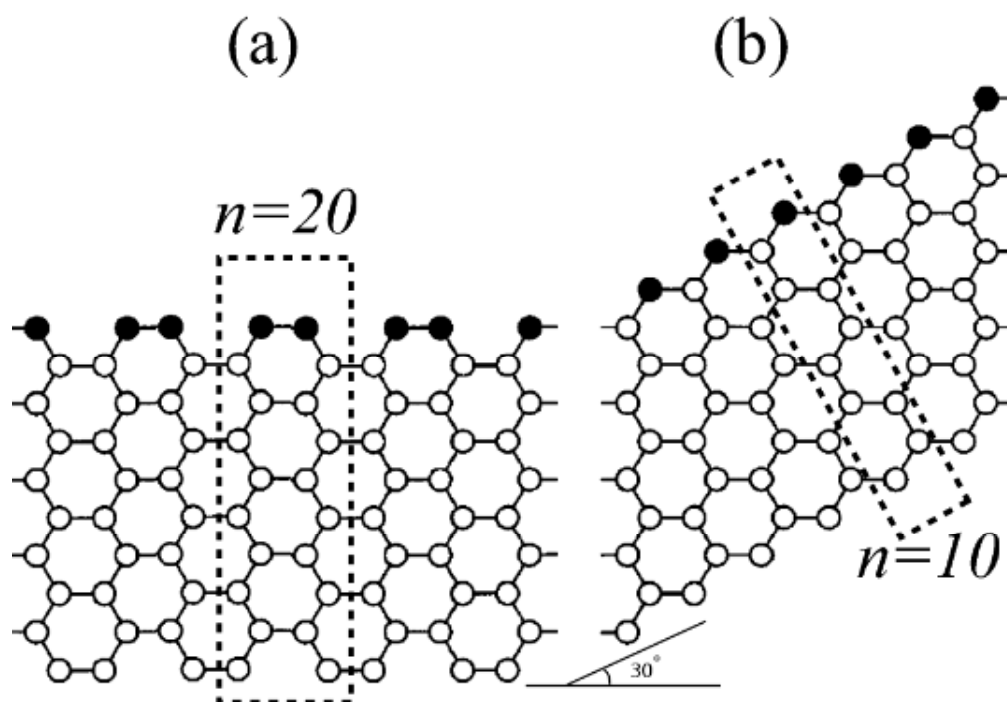


Figure 2.4: (Reproduced from Ref.[26]) Graphene nanoribbons terminated by (a) armchair edges and (b) zigzag edges, indicated by filled circles. The unitcells are emphasized by dashed lines. The width “N” of ribbons are defined as the number of carbon atoms in a unit cell.

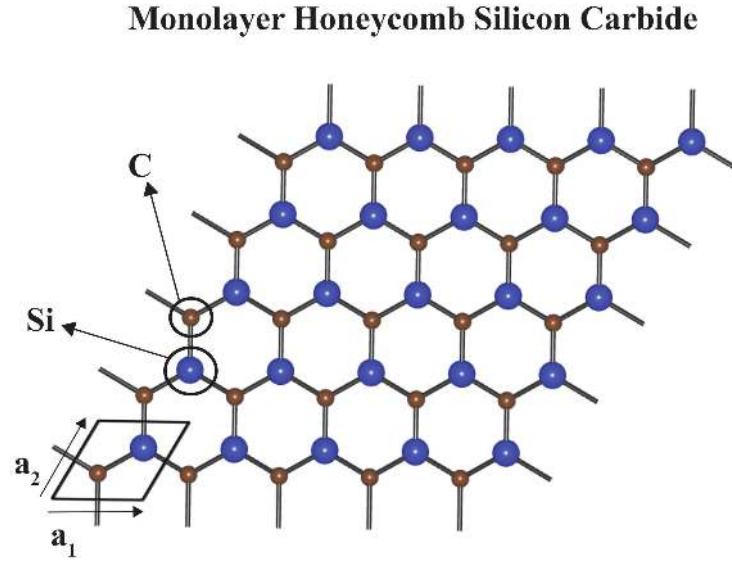


Figure 2.5: Unit cell and lattice vectors of planar honeycomb SiC

$$b_1 = \frac{2\pi}{d} \left(\frac{1}{\sqrt{3}}, -\frac{1}{3}, 0 \right) \quad b_2 = \frac{2\pi}{3d} (0, 2, 0) \quad (2.6)$$

There is no ambiguity in using different lattice vectors from generally used ones for graphene. They are rotationally translated forms of each other and yield the same lattice. General 2D honeycomb SiC plane and its lattice vectors are delineated in Fig. 2.5.

Bare armchair SiC nanoribbons have the similar pattern with those of graphene but have reconstruction at the edges. Upon hydrogen passivation reconstruction vanishes. Zigzag SiC nanoribbons are studied theoretically in a recent paper [41]. Therefore, we did not include them in our study.

Experimental studies of exfoliation and synthesis of single layer SiC do not have the same hasty pace as graphene. In fact, to the best of our knowledge the monolayer SiC has not been obtained up to now. Experimental research has been on honeycomb SiC patches of certain thicknesses.

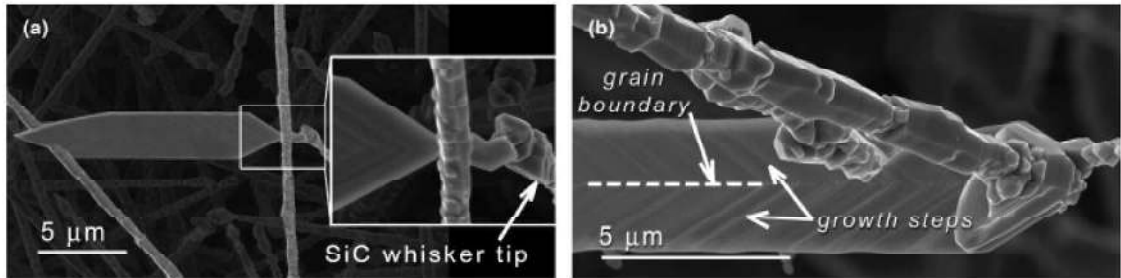


Figure 2.6: (Reproduced from Ref.[44]) Scanning electron microscopy micrographs of a short SiC ribbon growing from the tip of a SiC whisker (a) and a SiC whisker growing from the edge of a SiC ribbon (b).

Carbon-rich SiC nanoribbons were fabricated using a nanosecond pulsed laser direct-write and doping (LDWD) technique [42]. The LDWD technique permits synthesis of heterostructured nanoribbons in a single step without additional material or catalyst, and effectively eliminates the need for nanostructure handling and transferring processes. The resulting nanoribbons comprise three layers each being approximately 50-60 nm thick, containing 15-17 individual sheets.

SiC ribbons were also synthesized by a carbothermal process [44]. the width of the ribbons produced are between 500 nm and 5 μm. The ribbon thicknesses ranged from 50 to 500 nm. Fig. 2.6 and Fig. 2.7 are about the process and the resulting structure .

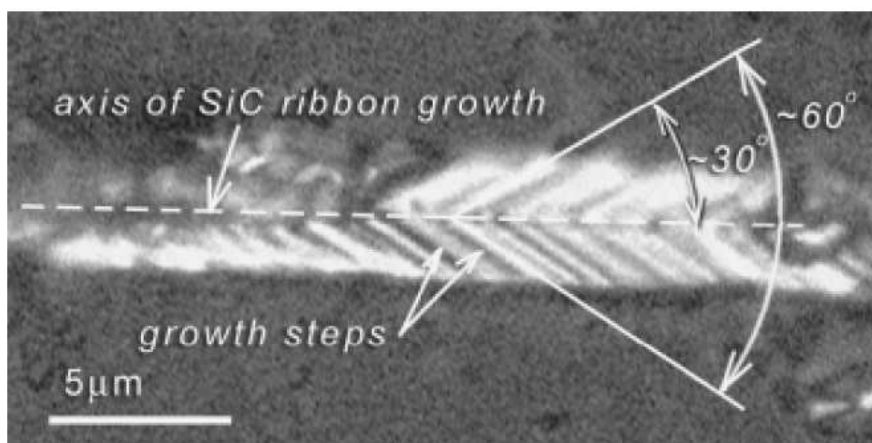


Figure 2.7: (Reproduced from Ref. [44]) Optical micrograph of a SiC ribbon attached to a flat substrate.

Chapter 3

Theoretical Background

3.1 Density Functional Theory

The initial work on DFT was reported in two publications: first by Hohenberg-Kohn in 1964 [47], and the next by Kohn-Sham in 1965 [48]. This was almost 40 years after Schrodinger (1926) had published his pioneering paper marking the beginning of wave mechanics. Shortly after Schrodinger's equation for electronic wave function, Dirac declared that chemistry had come to an end since all its content was entirely contained in that powerful equation. Unfortunately in almost all cases except for the simple systems like He or H, this equation was too complex to allow a solution. DFT is an alternative approach to the theory of electronic structure, in which the electron density distribution $\rho(\mathbf{r})$ rather than many-electron wave function plays a central role. In the spirit of Thomas-Fermi theory [45, 46], it is suggested that a knowledge of the ground state density of $\rho(\mathbf{r})$ for any electronic system uniquely determines the system.

3.1.1 Hohenberg-Kohn Formulation

The Hohenberg-Kohn [47] formulation of DFT can be explained by two theorems:

Theorem 1: The total energy of the system is univocally determined by the electronic density, except for a trivial additive constant.

Since $\rho(\mathbf{r})$ determines $V(\mathbf{r})$, then it also determines the ground state wave function and gives us the full Hamiltonian for the electronic system. Hence $\rho(\mathbf{r})$ determines implicitly all properties derivable from the hamiltonian H through the solution of the time-dependent Schrodinger equation.

Theorem 2: (Variational Principle) The minimal principle can be formulated in terms trial charge densities, instead of trial wavefunctions.

