

Research Article

Renormalization of The Band - Gap by Isotope in Graphene

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Abstract

Isotope investigation, manufacture, and application is highly variable and is determined by different science and technology areas. The new era of Nano electronics on the graphene basis needs the creation of the semiconducting graphene. Numerous attempts to elaborate the semiconducting graphene creation technology meet several difficulties: firstly, it is quite expensive; secondly it is technical difficult to produce. In the present paper the based-on principle new nuclear semiconducting graphene creation technology is described. The new method is based on the electronic excitations energy renormalization by the strong (nuclear) interaction. Suggested method provides an alternative way to experimentally tune the band - gap of graphene, which would be more efficient and more controllable than other methods that are used to open band - gap in graphene. This method not only opens the isotopical band - gap in graphene but also may throw light on the massless fermion renormalization in graphene.

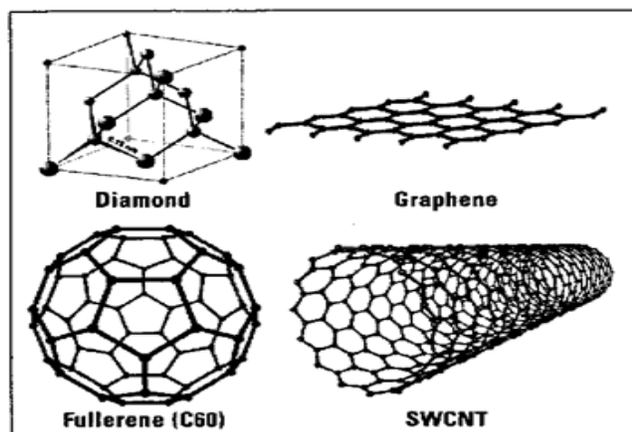
The science of the nuclear, atoms, simple molecules and the science of matter from microstructure to larger scales are well established. A remaining, extremely important [1, 2], size related challenge is at the atomic scale, roughly the dimensional scale between 1 and 10 molecular sizes, where the fundamental properties of materials are determined and can be engineered (see also [3]). This field of science - isotopetronics - is a broad and interdisciplinary field of emerging research and development. Isotopetronics is connected to materials, structures and systems which components, as in nanoscience, exhibit novel and significantly modified physical properties due to their small sizes [3]. The method of the isotope renormalization of the energy of elementary excitations in bulk solid and low - dimensional structures very often used in the last five decades and well documented in the scientific literature (see [1,2] and references quoted

therein). In this paper, current understanding of the band - gap opening in graphene is discussed along with associated experimental and theoretical investigations.

The richness of optical and electronic properties of graphene attracts enormous interest [4]. Carbon atom is built from 6 protons, A neutrons and 6 electrons, where A = 6 or 7, yield the stable isotopes ^{12}C and ^{13}C , respectively, and A =8 characterizes the radioactive isotope ^{14}C . The isotope ^{12}C , with nuclear spin I = 0, is the most common one in nature with 99% of all carbon atoms, whereas only ~ 1% are ^{13}C with nuclear spin I = 1/2. There are only traces of ^{14}C (10^{-12} of all carbon atoms) which β - decays into nitrogen ^{14}N . Although ^{14}C only occurs rarely, it is an important isotope used for historical dating (see, e.g. [5]).

Carbon, one of the most basic elements in nature, still gives a lot of surprises. It is found in many different forms - allotropes - from zero-dimensional fullerene, one-dimensional carbon nanotubes, two-dimensional graphene and graphite, to three-dimensional diamond (Figure 1) and the properties of the various carbon allotropes can vary widely [6]. For instance, diamond is the hardest material, while graphite is one of the softest: diamond is transparent to the visible part of spectrum, while graphite is opaque; diamond is an electrical insulator, while graphite is a conductor. Very important is that all these different properties originate from the same carbon atoms, simply with different arrangements of the atomic structure.

Figure 1: Structure of some representative carbon allotropes.

