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# Quantum Calculation in Prediction the Properties of Single-Walled Carbon Nanotubes

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#### 1. Introduction

Since the first discovery of single-walled carbon Nanotubes (SWCNTs) by Iijima and Bethune in 1993 (Bethune et al., 1993), many applications as molecular components for nanotechnology including conductivity and high-strength composites; energy storage and energy conversion devices; sensors; field emission displays and radiation sources; hydrogen storage media; and nanometer-sized semiconductor devices, probes, and interconnects are known (Ajayan et al., 1994; Saito et al., 1997; deHeer et al., 1995; Collins et al., 1997; Nardelli et al., 1998; Huang et al., 2006). SWCNTs have been considered as the leading candidate for Nan device applications because of their one-dimensional electronic bond structure, molecular size, biocompatibility, controllable property of conducting electrical current and reversible response to biological reagents. Hence SWCNTs make possible bonding to polymers and biological systems such as DNA and carbohydrates. Most SWCNTs have a diameter of close to 1 nanometer, with a tube length that can be many millions of times longer. The average diameter of a SWNT is 1.2 nm (Spires & Brown, 1996). However, Nanotubes can vary in size, and they aren't always perfectly cylindrical. As in Fig. 1, the average bond length and carbon separation values for the hexagonal lattice were shown. The carbon bond length of 1.42 Å was measured by Spires and Brown in 1996 (Spires & Brown, 1996) and later confirmed by Wilder et al. in 1998 (Wilder et al., 1998).

The structure of a SWNT can be formed by the rolling of a single layer of sp<sup>2</sup> carbon, called a graphene layer, into a seamless hollow cylindrical tube with Nan scale dimensions of 1-1.5 nm. The length is usually in the order of microns to centimeters. Besides their unique physical properties (elasticity, tensile strength, stiffness, and deformation), Nano tubes exhibit varying electrical properties (depending on the direction that the graphite structure spirals around the tube (quantified by the "Chiral vector"), and other factors, such as doping), and can be superconductor, conductor (metallic), semiconductor or, insulator. The band structure can even be further manipulated, by introducing defects into a tube. Single-

walled nanotubes exhibit electric properties that are not shared by the multi-walled carbon nanotube (MWNT) variants. In particular, their band gap can vary from zero to about 2 eV and their electrical conductivity can show metallic or semiconducting behavior, whereas MWNTs are zero-gap metals. The C-C tight bonding overlap energy is in the order of 2.5 eV. Wilder et al. estimated it to be between 2.6 eV - 2.8 eV (Wilder et al., 1998) while at the same time, Odom et al. estimated it to be 2.45 eV (Odom et al., 1998). Multi-walled carbon nanotubes have a layer of carbon shells with differing physics that can all potentially interact. It is shown that only the outer shell of MWCNTs contributes to electrical transport, and so only small diameter MWCNTs could be used to make transistor devices. SWCNTs are the most likely candidate for miniaturizing electronics beyond the micro electromechanical scale currently used in electronics. As this field continues to expand and grow, materials technology will produce products, components and systems that are smaller, smarter, multi-functional, environmentally compatible, more survivable, and customizable. These products will not only contribute to the growing revolutions of information and biology, but will also significantly impact manufacturing, logistics, and our culture as a whole. The development of scanning probe techniques has allowed not only the microscopy of surfaces with atomic resolution, but also the manipulation of atoms and molecules on surfaces, and many analytical techniques have been developed to allow detailed characterization of materials and structures on the atomic level with unprecedented accuracy. The utilization of materials with nanometer-sized structures will lead to innovative products which are smaller, smarter, and more multi-functional. Therefore, understanding of fundamental properties of structures at the nano scale with the aid of computational models is important to design the specific material properties.