The ground state energy E could be obtained by solving the Schrödinger equation directly or from the Rayleigh-Ritz minimal principle:

$$E = \min\left\{\frac{\langle\tilde{\Psi}|H|\tilde{\Psi}\rangle}{\langle\tilde{\Psi}|\tilde{\Psi}\rangle}\right\}. \quad (3.1)$$

Using $\tilde{\rho}(\mathbf{r})$ instead of $\tilde{\Psi}(\mathbf{r})$ was first presented in Hohenberg and Kohn. For a non-degenerate ground state, the minimum is attained when $\tilde{\rho}(\mathbf{r})$ is the ground state density. And energy is given by the equation:

$$E_V[\tilde{\rho}] = F[\tilde{\rho}] + \int \tilde{\rho}(\mathbf{r})V(\mathbf{r})d\mathbf{r}, \quad (3.2)$$

with

$$F[\tilde{\rho}] = \langle\Psi[\tilde{\rho}|\hat{T} + \hat{U}|\Psi[\tilde{\rho}]\rangle, \quad (3.3)$$

and $F[\tilde{\rho}]$ requires no explicit knowledge of $V(\mathbf{r})$.

These two theorems form the basis of the DFT. The main remaining error is due to inadequate representation of kinetic energy and it will be cured by the Kohn-Sham equations.

3.1.2 Kohn-Sham Equations

There is a problem with the expression of the kinetic energy in terms of the electronic density. The only expression used until now is the one proposed by

Thomas-Fermi, which is local in the density so it does not reflect the short-ranged, non-local character of kinetic energy operator. In 1965, W. Kohn and L. Sham [48] proposed the idea of replacing the kinetic energy of the interacting electrons with that of an equivalent non-interacting system. With this assumption density can be written in atomic units as

$$\rho(\mathbf{r}) = \sum_{s=1}^2 \sum_{i=1}^{N_s} |\varphi_{i,s}(\mathbf{r})|^2, \quad (3.4)$$

$$T[\rho] = \sum_{s=1}^2 \sum_{i=1}^{N_s} \langle \varphi_{i,s} | -\frac{\nabla^2}{2} | \varphi_{i,s} \rangle, \quad (3.5)$$

where $\varphi_{i,s}$'s are the single particle orbitals which are also the lowest order eigenfunctions of Hamiltonian non-interacting system

$$\left\{ -\frac{\nabla^2}{2} + v(\mathbf{r}) \right\} \varphi_{i,s}(\mathbf{r}) = \epsilon_{i,s} \varphi_{i,s}(\mathbf{r}). \quad (3.6)$$

Using new form $T[\rho]$ density functional takes the form

$$F[\rho] = T[\rho] + \frac{1}{2} \int \int \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}d\mathbf{r}' + E_{XC}[\rho], \quad (3.7)$$

where this equation defines the exchange and correlation energy as a functional of the density. Using this functional in Eq. 3.2, we finally obtain the total energy functional which is known as Kohn-Sham functional [48]

$$E_{KS}[\rho] = T[\rho] + \int \rho(\mathbf{r})v(\mathbf{r})d\mathbf{r} + \frac{1}{2} \int \int \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}d\mathbf{r}' + E_{XC}[\rho]. \quad (3.8)$$

In this way we have expressed the density functional in terms KS orbitals which minimize the kinetic energy under the fixed density constraint. In principle these orbitals are a mathematical object constructed in order to render the problem more tractable, and do not have a sense by themselves. The solution of KS equations has to be obtained by an iterative procedure, in the same way of Hartree and Hartree-Fock equations.

3.2 Exchange and Correlation

3.2.1 Local Density Approximation (LDA)

The local density approximation [49] has been the most widely used approximation to handle exchange-correlation energy. Its philosophy was already present in Thomas-Fermi theory. The main idea of LDA is to consider the general inhomogeneous electronic systems as locally homogeneous and then use the exchange-correlation corresponding to that of the homogeneous electron gas.

LDA favors more homogeneous systems. It over-binds molecules and solids but the chemical trends are usually correct.

3.2.2 Generalized Gradient Approximation (GGA)

Once the extent of the approximations involved in the LDA has been understood, one can start constructing better approximations. The most popular approach is to introduce semi-locally the inhomogeneities of the density, by expanding $E_{XC}[\rho]$ as a series in terms of the density and its gradients. This approximation is known as GGA [50] and its basic idea is to express the exchange-correlation energy in the following form:

$$E_{XC}[\rho] = \int \rho(\mathbf{r})\epsilon_{XC}[\rho(\mathbf{r})]d\mathbf{r} + \int F_{XC}[\rho(\mathbf{r}, \nabla\rho(\mathbf{r}))]d\mathbf{r} \quad (3.9)$$

where the functional F_{XC} is asked to satisfy the formal conditions.

GGA approximation improves binding energies, atomic energies, bond lengths and bond angles when compared the ones obtained by LDA.

3.3 Periodic Supercells

By using the represented formalisms observables of many-body systems can be transformed into single particle equivalents. However, there still remains two

difficulties: A wave function must be calculated for each of the electrons in the system and the basis set required to expand each wave function is infinite since they extend over the entire solid. For periodic systems both problems can be handled by Bloch's theorem [51].

3.3.1 Bloch's Theorem

Bloch theorem states that in a solid having translational periodicity each electronic wave function can be written as:

$$\Psi_i(\mathbf{r}) = u_i(\mathbf{r})e^{i\mathbf{k}\mathbf{r}}, \quad (3.10)$$

where u_k has the period of crystal lattice with $u_k(\mathbf{r}) = u_k(\mathbf{r}+\mathbf{T})$. This part can be expanded using a basis set consisting of reciprocal lattice vectors of the crystal.

$$u_i(\mathbf{r}) = \sum_G a_{i,G} e^{i(\mathbf{G})\mathbf{r}}. \quad (3.11)$$

Therefore each electronic wave function can be written as a sum of plane waves

$$\Psi_i(\mathbf{r}) = \sum_G a_{i,k+G} e^{i(\mathbf{k}+\mathbf{G})\mathbf{r}}. \quad (3.12)$$

3.3.2 k-point Sampling

Electronic states are only allowed at a set of \mathbf{k} -points determined by boundary conditions. The density of allowed \mathbf{k} -points are proportional to the volume of the cell. The occupied states at each \mathbf{k} -point contribute to the electronic potential in the bulk solid, so that in principle, an infinite number of calculations are needed to compute this potential. However, the electronic wave functions at \mathbf{k} -points that are very close to each other, will be almost identical. Hence, a single \mathbf{k} -point will be sufficient to represent the wave functions over a particular region of \mathbf{k} -space. There are several methods which calculate the electronic states at special \mathbf{k} points in the Brillouin zone [52]. Using these methods one can obtain an accurate approximation for the electronic potential and total energy at a small number of \mathbf{k} -points. The magnitude of any error can be reduced by using a denser set of \mathbf{k} -points.

3.3.3 Plane-wave Basis Sets

According to Bloch's theorem, the electronic wave functions at each \mathbf{k} -point can be extended in terms of a discrete plane-wave basis set. Infinite number of plane-waves are needed to perform such expansion. However, the coefficients for the plane waves with small kinetic energy $(\hbar^2/2m)|\mathbf{k} + \mathbf{G}|^2$ are more important than those with large kinetic energy. Thus some particular cutoff energy can be determined to include finite number of \mathbf{k} -points. The truncation of the plane-wave basis set at a finite cutoff energy will lead to an error in computed energy. However, by increasing the cutoff energy the magnitude of the error can be reduced.

3.3.4 Plane-wave Representation of Kohn-Sham Equations

When plane waves are used as a basis set, the Kohn-Sham (KS) [48] equations assume a particularly simple form. In this form, the kinetic energy is diagonal and potentials are described in terms of their Fourier transforms. Solution proceeds by diagonalization of the Hamiltonian matrix. The size of the matrix is determined by the choice of cutoff energy, and will be very large for systems that contain both valence and core electrons. This is a severe problem, but it can be overcome by considering pseudopotential approximation.

3.3.5 Nonperiodic Systems

The Bloch Theorem cannot be applied to a non-periodic systems, such as a system with a single defect. A *continuous* plane-wave basis set would be required to solve such systems. Calculations using plane-wave basis sets can only be performed on these systems if a periodic supercell is used. Periodic boundary conditions are applied to supercell so that the supercell is reproduced through out the space. As seen schematically in Fig. 3.1 even molecules can be studied by constructing a large enough supercell which prevents interactions between molecules.

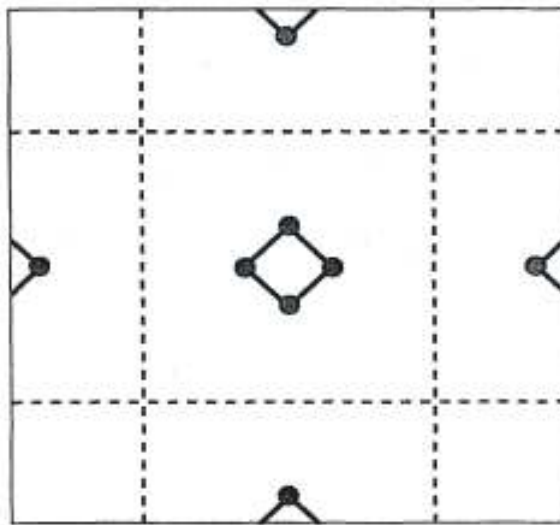


Figure 3.1: (Reproduced from Ref.[43]) Supercell geometry for a molecule. Supercell is chosen large enough to prevent interactions the molecules.

3.4 Pseudopotential Approximation

The act of using a pseudopotential constitutes for replacing the Coulomb potential of the nucleus and the inner core electrons by an effective potential on valence electrons. Once the pseudopotential is generated by an atomic calculation, it can be used to compute material characteristics of an atom in a molecule.

In plane wave calculations, valence wave functions are expanded in Fourier series. The more Fourier components are used the more the computational cost increases. The term “smoothness” means minimizing the range in Fourier space, which is needed to describe the valence properties to a given accuracy. Fig. 3.2 gives a comparison between all-electron and pseudopotential approaches.