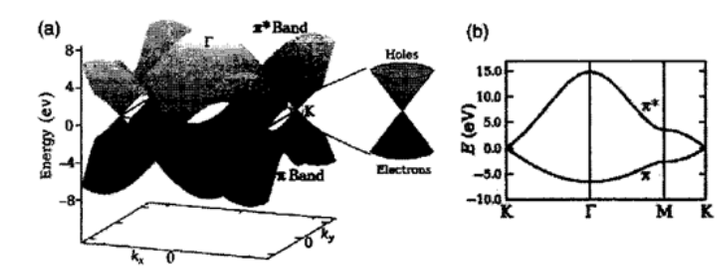


In two-dimensional graphene, carbon atoms are periodically arranged in an infinite honeycomb lattice (Fig.1(a) in [7]). Such an atomic structure is defined by two types of bonds within the sp^2 hybridization. From the four valence orbitals of the carbon atom (the $2s$, $2p_x$, $2p_y$, and $2p_z$ orbitals, where z is the direction perpendicular to the sheet), the (s, p_x, p_y) orbitals combine to form the in-plane σ (bonding or occupied) and σ^* (antibonding or unoccupied) orbitals. Three σ - bonds join a C atom to its three neighbors. They are quite strong, leading to optical - phonon frequencies much higher than observed in diamond (see below). Such orbitals are even with respect to the planar symmetry. The σ bonds are strongly covalent bonds determining the energetic stability and the elastic properties of graphene. The remaining p_z orbital, pointing out of the graphene sheet is odd with respect to the planar symmetry and decoupled from the σ states. From the lateral interaction with neighboring p_z orbitals (called the $pp\pi$ interaction), localized π (bonding) and π^* (antibonding) orbitals are formed [8]. Graphite consists of a stack of many graphene layers.

The unit cell in graphite can be primarily defined using two graphene layers translated from each other by a C-C

distance ($a_{c-c}=1.42 \text{ \AA}$). The three-dimensional structure of graphite is maintained by the weak interlayer van der Waals interaction between π bonds of adjacent layers, which generate a weak but finite out-of-plane delocalization [4]. The bonding and antibonding σ bands are actually strongly separated in energy $> 12 \text{ eV}$ at, and therefore their contribution to electronic properties is commonly disregarded, while the bonding and antibonding π states lie in the vicinity of the Fermi level (Fig. 2). The two remaining π bands completely describe the lowenergy electronic excitations in both graphene and graphite (see [4] and references therein). The bonding π and antibonding π^* orbitals produce valence and conduction bands (Figure 2) which cross at the charge neutrality point (Fermi level of undoped graphene) at vertices of the hexagonal Brillouin zone. Carbon atoms in a graphene plane are located at the vertices of a hexagonal lattice.

Figure 2: Energy dispersion of graphene obtained within the tight - binding approximation. a) Energy dispersion relation for graphene, drawn in the entire region of the Brillouin zone. Since in this approximation to ignore the coupling between the graphene sheets, the bands depend only on k_x and k_y . The π band is completely filled and meets the totally empty π^* band at the K points. Near these points both bands linear dispersion as described in the literature. b) The dispersion along the high symmetry points Γ MK.



This graphene network can be regarded as a triangular Bravais lattice with two atoms per unit cell (A and B). Each A- or B - type atom is surrounded by three atoms of the opposite type. In a simple neighbor model graphene is a semimetal with zero - overlap between valence and conduction bands. The energy dispersion of π electrons in graphene was first derived in 1947 by Wallace [8] within the tight - binding approximation. In this case, the wave function of graphene is a linear combination of Bloch function for sublattice A

$$\Phi_A = \frac{1}{\sqrt{N}} \sum_{\vec{R}_A} e^{i\vec{k}\vec{R}_A} \varphi(\vec{r} - \vec{R}_A), \quad (1) \quad \rightarrow$$

and equilibrium function ϕ_B for the B sublattice. Here N is the number of unit cells, R_A are the position of the atom A and $\varphi(\vec{r} - \vec{R}_A)$ is the $2p_z$ orbital of the atom at \vec{R}_A . The sum runs over all unit cells, i.e. all possible lattice vectors. In the nearest neighbor approximation (every A site has three nearest B sites, and vice versa), the energy eigenvalues can be obtained in a closed form [4,9]

$$\varepsilon(k_x, k_y) = \pm \gamma_0 \left[1 + 4\cos\frac{\sqrt{3}k_x a}{2} \cos\frac{k_y a}{2} + 4\cos^2\frac{k_y a}{2} \right]^{1/2}, \quad (2)$$

where γ_0 is the transfer integral between the nearest neighbors. The energy dispersion of two - dimensional graphene according to this formula is plotted in Fig. 2(a) as a function of the wave vector k. The upper half of the