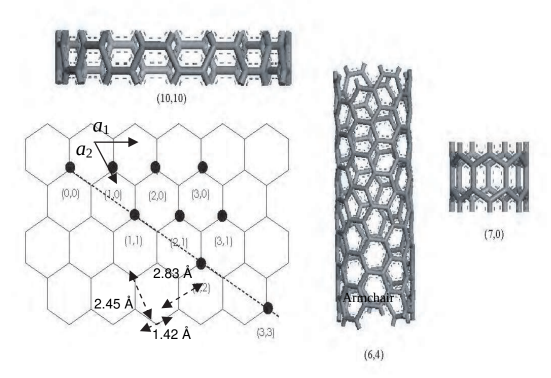


Fig. 1. The geometrical structure of SWCNT

Recently, theoretical and experimental work have predicted that the infinity length SWCNTs are Pi-bonded aromatic molecules that the electrical properties depending upon the tubular diameter and helical angle (Zhou et al., 2004; Baron et al., 2005). SWCNTs can be chiral or nonchiral, again depending on the way of the rolling up vector. As a graphene sheet was rolled in many ways in horizontal, vertical, and diagonal direction represent as arrow vectors,  $\vec{a}$ , as in Fig. 1, the different types of carbon nanotubes were produced. The three main types are armchair, zig-zag, and chiral nanotube. The geometrical and electronic structure of SWCNT can be described by a chiral vector, the angle between the axis of its hexagonal pattern and the axis of the tube which is presented by a pair of indices ( $n_1$ , $n_2$ ) called the chiral vector. The integers  $n_1$  and  $n_2$  denote the number of unit vectors along two directions in the honeycomb crystal lattice of graphene. When the indices are ( $n_1$ ,0) called zig-zag, ( $n_1$ ,  $n_1$ ) called armchair, and ( $n_1$ ,  $n_2$ ) where  $n_1 \neq 0$  and  $n_2 \neq 0$  known as chiral SWCNT. For (2  $n_1 + n_2$ )/3 = integer, SWCNTs are metallic and others are semiconductors (Saito et al.,

1992a, 1992b). For large diameter SWCNTs defined by 
$$d = \frac{\sqrt{3(n_1^2 + n_2^2 + n_1 n_2)}}{\pi} a_{c-c}$$
, where  $a_{c-c}$ 

is the distance between neighboring carbon atoms in the flat sheet, armchair SWCNTs are always metallic which is good for nanotechnology application. A zigzag carbon nanotube  $(n_1, 0)$ , is a semiconductor when  $n_1/3 \neq \text{integer}$ . Such semiconductor zigzag carbon nanotubes have the ability to become base of many nanoelectronic devices and transistors. Although, scientific efforts focused on the electrostatics properties and commercial applications of these materials (Ouyango et al., 2002; Kane & Mele, 1997; Hartschuh et al. 2005), there have been no experimental structural data sufficiently accurate for the identification of the chirality indices of SWCNTs, especially for the kind of smaller diameter nanotubes. In all experimental methods for the identification commonly utilized so far is Raman spectroscopy and phonon dispersion. Phonon dispersion relations in one dimension of this system have been studied by using zonefolding along one direction of Brillouin zone considering the tube symmetry (Eklund et al., 1995). The tight binding electronic band structure and the reverse of the diameter (1/d) dependence of the frequency of the radial breathing mode (RBM) were employed (Jorio et al., 2001; Bachilo et al, 2002; Pfeiffer et al. 2003; Kurti et al., 2004; Maultizsch et al., 2005). The size and chirality of the carbon nanotubes were typical determined from the SWCNT Raman energy spectra of a peak around 150-300 cm<sup>-1</sup>, due to the radial breathing mode (Maultizsch et al., 2005; Jorio et al., 2005). Further Raman studies of SWCNT modified by various reactions e.g. oxidation reactions, ozonolysis, fluorination, residues modification (Srano et al., 2003; Umek et al., 2003; Peng et al., 2003; Bahr et al., 2001; Holzinger et al., 2003; Mickelson et al., 1998; Cai et al., 2002; Banerjee & Wong, 2002; Herrera & Resasco, 2003; Martinez et al., 2003) have revealed that covalent functionalization mainly affects the intensity of the Raman bands. Characterization of nanotube in adsorption gas has been studied by Monte Carlo and Langevin Dynamic Simulation (Monajjemi et al., 2008b). It is also important to investigate the effects of diameter on a SWCNT structure how the diameter depends on geometrical parameters such as the C-C bond lengths and some of the dihedral angles of SWCNTs.

For a better understanding of the physical and electronic properties of SWCNT, a challenging task in theoretical calculation is needed to specify the material properties because of the large size of the SWCNTs and their complicated (and size-dependent) electronic structure. Quantum calculation in prediction the properties of single-walled carbon nanotubes (SWCNTs) will be discussed.

#### 2. Vibrational mode of SWCNT

Normal mode analysis has become one of the standard techniques in the study of the dynamics of nanotubes. It is primarily used for identifying and characterizing the slowest motions in a poly system, which are inaccessible by other methods. This text explains what normal mode analysis is and what one can do with it without going beyond its limit of validity. By definition, normal mode analysis is the study of harmonic potential wells by analytic means. The first section of this study will therefore deal with potential wells and harmonic approximations. This study is about normal mode approaches to different physical situations, and it discusses how useful information can be extracted from normal modes. Normalmode coordinates are obtained by a linear combination of Cartesian coordinates. Thus, there are no couplings in the kinetic part; that is, they diagonalize the kinetic energy as well the quadratic part of the potential energy operator. They include simultaneous motion of all atoms during the vibration, which leads to a natural description of molecular vibrations. Therefore, they are good candidates for representation of the molecular Hamiltonian. Since a transformation between different sets of coordinates is possible, the anharmonic terms can be calculated in one representation, and then transformed into another one.