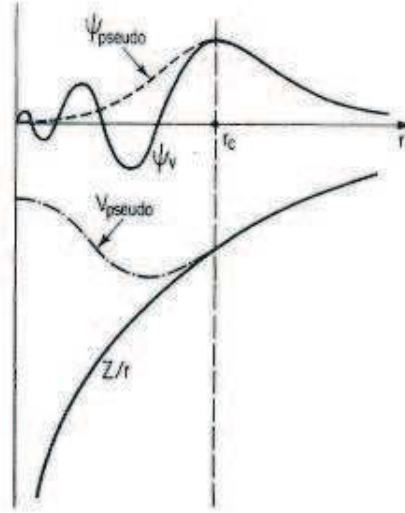


Figure 3.2: (Reproduced from Ref.[43]) Illustration of all-electron (solid lines) and pseudoelectron (dashed lines) potentials and their corresponding wave functions.

3.4.1 Projector Augmented Waves (PAWs)

The projector augmented wave (PAW) method is a reformulation of the OPW (Orthogonalized Plane Wave) method with an adaptation to modern techniques for calculation of total energy, forces etc. It comprises projectors and auxiliary localized functions. The difference with the ultrasoft pseudopotential is that PAW keeps the full all-electron wave function. Due to rapid oscillation of the full wave function around the nucleus, all integrals are evaluated as a combination of integrals of smooth functions.

Main ideas of PAW can be sketched as follows:

Let $|\tilde{\psi}_i^v\rangle$ be smooth part of valence and $|\psi_i^v\rangle$ be all electron valence wave-function. (omit superscript v from now on) The unitary transformation between them is $|\psi_i\rangle = \tau|\tilde{\psi}_i\rangle$.

each of smooth function ($|\tilde{\psi}_i\rangle$'s) in partial waves m can be written

$$|\tilde{\psi}_i\rangle = \sum_m c_m |\tilde{\psi}_m\rangle \quad (3.13)$$

The transformation to all electron function :

$$|\psi_i\rangle = \tau |\tilde{\psi}_i\rangle = \sum_m c_m |\psi_m\rangle \quad (3.14)$$

Full wavefunction in all space :

$$|\psi\rangle = |\tilde{\psi}\rangle + \sum_m c_m \{|\psi_m\rangle - |\tilde{\psi}_m\rangle\} \quad (3.15)$$

τ is linear. The coefficients are :

$$c_m = \langle \tilde{p}_m | \tilde{\psi} \rangle \quad (3.16)$$

with \tilde{p}_m being a projection operator.

Transformation operator becomes:

$$\tau = 1 + \sum_m \{|\psi_m\rangle - |\tilde{\psi}_m\rangle\} \quad (3.17)$$

Any operator A can be transformed to \tilde{A} for smooth part of wavefunctions as :

$$\tilde{A} = \tau^\dagger A \tau = A + \sum_{mm'} |\tilde{p}_m\rangle \{ \langle \psi_m | A | \psi_{m'} \rangle - \langle \tilde{\psi}_m | A | \tilde{\psi}_{m'} \rangle \} \langle p_{m'} | \quad (3.18)$$

Expressions for physical quantities can be derived using Eqs. (3.17) and (3.18)

PAW method can provide the core wavefunctions. From these core wavefunctions, full wavefunctions can be developed [53].

3.5 Phonon Calculations

3.5.1 Phonon frequencies

The central quantity in the calculation of the phonon frequencies is the force-constant matrix $\Phi_{is\alpha,jt\beta}$, since the frequencies at wavevector \mathbf{k} are the eigenvalues of the dynamical matrix $D_{s\alpha,t\beta}$, defined as:

$$D_{s\alpha,t\beta}(\mathbf{k}) = \frac{1}{\sqrt{M_s M_t}} \sum_i \Phi_{is\alpha,jt\beta} \exp(i\mathbf{k}(\mathbf{R}_j^0 + \tau_t - \mathbf{R}_i^0 - \tau_s)), \quad (3.19)$$

where \mathbf{R}_i^0 is a vector of the lattice connecting different primitive cells and τ_s is the position of the atom s in the primitive cell. If we have the complete force-constant matrix, then $D_{s\alpha,t\beta}$ and hence the frequencies $\omega_{\mathbf{k}s}$ can be obtained at any \mathbf{k} . In principle, the elements of $\Phi_{is\alpha,jt\beta}$ are non-zero for arbitrarily large separations $|\mathbf{R}_j^0 + \tau_t - \mathbf{R}_i^0 - \tau_s|$, but in practice they decay rapidly with separation, so that a key issue in achieving our target precision is the cut-off distance beyond which the elements can be neglected.

3.5.2 Calculation of the force constant matrix

We calculate $\Phi_{is\alpha,jt\beta}$ by the small-displacement method. In harmonic approximation the α Cartesian component of the force exerted on the atom at position $\mathbf{R}_i^0 + \tau_s$ is

$$F_{is\alpha} = - \sum_{jt\beta} \Phi_{is\alpha,jt\beta} u_{jt\beta} \quad (3.20)$$

where $u_{js\beta}$ is the displacement of the atom in $\mathbf{R}_j^0 + \tau_t$ along the direction β . The force constant matrix can be calculated by

$$\Phi_{is\alpha,jt\beta} = -\frac{F_{is\alpha,jt\beta}}{u_{jt\beta}} \quad (3.21)$$

by displacing once at a time all the atoms of the lattice along the three Cartesian components by $u_{jt\beta}$, and calculating the forces $F_{is\alpha,jt\beta}$ induced on the atoms in $\mathbf{R}_i^0 + \tau_s$. Since the crystal is invariant under translations of any lattice vector, it is only necessary to displace the atoms in one primitive cell and calculate the forces induced on all the other atoms of the crystal. In what follows we will assume this as understood and put simply $j = 0$.

It is important to appreciate that the $\Phi_{ls\alpha,l't\beta}$ in the formula for $D_{s\alpha,t\beta}(\mathbf{k})$ is the force-constant matrix in the infinite lattice, with no restriction on the wavevector \mathbf{k} , whereas the calculations of $\Phi_{ls\alpha,l't\beta}$ can only be done in supercell geometry. Without a further assumption, it is strictly impossible to extract the infinite-lattice $\Phi_{ls\alpha,l't\beta}$ from supercell calculations, since the latter deliver information only at wavevectors that are reciprocal lattice vectors of the superlattice. The further assumption needed is that the infinite-lattice $\Phi_{ls\alpha,l't\beta}$ vanishes when the separation $\mathbf{R}_{l't} - \mathbf{R}_{l_s}$ is such that the positions \mathbf{R}_{l_s} and $\mathbf{R}_{l't}$ lie in different Wigner-Seitz (WS) cells of the chosen superlattice. More precisely, if we take the WS cell centred on $\mathbf{R}_{l't}$, then the infinite-lattice value of $\Phi_{ls\alpha,l't\beta}$ vanishes if \mathbf{R}_{l_s} is in a different WS cell; it is equal to the supercell value if \mathbf{R}_{l_s} is wholly within the same WS cell; and it is equal to the supercell value divided by an integer P if \mathbf{R}_{l_s} lies on the boundary of the same WS cell, where P is the number of WS cells having \mathbf{R}_{l_s} on their boundary. With this assumption, the $\Phi_{ls\alpha,l't\beta}$ elements will converge to the correct infinite-lattice values as the dimensions of the supercell are systematically increased.

It is not always necessary to displace all the atoms in the primitive cell, since the use of symmetries can reduce the amount of work needed. This is done as follows. We displace one atom in the primitive cell, let's call it 'one', and we calculate the forces induced by the displacement on all the other atoms of the supercell. Then we pick up one other atom of the primitive cell, atom 'two'. If there is a symmetry operation S (not necessarily a point group symmetry operation) such that, when S is applied to the crystal atom two is sent into

atom one and the whole crystal is invariant under such transformation, then it is not necessary to displace atom two, and the part of the force constant matrix associated with its displacement can be calculated using :

$$\Phi_{is,02} = B(S)\Phi_{\lambda_{is}(S),01}B(S^{-1}) \quad (3.22)$$

where $\mathbf{B}(S)$ is the 3×3 matrix representing the point group part of S in Cartesian coordinates, and $\lambda_{is}(S)$ indicates the atom of the crystal where the atom in $\mathbf{R}_i^0 + \tau_s$ is brought because of the action of the symmetry operation S . If there is no symmetry operation connecting atom two to atom one then atom two is displaced and all the induced force field is calculated. The procedure is repeated for all the atoms of the primitive cell.

In principle each atom has to be displaced along the three Cartesian directions. It is sometimes convenient to displace the atoms along some special directions so as to maximize the number of symmetry operations still present in the 'excited' supercell, in this way the calculations of the forces are less expensive. This can always be done, as long as one displaces the atoms along three linearly independent directions. The forces induced by the displacements along the three Cartesian directions is easily reconstructed by the linear combination :

$$F_{is,0t\alpha} = \sum_l A_{l\alpha} \tilde{F}_{is,0tk} \quad (3.23)$$

where $\tilde{F}_{is,0tk}$ is the force induced on the atom in $\mathbf{R}_i^0 + \tau_s$ due to a displacement of the atom in τ_t along the direction \mathbf{u}_k , and $\mathbf{A} = \left(\frac{\mathbf{u}_1}{|\mathbf{u}_1|}, \frac{\mathbf{u}_2}{|\mathbf{u}_2|}, \frac{\mathbf{u}_3}{|\mathbf{u}_3|} \right)^{-1}$ is the inverse of the 3×3 matrix whose columns are the normalized displacements in Cartesian coordinates.

Using symmetries it is possible to reduce the number of displacements even further: if applying a point group symmetry operation U to the displacement vector \mathbf{u}_1 one obtains a vector \mathbf{u}_2 which is linearly independent from \mathbf{u}_1 , then the force field that would be induced by the displacement \mathbf{u}_2 can be calculated by :

$$\mathbf{F}_{is,0t2} = \mathbf{B}(U)\mathbf{F}_{\lambda_{is}(U^{-1}),0t1}. \quad (3.24)$$

a linearly independent direction cannot be found one has to displace the atom along a chosen independent direction and perform an other calculation. This is done until a set of three independent directions is found.