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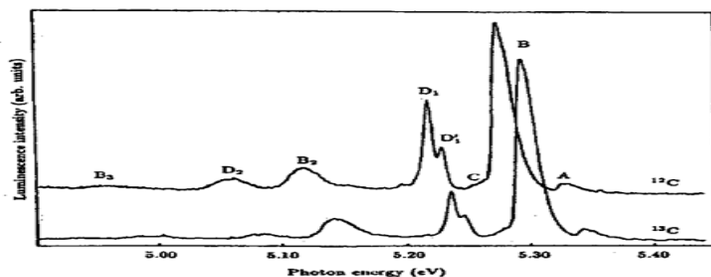
curves is called the π^* or the antibonding band while the lower one is π or the bonding band. The two bands degenerate at the two K points given by the reciprocal space $K = (2\pi/a)(1/3, 1/\sqrt{3})$ and $K = (2\pi/a)(-1/3, 1/\sqrt{3})$ points where the dispersion vanishes (see above).

Basically, graphene has redefined the limits of what a material can do: it boasts record thermal conductivity and the highest current density at room temperature ever measured (a million times that of copper!); it is the strongest material known (a hundred times stronger than steel!) yet is highly mechanically flexible; it is the least permeable material known (not even helium atoms can pass through it!); the best transparent conductive film; the thinnest material known; and the list goes on ...[9]. In the vicinity of K - points (as it can be seen from Fig. 2), the low - energy electron/hole dispersion relation is proportional to momentum, rather than its square. This is analogous to the energy dispersion relation of massless relativistic electrons, so the electrons/holes of graphene are described as Dirac fermions having no mass. In a simple neighbor model graphene is a semimetal with zero - overlap between valence and conduction bands. In order to make graphene a real technology, a special issue must be solved: creating an energy gap at K - points in the Brillouin zone [10]. Different attempts have been made by researchers, such as patterning graphene into nanoribbon [11], forming graphene quantum dots [12-14], making use of multilayer graphene sheets [15, 16] and applying an external electric field [17]. It was shown that the uniaxial strain can open a band - gap in a metallic carbon nanotubes as well as carbon nanoribbon [18].

Further we will briefly discuss dependence of the electronic gap (E_g) as well as phonon states of diamond with its isotopic composition. Fig. 3 compares the edge luminescence for a natural diamond with that for a synthetic (^{13}C) diamond. The peaks labeled A, B and C due, respectively, to the recombination of a free exciton with emission of transverse - acoustic, transverse - optic and longitudinal - optic phonons having wave vector $\pm k_{\min}$ [1,2].

Figure 3 compares the edge luminescence for a natural diamond with that for a synthetic (^{13}C) diamond. The peaks labeled A, B and C due, respectively, to the recombination of a free exciton with emission of transverse - acoustic, transverse - optic and longitudinal - optic phonons having wave vector $\pm k_{\min}$ [1,2].

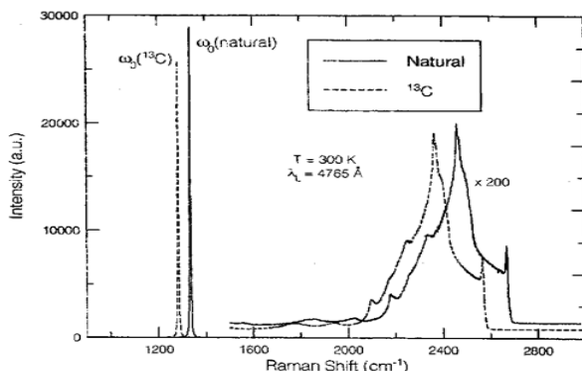
Figure 3: Cathodoluminescence spectra of ^{12}C and ^{13}C at 77 K [1].