The force constant matrix is invariant under the point group symmetry operations of the crystal. This is not automatically guaranteed by the procedure just described, because in general the crystal is not harmonic. So, the force constant matrix must be symmetrized with respect to the point group operations of the crystal:

$$\Phi_{is,0t} = \frac{1}{N_G} \sum_U \mathbf{B}(U)\Phi_{\lambda_{is}(U),0t}\mathbf{B}(U^{-1}). \quad (3.25)$$

The symmetrization of the force constant matrix removes all even-order anharmonicities. The harmonic approximation becomes better and better as the displacement are made smaller and smaller. However, if the displacements are small, also the force induced are small, but there is a limit in the accuracy achievable in the calculations, so one cannot make too small displacements [55].

Chapter 4

Results

4.1 Introduction

Synthesis of a single atomic plane of graphite, i.e. *graphene* with covalently bonded honeycomb lattice has been a breakthrough for several reasons [23, 56, 57]. Firstly, electrons behaving as if massless Dirac fermions have made the observation of several relativistic effects possible. Secondly, stable graphene has disproved previous theories, which were concluded that two-dimensional structures cannot be stable. Graphene displaying exceptional properties, such as high mobility even at room temperature, ambipolar effect, Klein tunnelling, anomalous quantum hall effect etc. seems to offer novel applications in various fields [58]. Not only 2D graphene, but also its quasi 1D forms, such as armchair and zigzag nanoribbons have shown novel electronic and magnetic properties [59, 60, 61], which can lead to important applications in nanotechnology. As a result, 2D honeycomb structures derived from Group IV elements and Group III-V and II-VI compounds are currently generating significant interest owing to their unique properties.

Boron nitride (BN) in ionic honeycomb lattice which is the Group III-V analogue of graphene has also been produced having desired insulator characteristics [62]. Nanosheets [63, 64], nanocones [65], nanotubes [66], nanohorns [67], nanorods [68] and nanowires [69] of BN have already been synthesized and these

systems might hold promise for novel technological applications. Among all these different structures, BN nanoribbons, where the charge carriers are confined in two dimension and free to move in third direction, are particularly important due to their well defined geometry and possible ease of manipulation.

BN nanoribbons posses different electronic and magnetic properties depending on their size and edge termination. Recently, the variation of band gaps of BN nanoribbons with their widths and Stark effect due to applied electric field have been studied [70, 71]. The magnetic properties of zigzag BN nanoribbons have been investigated [72]. Half-metallic properties have been revealed from these studies which might be important for spintronic applications. Production of graphene nanoribbons as small as 10 nm in width has been achieved [73, 74] and similar techniques are expected to be developed for BN nanoribbons.

Another example of graphene-like structures is monolayer honeycomb zinc oxide (ZnO). The monolayer of ZnO is a III-V ionic compound and has the honeycomb structure similar to graphene and BN. Very thin nanosheets [75], nanobelts [76], nanotubes [77] and nanowires [78] of ZnO have already been synthesized. Two monolayer thick ZnO(0001) films grown on Ag(111) were reported [79].

Quasi one dimensional (1D) ZnO nanoribbons show diverse electronic and magnetic properties depending on their size and edge termination. Recently, it has been predicted that the ferromagnetic behavior of ZnO nanoribbons due to unpaired spins at the edges is dominated by oxygen atoms [80, 81]. Apparently, 2D and 1D honeycomb structure of ZnO can provide us with exceptional electronic and magnetic properties and hence hold the promise of novel technological applications.

In a recent paper, a comprehensive analysis of the atomic, electronic and magnetic properties of monolayer, bilayer and nanoribbons of ZnO are carried out using first-principles calculations. Having presented the analysis on the stability of 2D and 1D ZnO in honeycomb structures, electronic and magnetic properties of single and bilayer ZnO in honeycomb structures were investigated. They are nonmagnetic semiconductors, but attain magnetic properties upon creation of Zn-vacancy defect. ZnO nanoribbons exhibit interesting electronic and magnetic

properties depending on their orientation. While armchair ZnO nanoribbons are nonmagnetic semiconductors with band gaps varying with their widths, bare zigzag nanoribbons are ferromagnetic metals. These electronic and magnetic properties show dramatic changes under elastic and plastic deformation. Hence, ZnO nanoribbons can be functionalized by plastic deformation [82].

Earlier, planar honeycomb structure of carbon was exfoliated and its physical properties were analyzed [23, 56, 57]. The honeycomb structure is common to 2D single layer of SiC and graphene. However, despite graphene being strictly planar, honeycomb structure of Si is unstable, but it is stabilized by puckering. The puckered honeycomb structure of Si is referred to as silicene [27]. The first theoretical study of single layer honeycomb SiC and its selected zigzag and armchair nanoribbons are presented recently [41].

In this thesis, a comprehensive analysis of the atomic, electronic and magnetic properties of 2D monolayer of SiC honeycomb structure and its bare and hydrogen passivated armchair nanoribbons (A-SiCNR) are carried out using first-principles calculations. We started with the discussion of 3D zincblende and wurtzite crystals, as well as SiC atomic chain as an ultimate 1D system; we presented an analysis of optimized atomic structures with corresponding phonon dispersion curves and electronic energy band structures and effective charges. Then we provided an extensive analysis of single layer 2D and quasi-1D (nanoribbon) SiC in terms of optimized atomic structures and their stability, electronic and magnetic structures. We revealed elastic constants, such as in-plane stiffness and Poisson's ratio using our method developed for honeycomb structures. Having made the results for 1D, 2D and 3D structures, we presented a comprehensive discussion of dimensionality effects. Then we investigated the effect of vacancy defects (such as Si and C vacancy, SiC vacancy and C+Si antisite defect) on the electronic and magnetic properties of monolayer SiC and its armchair nanoribbons. Furthermore, we showed that SiC can be functionalized through adsorption of a foreign atom to the surface of 2D SiC or through substitution of either C or Si with a foreign atom. We found that 2D SiC and its ribbons provide new physical properties, which extend those of 3D SiC crystals for technological applications. For example, while various allotropic forms of SiC including its honeycomb structures

are normally nonmagnetic semiconductors, a Si vacancy gives rise to spin polarization. Significant variation of the band gap of narrow A-SiCNRs may be crucial in optoelectronic nanodevices.

4.2 Method of calculations

We have performed first-principles plane wave calculations within Density Functional Theory (DFT) using PAW potentials [83]. The exchange correlation potential has been approximated by Generalized Gradient Approximation (GGA) using PW91 [84] functional both for spin-polarized and spin-unpolarized cases. For the sake of comparison calculations are also carried out using different potentials and exchange-correlation approximations. All structures have been treated within supercell geometry using the periodic boundary conditions. A plane-wave basis set with kinetic energy cutoff of 500 eV has been used. The interaction between SiC monolayers in adjacent supercells is hindered by a minimum 12 Å vacuum spacing. In the self-consistent potential and total energy calculations, when finding the optimum lattice constants the Brillouin zone (BZ) is sampled by, respectively (5×5×5), (11×11×1) and (11×1×1) special \mathbf{k} -points for 3D bulk, 2D honeycomb and 1D nanoribbons of SiC. Further relaxation is made with (11×11×11), (31×31×1) and (25×1×1) special \mathbf{k} -points in order to find the ultimate structure and charge density. All atomic positions and lattice constants are optimized by using the conjugate gradient method, where the total energy and atomic forces are minimized. Pressure on the cell was minimized automatically during this process. The convergence for energy is chosen as 10^{-5} eV between two steps and the maximum Hellmann-Feynman forces acting on each atom is less than 0.04 eV/Å upon ionic relaxation. The pseudopotentials corresponding to 4 valence electrons of Si (Si:3s² 3p²) and C (C:2s² 2p²) are used. Numerical plane wave calculations are performed by using VASP [85, 86]. Part of the calculations have also been repeated by using SIESTA [88] software. The cohesive energy of any structure SiC is found as $E_C = E_T[SiC] - E_T[Si] - E_T[C]$ in terms of the optimized total energy of any SiC structure, and the total energies of free Si and C atoms, all calculated in the same supercell using the same parameters. Phonon calculations

Table 4.1: Si-C bond length (d), lattice constant (a), kink angle (α), bandgap (E_G), cohesive energy (E_c) values for two different types of SiC chains

Type	d (Å)	a (Å)	α (degree)	δq (e)	E_G (eV)	E_c (eV)
Linear	1.649	3.298	180	2.28	1.996	8.923
Wide Zigzag	1.673	3.268	155.2	2.15	1.778	8.963

were carried out using PHON program [87] implementing force constant method.

4.3 1D atomic chain and 3D bulk crystal of SiC

In this section, we present a brief discussion of 1D SiC atomic chain and 3D bulk crystal based on our structure optimized total energy, phonon and electronic energy calculations. Studies [3, 2] on SiC bulk lattice and atomic chains [89] already exist in the literature. However, our purpose is, however to carry out calculations with same parameters as used in 2D single layer SiC honeycomb structure and provide a consistent comparison of dimensionality effects.

4.3.1 1D SiC Chains

Earlier, the first theoretical study of Group IV and III-V binary compounds were reported by Durgun et. al [89]. They examined SiC atomic chain as a function of lattice parameter and found that the wide zigzag atomic chain of SiC with bond angle of $\sim 147^\circ$ is energetically more favorable than the linear and narrow angle zigzag chains. Present calculations find that the atomic chains of SiC are nonmagnetic. Calculated structural parameters, cohesive energies, band gap and phonon modes of linear and zigzag atomic chains, which are relevant for the present study are given in Fig. 4.1. The lattice constant of the linear SiC chain, $a = 3.298$ Å, Si-C bond distance, $d = 1.649$ Å. The charge transfer from Si to C is $\delta q = 2.28$ electrons, calculated by using Bader analysis [90]. Important characteristics of SiC chains are summarised in Table 4.1.

Phonon modes calculated with force constant method have imaginary frequencies : two acoustical and two optical branches of linear SiC chain have imaginary frequencies. Wide angle zigzag SiC chain has one optical and one acoustical branches with imaginary frequencies. These results indicate that free standing SiC chains are unstable.

4.3.2 3D SiC Crystals

Our work on bulk SiC includes wurtzite (wz) and zincblende (zb) structures. Atoms in wz- and zb-SiC are four fold coordinated through tetrahedrally directed sp^3 -orbitals. Calculated structural parameters, cohesive energies(eV), energy band structures and phonon modes are given in Fig. 4.2. Zincblende SiC structure in T_d symmetry has cubic lattice constants are $a_1 = a_2 = a_3 = 3.096 \text{ \AA}$. Si-C bond distance(\mathbf{d}) is 1.896 \AA . There are four tetrahedrally coordinated bonds from each atom, all of which are equal. Charge transfer from Si to C is $\delta q = 2.59$ electrons calculated via Bader analysis [90]. While the GGA band gap is 1.406 eV, it increases to 2.4 eV after GW_0 [91] correction is taken into account. As for wz-SiC crystal, the hexagonal lattice constants of the optimized structure in equilibrium are $a_1 = a_2 = 3.093 \text{ \AA}$, $c/a = 1.63307$. The deviation of c/a from the ideal value of 1.633 imposes a slight anisotropy on the lengths of tetrahedrally directed Si-C bonds. While the length of three short bonds is 1.892 \AA , the fourth bond is slightly longer and has the length of 1.9 \AA . Charge transfer from Si to C is $\delta q = 2.63$ electrons. The GGA band gap is 2.316 eV, but it increases to 3.32 eV after GW_0 correction. Corrected badgaps are exactly the same as those from the experiment [92]. The calculated structural parameters and energy band gaps are in reasonable agreement with the earlier calculations and experimental measurements.

Five out of six phonon modes of SiC zincblende are in complete agreement with literature [93]. The highest optical mode goes down and couples with nearest optical mode. The same problem is present in the wurtzite SiC. Again other five phonon modes of wurtzite structure agrees with literature [94].

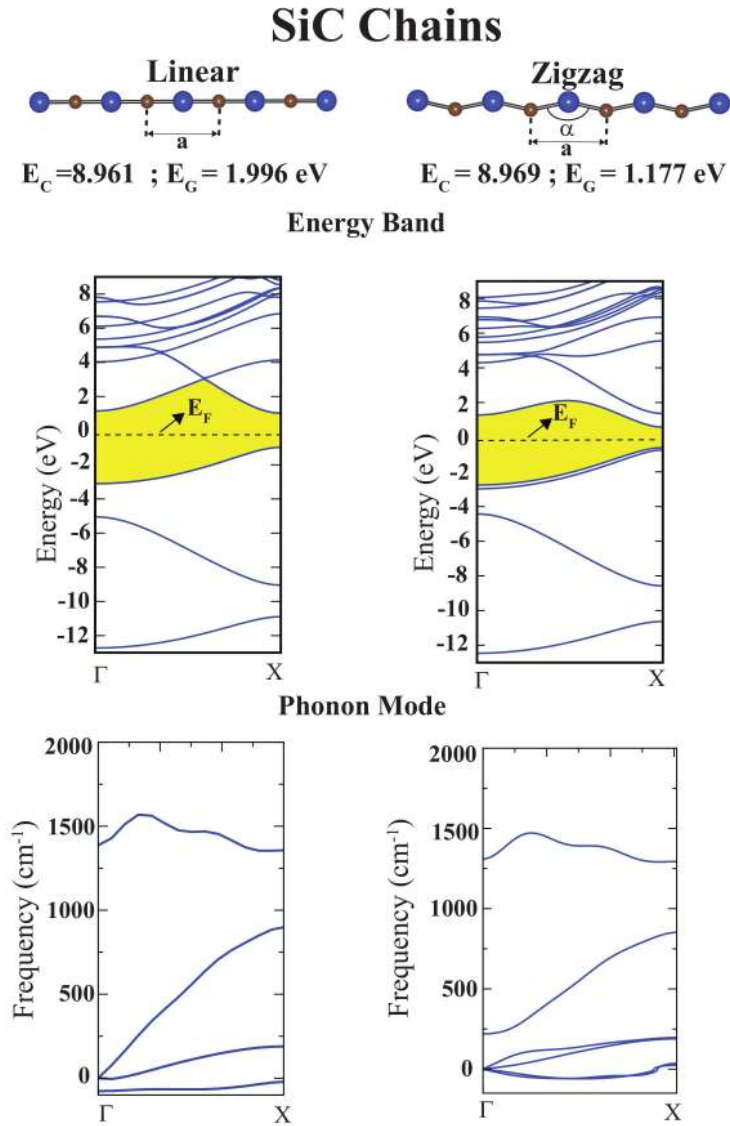


Figure 4.1: Atomic structures, electronic energy bands and dispersion of phonon modes of linear and zigzag SiC atomic chains. E_c and E_G are cohesive and band gap energies, respectively. Si and C atoms are shown by blue-large and black small balls, respectively.

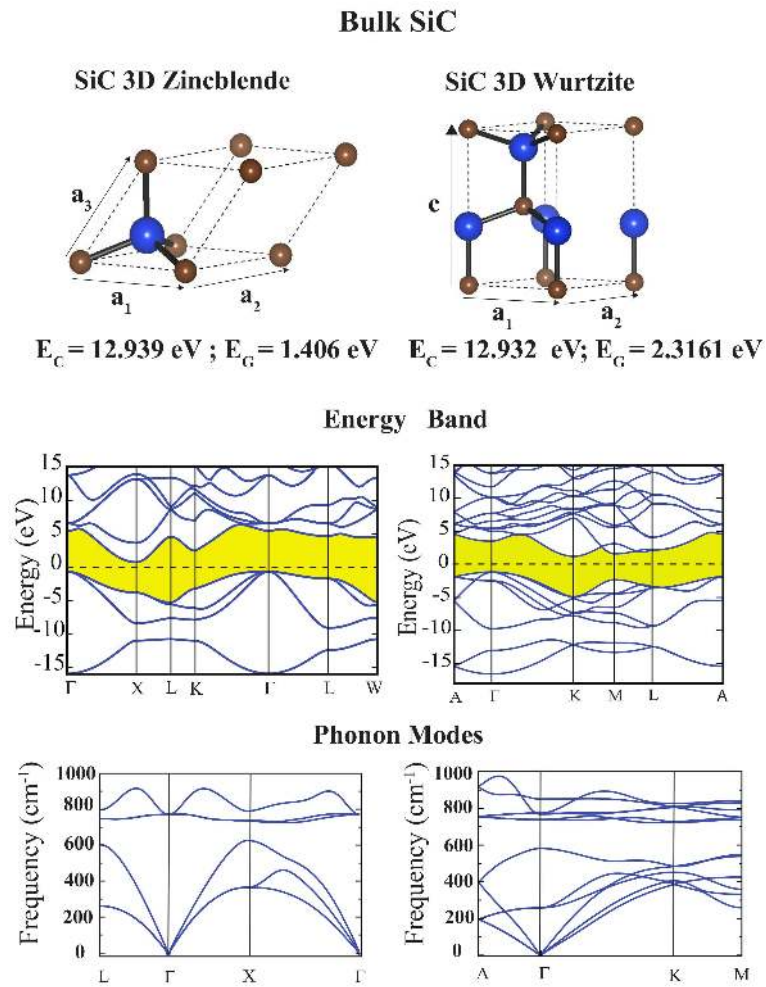


Figure 4.2: Optimized atomic structure with relevant structural parameters, corresponding energy band structure and frequencies of phonon modes of 3D bulk SiC in zincblende and wurtzite structures. Zero of energy of the band structure is set at the Fermi level, and band gap is shaded.

Table 4.2: Si-C bond length(d), lattice constant (a), bandgap (E_G), cohesive energy (E_c) values for the monolayer SiC calculated with different potentials

Potential	d (Å)	a (Å)	E_G (eV)	E_c (eV)
PAW + GGA	1.786	3.094	2.53	11.94
PAW + LDA	1.77	3.07	2.51	13.54
US + GGA	1.776	3.079	2.542	11.97
US + LDA	1.759	3.048	2.532	13.47

4.4 2D Honeycomb SiC

Two-dimensional hexagonal structure of SiC is optimized using periodically repeating supercell having 15 Å spacing between SiC planes. The minimum of total energy occurred when Si and C atoms are placed in the same plane forming a honeycomb structure. The magnitude of the Bravais vectors of the hexagonal lattice is found to be $a_1 = a_2 = 3.094$ Å (see Fig. 4.3), and the Si-C bond to be $d = 1.786$ Å. The planar structure of 2D SiC is tested by displacing Si and C atoms arbitrarily from their equilibrium positions and then reoptimizing the structure. Upon optimization, the displaced atoms returned to their original positions in the same plane. 2D honeycomb SiC is found to be a semiconductor with a GGA band gap of 2.53 eV using LDA as well as GGA. Furthermore, in Table 4.2, we give lattice constant, bond length, cohesive energy, energy gap values of SiC honeycomb structure calculated with different parameters. Since DFT usually underestimates the band gap of semiconductors, we corrected the GGA band gap using GW_0 correction and found it to be 3.90 eV. The charge transfer from Si to C in 2D honeycomb SiC is calculated to be $\delta q = 2.53$ eV electrons. Earlier DFT calculations [41] on 2D honeycomb SiC and its ribbons used PAW [83] potentials and GGA approximation via VASP [85, 86]. Their results are in reasonable agreement with ours.