As it can be seen from Fig. 3 the band gap of ^{13}C has increased by 13.6 meV. Numerous examples of band gap increase at hard isotope substitution were collected in the papers [1, 2]. The effect of the isotopic ^{12}C to ^{13}C ratio on the first and second - order Raman scattering of light in the diamond has been investigated in [18]. As the ^{13}C content is increased from the natural ratio ($^{12}\text{C}/^{13}\text{C} = (1 - x) / x$, where $x = 0.011$), to the almost pure ^{13}C ($x = 0.987$), the whole spectrum has shifted towards longer wavelengths (see Fig. 4) in good agreement with the expected

$M^{-0.5}$ frequency dependence on the reduced mass M . For an approximately equal mix of the two isotopes, the authors reported that the features seen in the above two - phonon spectra were either broadened or unresolvable. We should stress that the main line in the first - order Raman scattering spectrum of light at $\omega = 1332 \text{ cm}^{-1}$ also shifts to the short-wavelength side on the 52.3 cm^{-1} [1,2].

Figure 4: The Raman spectrum of a natural and a ^{13}C diamond. The spectra show the dominant first - order, Raman - active F_{2g} line and the significantly weaker, quasi - continuous multi - phonon features [19].



Elastic and inelastic light scattering are powerful tools for investigating graphene [19-25]. Raman spectroscopy allows monitoring of doping, defects, disorder, chemical and isotope [1,2] modifications, as well as edges and uniaxial strain. All sp^2 - bonded carbons show common features in their Raman spectra, the so - called G and D peaks (see, e.g. Fig. 8 in [7]), around 1580 and 1360 cm^{-1} (see, e.g. [21,22]). The G peak (see, also below Fig. 6) corresponds to the E_{2g} phonon at the Brillouin zone center ($\Gamma - point$). The D peak is due to the breathing modes of six - atom rings and requires a defect for its activation. It comes from TO phonons around the Brillouin zone K point and it is activated by an intravalley scattering process [21]. The 2D peak is the second order of the D peak. This is a single peak in monolayer graphene, whereas it splits into four bands in bilayer graphene, reflecting the evolution of the band structure [7, 22]. The Raman spectrum of graphene also shows significantly less intensive defect - activated peaks such as the D' peak, which lies at $\sim 1620 \text{ cm}^{-1}$. This is activated by an intravalley process i.e. connecting two points belonging to the same cone around K (see, Fig. 2) [22]. The second order of the D' peak is called 2D' peak. Since 2D and 2D' peaks originate from a Raman scattering process where momentum conservation is obtained by the participation of two phonons with opposite wave vector (q and $-q$), they do not require the presence of defects. Thus, they are always visible in the \vec{R} aman spectrum (see cited papers [19 -25] and references therein).

Graphene is one unique material which shows properties not found in other materials. One of these unique features of graphene is the influence of long-range strains on the electronic properties. The possibility of tuning the dynamics of carriers as well as phonons by appropriately designed strain patterns opens the way for novel applications of graphene, not possible with any other materials (see, e.g. [26] and references therein). At present time we have several reports, which have examined graphene properties under uniaxial deformation [18,25,26,27].

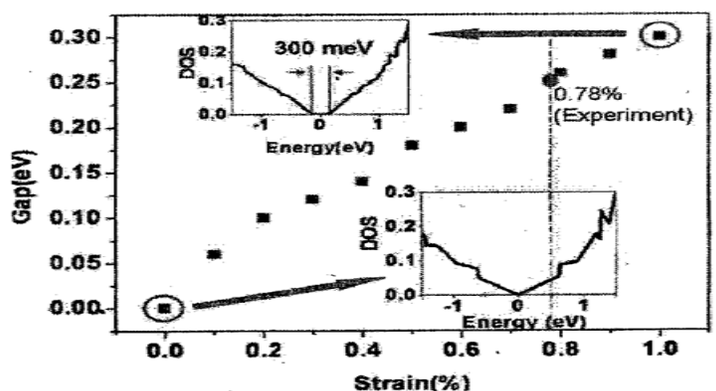
Strain can be very efficiently studied by Raman spectroscopy since this modifies the crystal phonon frequency, depending on the anharmonicity of the interatomic potentials of the atoms. Raman spectra of strained graphene show significant redshifts of 2D and G band (see Table 1) because of the elongated of the carbon - carbon bonds.