Our phonon modes were calculated with force constant method [87]. Since all the modes are positive, a single layer SiC sheet is stable. Forces were found by displacing a single atom in a $7 \times 7 \times 1$ supercell by 0.4 Å. We use a small displacement in order to stay in the harmonic region. We increased default grids used

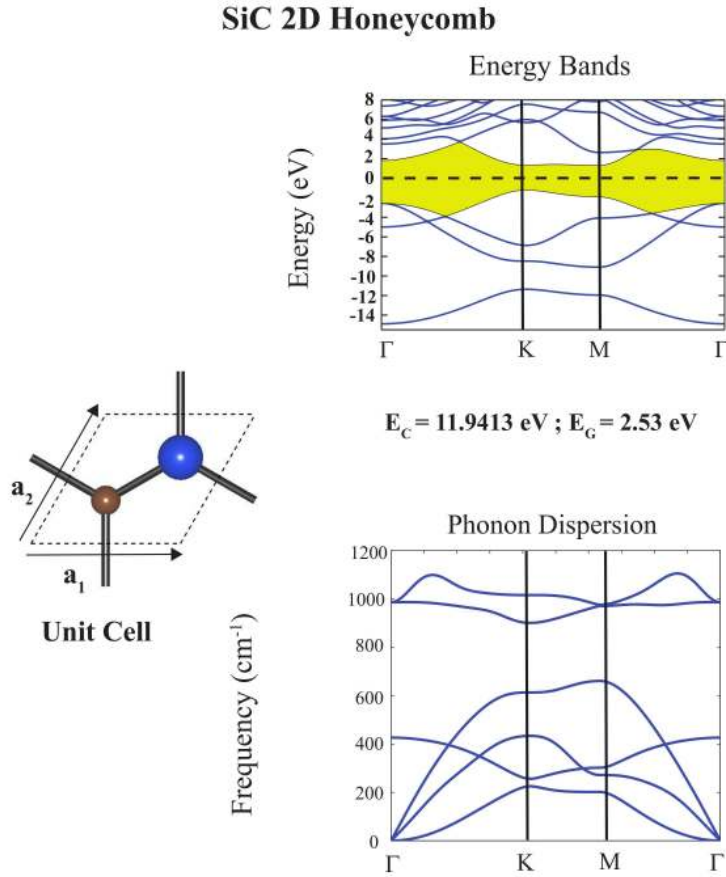


Figure 4.3: Optimized atomic structure, energy band structure and phonon modes of 2D SiC in honeycomb structure. The primitive unit cell is delineated. The zero of energy in the band structure is set to the Fermi level.

by VASP until calculations converge to a sensible result . Since problem arises due to the lowest acoustic mode which presents an out of plane displacement. A fictitious dip near Γ point occurs in crude calculations, but it can be overcome by refining the mesh in the z - axis (perpendicular to plane) as much as possible. By this way, force in that direction is calculated more rigorously.

4.4.1 Dimensionality effects

In Table 4.3, we compare the variation of the effective charge on Si and C atoms, namely Z_{Si}^* and Z_C^* respectively; charge transfer from Si to C, $\delta q = Z_{Si}^* - 4$;

Table 4.3: Si-C bonding type, bond length(d), lattice constant (a), charge transfer (δq), bandgap (E_G), cohesive energy (E_c) comparison for SiC polymorphs

Material	Bonding	d (Å)	a (Å)	δq (e)	E_G (eV)	E_c (eV)
Linear Chain	sp	1.649	3.298	2.28	1.996	8.923
Wide Zigzag Chain	sp	1.673	3.268	2.15	1.778	8.963
2D Honeycomb	sp^2	1.786	3.094	2.53	2.53	11.94
Zincblende	sp^3	1.896	3.096	2.59	1.406	12.939
Wurtzite	sp^3	1.892 (3), 1.9 (1)	3.093	2.63	2.3161	12.932

Si-C bond length d ; lattice constant a , cohesive energy per Si-C, E_c ; band gap E_G calculated for SiC in different dimensionalities. It should be noted the length of Si-C bonds of 2D SiC honeycomb structure is smaller than that in the 3D bulk (wz, zb) crystals, but larger than that in zigzag atomic chains. Here we see that the dimensionality effect is reflected to the strength of the bonding through sp^n hybridization, where n coincides with the dimensionality. While sp^2 hybrid orbitals of 2D planar honeycomb structure form stronger bonds than tetrahedrally coordinated sp^3 orbitals of 3D bulk, they are relatively weaker than sp hybrid orbitals of 1D chain.

4.5 Bare and Hydrogen Passivated SiC Nanoribbons

In this section, we consider bare and hydrogen passivated (H-pass) armchair SiC nanoribbons (A-SiCNR). These nanoribbons are specified according to their width given in terms of N number of Si-C atom pairs in their unit cells. Hence, A-SiCNR(N) indicates armchair SiC nanoribbons having N Si-C pairs in their unit cell. We have analyzed odd numbered A-SiCNR (both bare and H-passivated) from 5 to 21, which have reflection symmetry with respect to their axis. We investigated their atomic structure; electronic, magnetic properties. Bare armchair

SiC ribbons are ferromagnetic in ideal honeycomb form. However, upon structural relaxation, reconstruction occurs at the edges resulting in a considerable gain of energy and the structure becomes nonmagnetic.

The band gap of the bare A-SiCNR increases by ~ 0.7 eV upon saturation with hydrogen atoms, since the edge states are removed from the gap. The band gap variation of both bare and hydrogen saturated armchair SiC ribbons cannot be reconciled with the quantum size effects, since E_G increases with increasing N . This is due to other effects which overcome the quantum confinement effect. For $N > 9$ the variation of E_G of bare SiC armchair ribbons is not significant. Band gaps of both bare and H-passivated SiC armchair ribbons are depicted in Fig. 4.4.

The bare A-SiCNR(9) is an indirect band gap semiconductor. Two bands at the conduction band edge are due to edge states, which are split due to edge-edge interaction. These bands are removed upon H-saturation of dangling bonds of atoms at the edges leading to the widening of the band gap. As for the other edge state band, it is located in the valence band. This band is also removed upon H-saturation, but the valence band edge is not affected. The band gap of H-saturated A-SiCNR(9) is direct. Energy bands and band decomposed charge densities are in Fig. 4.5.

Lengths of Si-H and C-H bonds of H-passivated A-SiCNR's are 1.49 Å and 1.09 Å, which is present in the previous work [41]. Our bandgap variations with respect to ribbon width of H-passivated SiC armchair ribbons are similar and they call it three-family behavior [41]. The main difference with graphene is that the band gaps of H-passivated A-SiCNR's are increasing. Hypothetically the bandgap will reach the value of 2D planar SiC (2.53 eV) for an infinitely large nanoribbon.

In H-saturated A-SiCNR's, Si-C bonds get shortened at most by 0.05 Å at the edges; while in bare A-SiCNR's this shortening is up to 0.9 Å. Hence, edge reconstruction exist in each case but is more pronounced in bare nanoribbons.

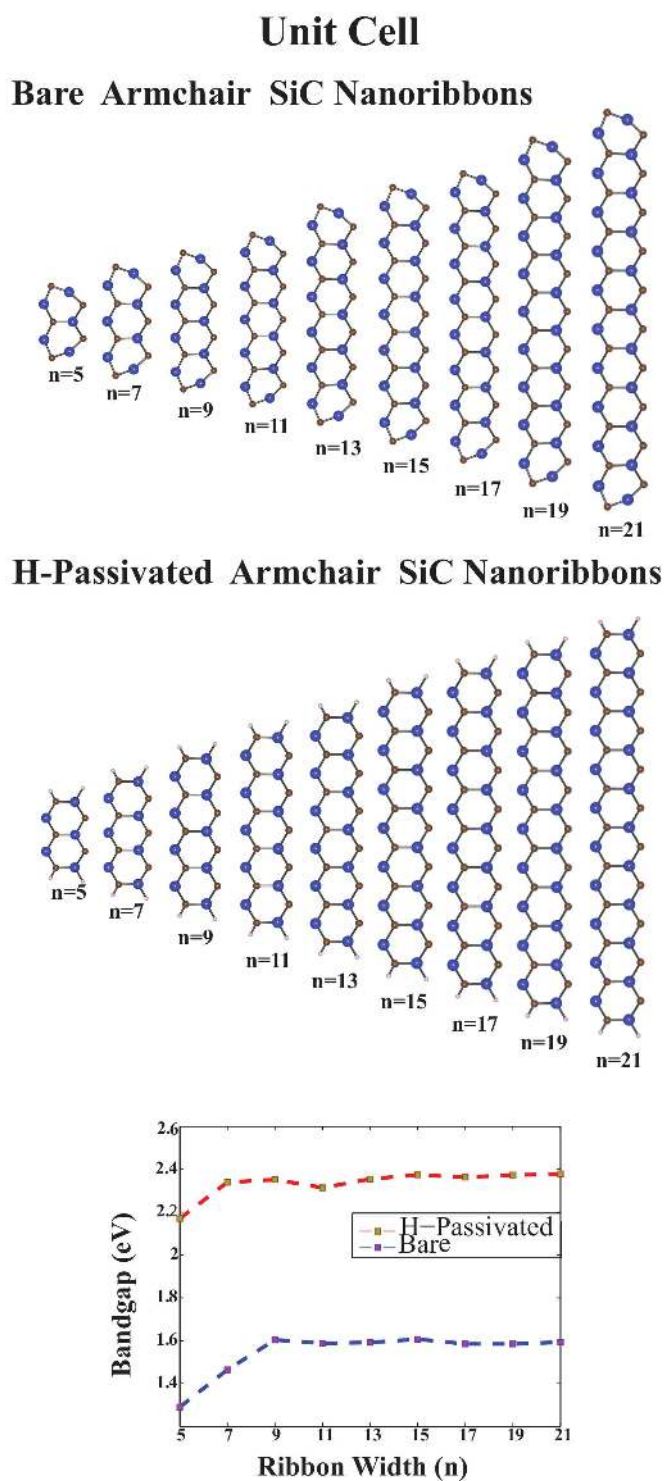


Figure 4.4: Optimized atomic structures and band gap variations of bare and H-passivated SiC nanoribbons with $5 < N < 21$.

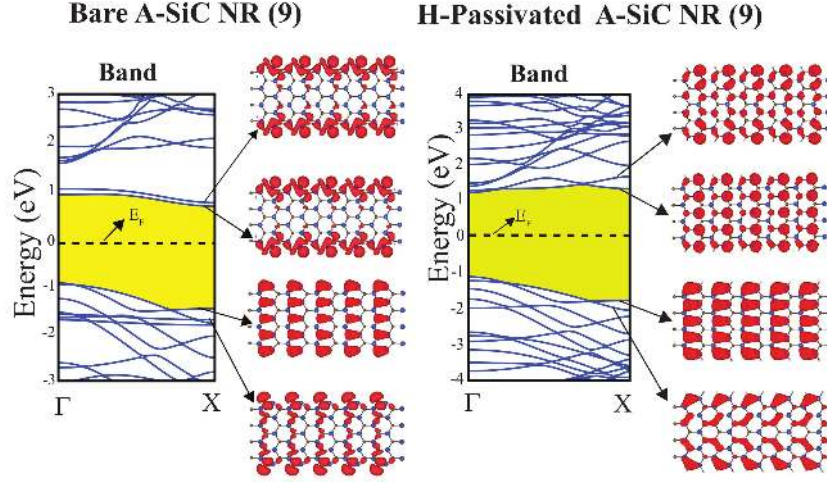


Figure 4.5: Energy band structure of bare and hydrogen saturated armchair SiC nanoribbons, A-SiCNR (N) with $N = 9$ and isosurfaces of charged densities of selected states. Zero of energy is set at the Fermi level.