Table 1: Red shift of the G and 2D bands in the Raman spectra in graphene monolayers under uniaxial tensile stress.

Ref.	Shift of G (G ⁺ and G ⁻) band cm ⁻¹ /%	Shift of 2D bands cm ⁻¹ /%	E _g , meV
15	14.2	27.8	300
25	5.6; 12.5	21	
27	10.8; 31.7	64	
26 theory			= 500

The authors of the paper [18] have proposed that by applying uniaxial strain on graphene, tunable band - gap at K - point can be realized. First principle calculations predicted a band - gap opening of = 300 meV for graphene under 1% uniaxial tensile strain (Fig. 5). Thus, the strained graphene provides an alternative way to experimentally tune the band - gap of graphene, which would be more efficient and more controllable than other methods (see, above) that are used to open band - gap in graphene.

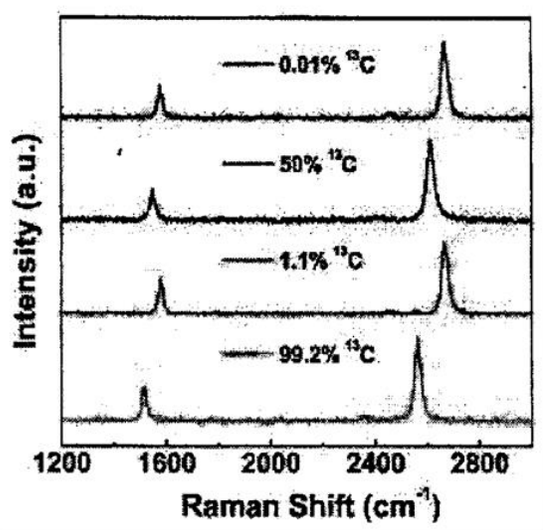
Figure 5: The band - gap of strained graphene with the increase of uniaxial tensile strain on graphene. The magnitude of gap is determined by the gap opening of density of states. The inserts show the calculated density of states of unstrained and 1% tensile strained graphene. The dash line and solid dot indicate the calculated bandgap of graphene under the highest strain (0.78 %) [18].



The method of the isotope renormalization of the energy of elementary excitations in solid very often used in last five decades and well described in the scientific literature (see, for example reviews [1, 2]). At now there is a large list of the paper devoted to investigation of the isotope - mixed graphene [10, 14, 24, 28 - 32]. Chen at al. [23] have reported the first experimental study of the isotope effect on the thermal properties of graphene. The thermal conductivity K, of isotopically pure ¹²C (0.01 of ¹³C) graphene determined was higher than 4000 W/mK (approximately two times more than it in diamond [17]) at the measured temperature T_m ~ 320K, and more than a factor of two higher than the value of K in a graphene sheets composed of a 50% - 50% mixture of ¹²C and ¹³C. Raman spectroscopy transferred to the 285 nm SiO₂/Si wafer was performed under 532 nm laser excitation [23]. The

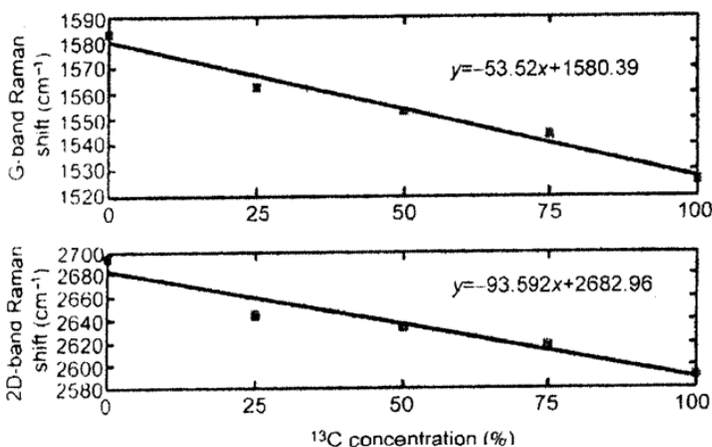
G peak and 2D band positions in Raman spectra of graphene with 0.01%, 1.1% , 50% and 99.2% ^{13}C - isotope are presented in Fig. 6.