4.6 Vacancy Defects and Antisite

It has been shown that the vacancy defects have remarkable effects on 2D graphene honeycomb structure and its nanoribbons [97, 98, 99, 100, 101, 102]. Non magnetic graphene sheets or nanoribbons can attain spin polarized states due to vacancy defects. We expect that similar effects of vacancy defects can occur in the electronic and magnetic properties of SiC honeycomb structure.

4.6.1 2D Honeycomb SiC

We investigated the effects of Si and C vacancies, Si+C-divacancy and C-Si-antisite in periodically repeating (7×7) supercells. The size of supercell is optimized to allow negligible defect-defect interaction between adjacent cells. Here the width of the flat bands derived from the states of periodically repeating vacancies is taken as the measure of the strength of vacancy-vacancy coupling. A (7×7) supercell led to rather flat defect bands which are suitable for our purpose, but allowed us to carry out numerical calculations. Our results are presented in Fig. 4.6 for single C, Si, Si+C and C+Si antisite vacancy defects.

Vacancies and Antisite in 7x7 Honeycomb SiC

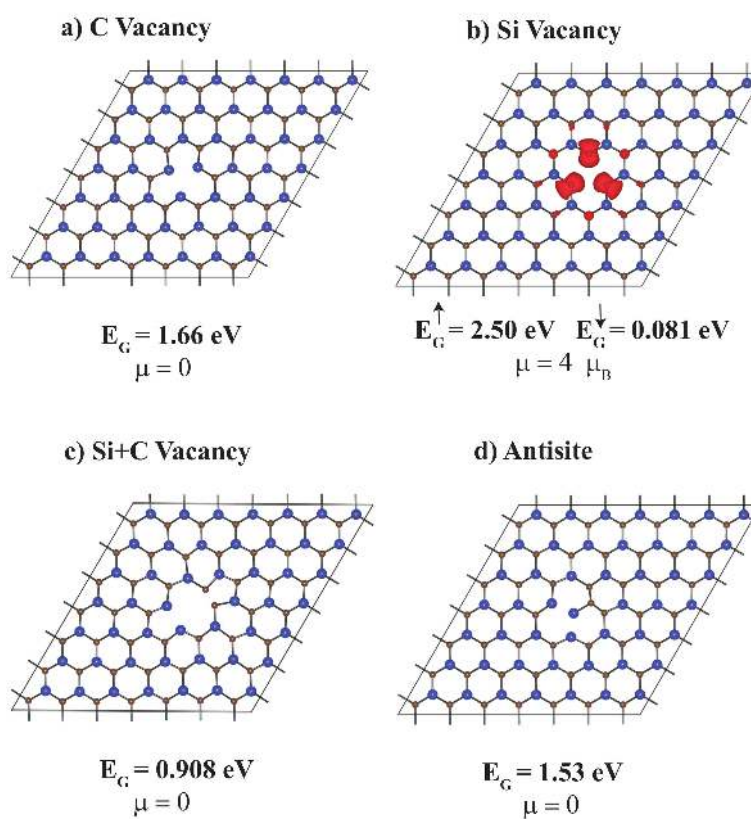


Figure 4.6: Energy band gap and magnetic moment of vacancy defects calculated in a (7×7) supercell of 2D SiC honeycomb structure. (a) C vacancy; (b) Si vacancy; (c) Si+C divacancy; (d) C-Si antisite. In (b) the difference of spin up and spin down charges are shown. Large/blue and small/gray balls represent Si and C atoms respectively.

A vacancy is generated first by removing a single atom C or Si atom from each supercell of the monolayer of SiC in honeycomb structure as shown in Fig. 4.6 (a) and (b). Subsequently, the atomic structure is optimized. Single C vacancy in 2D monolayer of SiC is nonmagnetic; Si atoms around vacancy with coordination number 2 are displaced in the transversal direction and do not induce any magnetic moment. The band gap is altered to 1.66 eV, as three obvious defect states occur in the original band gap.

As for Si vacancy, three C atoms around vacancy remained planar. Similar to the vacancies in graphene and BN, Si-vacancy induces a local magnetization in the system. Isovalue surfaces of the difference between spin up and spin down charge densities i.e. $\Delta\rho^{\uparrow\downarrow}$ clearly shows a spin polarization and a net magnetic moment constructed therefrom. The calculated total magnetic moment is $4 \mu_B$ per supercell. The Si-vacancy in a repeating (7×7) supercell also modifies the electronic structure. While the band gap of spin unpolarized (defect free) SiC is practically unaltered for the spin-up states, it is lowered from 2.53 eV to 0.081 eV for the spin down states. This can be understood as a spin up donor stable near the conduction band. As for Si+C divacancy in Fig. 4.6 (c), it is again nonmagnetic. However, two C atoms around the vacancy choose to make a bond with each other. The band gap is also modified. Two totally separate states occur in the original band gap and band decomposed charge densities of these states are located around the vacancy. Finally, we consider the antisite effect. The resulting relaxed structure is given in Fig. 4.6 (d). Lattice is distorted as C-C bond is shorter than Si-Si bond in the antisite case. It is noted that the calculated magnetic moments for single Si- and C-vacancy do not agree with Lieb's theorem [103], which normally predicts $1 \mu_B$ net magnetic moment both for Si and C vacancies in Fig. 4.6. We attribute the discrepancy between the results of first principles calculations and Lieb's theorem to the structural relaxation after the generation of vacancy into account.

4.6.2 Vacancy defects in SiC Nanoribbons

We examined the effect of the vacancy defects on the electronic and magnetic properties of hydrogen passivated A-SiCNR(9) using a (4×1) repeating supercell. Our main motivation was to investigate what differences would occur in a ribbon. Final structures, band gaps, total magnetic moments are given in Fig. 4.7. Overall effects are similar to those in 2D single layer SiC honeycomb structure except the antisite case in which 4×1 hydrogen passivated SiC armchair nanoribbons where the exchanged Si atom moves into the plane about 0.9 \AA . Two separate states, one in the conduction band and one in the valence band occur due to antisite and decrease the band gap. A straightforward conclusion is that vacancy and antisite states are localised and do not couple to the hydrogen states at the edges.

4.7 Functionalization of SiC honeycomb structure by adatoms

Like vacancy, adatom adsorption as well as substituting Si or C atoms in the honeycomb structure by foreign atoms can modify the properties of 2D monolayer of SiC and its nanoribbons. This way SiC honeycomb structure can be functionalized. Adatom adsorption and substitution are considered within the periodically repeating (7×7) supercell geometry to minimize the interaction between them. That the coupling between adatoms are negligible are tested using (10×10) supercell geometry.

4.7.1 Adatom adsorption

As adatom we considered Al, Co, Fe, N, P, Ti by placing each of them on four different positions (Fig. 4.9) in the (7×7) monolayer honeycomb SiC surface and then by fully relaxing the whole system. The initial positions of the adsorption are on top of silicon atom (TS), on top of carbon atom (TC), at the center

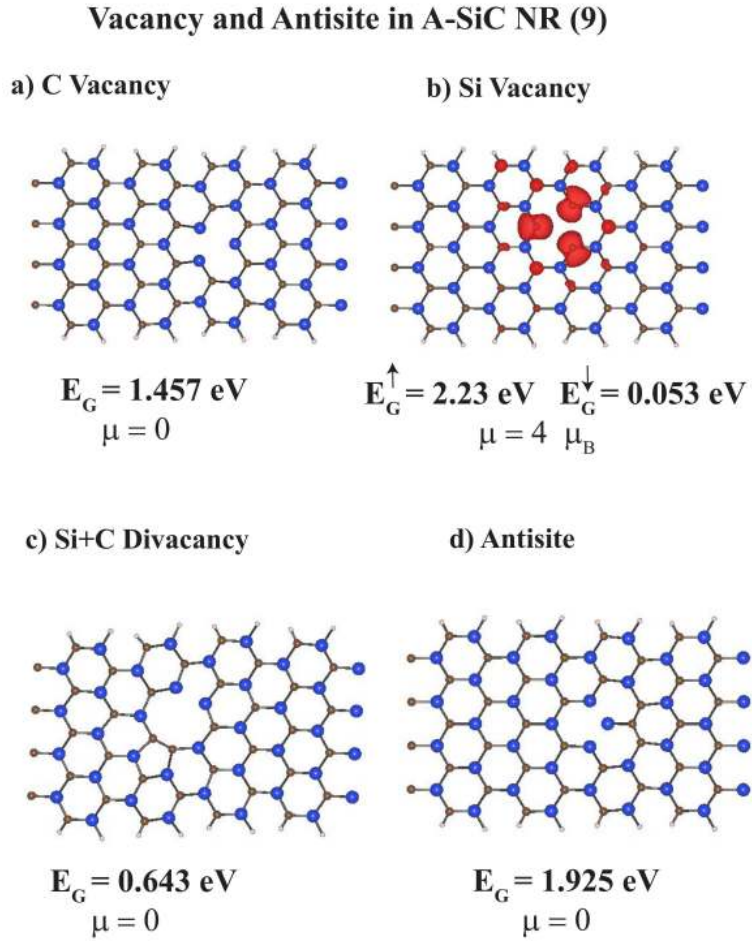


Figure 4.7: Energy band gap and magnetic moment of vacancy defects calculated in a (4×1) supercell of quasi 1D SiC armchair nanoribbon with $N=9$, i.e. A-SiCNR(9). Dangling bonds at both edges are saturated by hydrogen atoms. (a) C vacancy; (b) Si vacancy; (c) Si+C divacancy; (d) C-Si antisite. In (b) the difference of spin up and spin down charges are shown. Large/blue and small/gray balls represent Si and C atoms respectively.

Table 4.4: Final positions, binding energies (E_B), total magnetic moments (μ), final distance of the dopant values (h) for doped planar SiC

Atom	Position	E_B (eV)	μ (μ_B)	h (Å)
Al	TS	1.23	1	1.408
Co	HS	2.099	1	1.392
Fe	HS	1.928	2	1.465
N	TC	3.052	0	0.896
P	BS	1.758	1	1.743
Ti	TS	2.605	2	1.355

of hollow (HS), above the middle of the Si-C bond (BS). To hinder adatom-adatom interaction the distance between adjacent SiC layers are more than 12 Å, and all calculations are carried out spin polarized. Whether the adatoms are bound to the surface are examined by calculating the binding energies of these six different individual atoms in terms of the calculated total energies as $E_B = E_T[\text{SiC} + \text{adatom}] - E_T[\text{SiC}(\text{bare})] - E_T[\text{adatom}]$. We found that all of these adatoms are bound with a binding energy larger than 1 eV. Energy bands and band decomposed charge densities of impurity states of Al adsorbed to SiC monolayer are given in Fig. 4.8. Flat bands indicate that states induced by the adatom are rather localized and hence adatom-adatom interactions are negligible. Optimized adsorption sites, total magnetic moments, distance from the SiC plane are given in Table 4.4 .

4.7.2 Substitution of Si and C by foreign atoms

Here we examined the substitution of single Si or C atoms in 2D SiC honeycomb structure by various foreign (dopant) atoms (B and N substituting C atom; Al, As, Ga, P substituting Si atom). Similar to adatoms, the substitution process is treated within periodically repeating (7×7) supercell. Because of periodic boundary condition the localized states appear as flat bands. The localization of all dopant atoms are attained, except for Ga and Al which required larger supercell. We found that B, N, As and P atoms have a net magnetic moment $\mu =$

Al Adsorption on 7x7 Honeycomb SiC

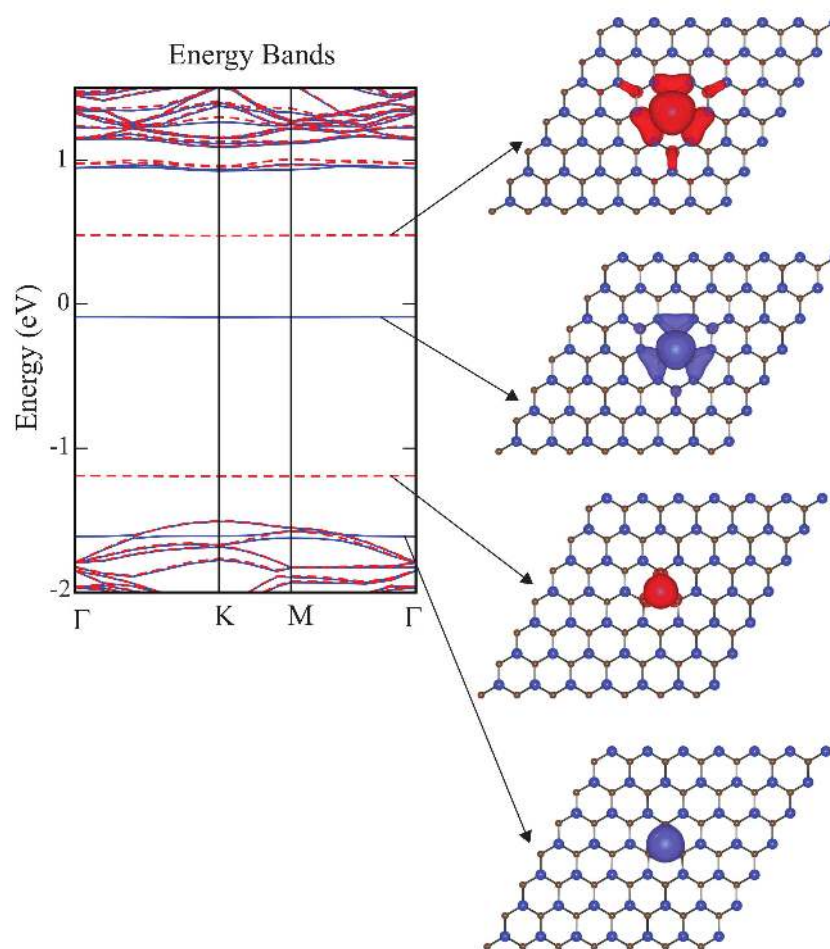


Figure 4.8: Calculated energy bands of single Al atom adsorbed on the top of a Si atom in the (7×7) supercell of 2D SiC honeycomb structure with **solid/blue** and **dashed/red** lines showing **spin up** and **spin down** bands, respectively. Flat bands are associated with the localized states of adsorbed Al atom. Band decomposed charge densities reveal that these states are localized around Al atom.

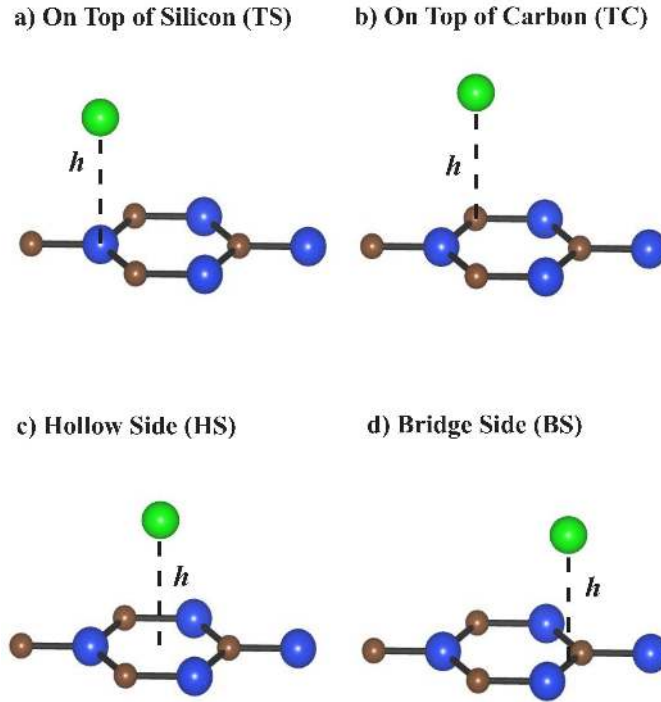


Figure 4.9: Starting positions of adatoms in 7×7 honeycomb SiC

$1 \mu_B$. Ga and Al do not create any magnetic moment as substituted atoms in SiC surface. During structural relaxation, each atom finds its most energetically favorable position in the lattice and there occurs different numbers of dangling electrons. This is the reason why atoms of different valence electrons induce the same magnetic moment. Energy bands and band decomposed charge densities of impurity states of N substituted SiC monolayer are given in Fig. 4.10.

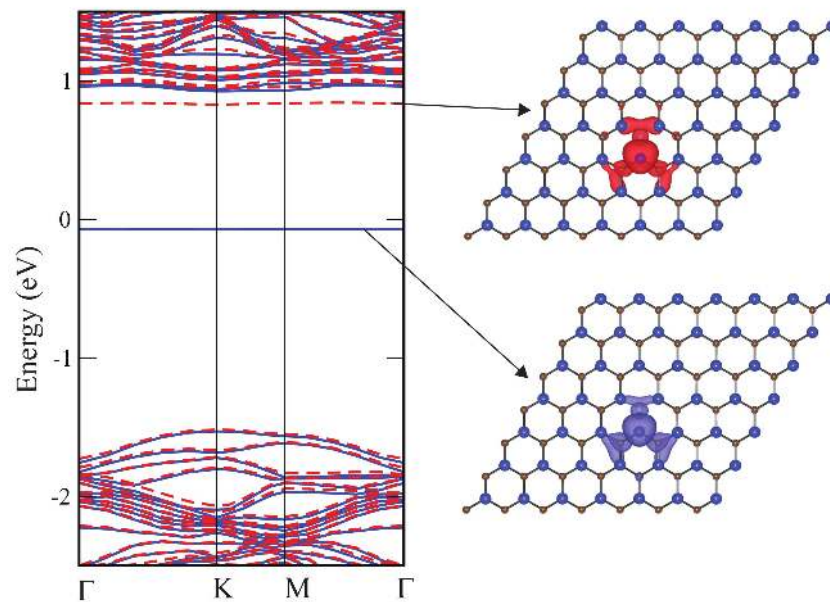
N Substitution in 7x7 Honeycomb SiC

Figure 4.10: Calculated energy bands of single N atom substituting a single C atom in the (7×7) supercell of 2D SiC honeycomb structure with solid/blue and dashed/red lines showing spin up and spin down bands, respectively. Flat bands are associated with the localized states of dopant N atom. Band decomposed charge densities reveal that these states are localized around N atom.

Chapter 5

Conclusions

We present an extensive study on 1D SiC chains, 2D monolayer SiC, 3D bulk SiC and SiC armchair nanoribbons together with the stability analysis of them. The monolayer of SiC is an ionic compound with charge transfer from silicon atoms to carbon atoms. It has hexagonal lattice forming a honeycomb structure and nonmagnetic semiconductor. However, it acquires net magnetic moment through single Si-vacancy. Single C-vacancy, Si+C divacancy and Si-C antisite defects do not give rise to any magnetic moment in the system. Our stability analysis based on phonon frequency calculations indicate that 2D SiC is stable.

Armchair SiC nanoribbons are found to be nonmagnetic semiconductors. The band gaps vary with their widths. They neither show quantum size effect nor have the same family behaviour of graphene. We have both tried bare and H-passivated cases and depicted their bandgap dispersions with respect to widths of nanoribbons.

Adding properties to a bulk material via doping is widely utilised in optical and electronic applications. We have tested similar cases in two dimensional SiC and obtained reasonable results.

After graphene and 2D BN; we have listed electronic, magnetic and structural properties of one additional 2D material. We assume our work becomes more

salient as experimental studies about honeycomb SiC emerge in the following years.

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