Figure 6: Raman spectra of graphene with different isotope concentration at room temperature [23].



Isotope shift of the G and 2D bands in the Raman spectra depicted on the Fig.7 [33].

Figure 7: Peak position of G and 2D bands in Raman spectra as a function of the concentration of ^{13}C [33].



As in the case of isotope - mixed diamond [1, 2] the Brillouin - zone - center optical - phonon frequency c varies with the atomic mass M as $c \sim M^{-1/2}$ making the Raman shift for ^{13}C approximately $(12/13)^{-1/2}$ times smaller than that for ^{12}C . The experimental difference between the lowest 99.2% ^{13}C and the highest 0.01% ^{13}C peak is $\sim 64 \text{ cm}^{-1}$ which is according [23] in agreement with the theory, and attests for the high quality of isotopically modified graphene. By the way we should indicate that in the Raman spectra in diamond (with sp^3 - bond) analogous shift of first - order line in Raman spectrum is equal 52.3 cm^{-1} [34], which is consistent with the isotope mass ratio. Substituting a light isotope (^{12}C , H) with a heavy one increases the interband transition energy in the case $^{12}\text{C}_x^{13}\text{C}_{1-x}$ on 14.7 meV and $\text{LiH}_x\text{D}_{1-x}$ on 103 meV [1]. Taking into account a more soft bond (sp^2 - bond instead sp^3 - bond in diamond) isotope - induced band - gap opening in graphene of some hundreds meV (up to E_g of Si) was predicted in

paper [14]. Such estimation of the value of isotopical band - gap opening in graphene agrees with not only the results of paper [34] but with very small value $C_{44} = 0.5 \cdot 10^{10}$ dyn/cm². Such small value indicates on the strong electron - photon interaction - main reason renormalization of electron excitation energy (for the details, see, e.g. [35]). Very close to isotopically renormalization of electronic excitation energy is the hydrogenation of graphene [7,10]. In last mechanism there is observable band - gap opening in graphene. We should add that use deuterium instead of hydrogen we may increase the value of E_g [1]. Thus, isotope substitution will be very useful method for renormalization of the band - gap in graphene - future semiconducting material. Moreover, this method allows to control not only of the strong nuclear interaction (quantum chromodynamics) but taking it into account at the renormalization of the electromagnetic interaction (quantum electrodynamics) [1]. Adding ¹³C makes magnetic materials isotope out of graphene.

In conclusion lets discuss neutron - electron interaction [36]. There is a common place in Standard Model of modern physics that the strong nuclear force does not act on leptons [37]. Numerous experimental results of the isotope effect study in solids [1-3] show the violation of this strong conclusion. Really, traditionally nuclear - electron (in our case an neutrino - electron) interaction taking into account the solving of Schrödinger equation (see, e.g. [35]) using Born - Oppenheimer (adiabatic) approximation [38]. This approximation results the omission of certain small terms which result from the transformation. As was shown in [35] the eigenvalue (energy of the electronic Schrödinger equation (equation 6 in [35])) depends on the nuclear charges through the Coulomb potential, but it does not include any references to nuclear mass and it is the same for the different isotopes. This result is forcing us to search for new models and mechanisms of nuclear - electron interaction including the results of subatomic physics [39], e.g. hadron - lepton interaction (see also [40]). We should remind that the Standard Model of particle physics (theory of strong interaction) assumes the conservation of lepton and baryon numbers separately, and there no processes that convert quarks to leptons. This means that the Standard Model itself does not prohibit the possibility that the charge of the electron and the proton are slightly different [39]. On the other hand, the neutron comprising quarks can decay into a proton, an electron, and a neutrino. Therefore, a charge asymmetry between the proton and electron would be linked to the charge of the neutron. (see, also [41]). The experimental results of isotope effect evidence the long - range strong interaction [42]. Thus, the use of the method of isotope effect in graphene may throw light on the renormalization of the mass of massless fermion in graphene as well as unification of forces (see, e.g. [43, 44]).

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