#### 2.1 Symmetry of SWCNT

Because a single carbon nanotube may be thought of as a graphene sheet rolled up to form a tube, carbon nanotubes should be expected to have many properties derived from the energy bands and lattice dynamics of graphite. For the very smallest tubule diameters, however, one might anticipate new effects stemming from the curvature of the tube wall and the closing of the graphene sheet into a cylinder. A method for identifying the Raman modes of single-wall carbon nanotubes (SWNT) based on the symmetry of the vibration modes has been widely used. The Raman intensity of each vibration mode varies with polarization direction, and the relationship can be expressed as analytical functions. Each Raman-active mode of SWNT can be distinguished from the group theory principle. The symmetry properties of periodic lattices of carbon nanotubes and the symmetry operations of chiral and achiral nanotubes (Damnjanović et al., 1999; Damnjanović et al., 2001; Alon, 2001, 2003) are usually described in terms of the group of the wavevector (Dresselhaus et al., 2006). However, since nanotubes can be viewed as quasi-1D systems, the line groups approach by Damnjanović et al. is suited to describe nanotube properties (Damnjanović et al., 1999).

As described earlier, the properties of nanotubes are determined by their diameter and chiral angle, both of which depend on  $n_1$  and  $n_2$ . Typically, SWCNT is presented by a pair of integers  $(n_1, n_2)$ . Its geometrical structure as shown in diagram can be represented in term of a chiral vector  $\vec{C}$  on a two-dimensional sp²-carbon sheet where  $\vec{C} = n_1 \vec{a}_1 + n_2 \vec{a}_2$  with integer  $n_1$  and  $n_2$ . Here,  $\vec{a}_1$  and  $\vec{a}_2$  represent the unit vectors of the hexagonal graphene lattice. This sheet is then rolled up to a cylinder so that  $\vec{C}$  becomes the circumference of the tube. The direction of the nanotube axis is naturally perpendicular to  $\vec{C}$ . The diameter, d, is simply the length of the chiral vector divided by  $\frac{1}{4}$ , and  $d = \left(\sqrt{3}/\pi\right)a_{c-c}(n_1^2 + n_2^2 + n_1n_2)^{1/2}$ , where  $a_{c-c}$  is the distance between neighbouring carbon atoms in the flat sheet. In turn, the chiral angle  $(\theta)$  is given by  $\tan^{-1}\left(\sqrt{3n/(2n_2+n_1)}\right)$ .

The translational period, a, is the shortest possible lattice vector along z direction. The translatory unit cell of a nanotube is a cylinder with a length in tube axis direction equal to the magnitude of the translation vector  $\vec{T}$  as shown in Fig. 1 which can be calculated as following equation:

$$a = -\frac{2n_2 + n_1}{nR}a_1 + \frac{2n_1 + n_2}{nR}a_2$$

with

$$a = |a| = \frac{\sqrt{3(n_1^2 + n_2^2 + n_1 n_2)}}{nR} a_0$$

where n is the greatest common divisor of  $n_1$  and  $n_2$ ,

if 
$$(n1 - n2)/3n \neq \text{integer}$$
, then R = 1

if 
$$(n1 - n2)/3n = integer$$
, then R = 3

and  $\vec{a}_1$  and  $\vec{a}_2$  form an angle of 60° and their length is  $|a_1| = |a_2| = a_0 = 2.461 \,\text{Å}$  Since the translational period, a, depends inversely on n and R the translation periodicity and thus the number of carbon atoms varies strongly for tubes with similar diameter. The number of graphene cells in the nanotube unit cell  $(n_c)$  obtained from:

$$n_c = 2q = 4\frac{n_1^2 + n_2^2 + n_1 n_2}{nR}$$

The groups of infinite line L are products L = ZP, where P is a point group and Z is the group of translations (screw axis, pure translations, and glide planes). Applying the above symmetry formulation to armchair ( $n_1 = n_2$ ) and zigzag ( $n_2 = 0$ ) nanotubes, such nanotubes with no caps have a isogonal point groups given by q (the number of graphene cells in the unit cell of the nanotubes) namely,  $D_{nd}$  when n is odd,  $D_{nh}$  when n is even, or Dqh = D2nh for achiral and Dq for chiral tubes. Whether the symmetry groups for armchair and zigzag tubules are taken to be  $D_{nd}$  or  $D_{nh}$ , the calculated vibrational frequencies will be the same; the symmetry assignments for these modes, however, will be different. It is, thus, expected that modes that are Raman or IR-active under  $D_{nd}$  (or  $D_{nh}$ ) but are optically under D2nh will only show a weak activity resulting from the fact that the existence of caps lowers the symmetry that would exist for a nanotube of infinite length.

#### 2.2 Active modes of Raman and IR

The phonon symmetries are found by decomposing the dynamical representation into its irreducible representations using symmetries of carbon and other nanotubes studied for line groups (Damnjanović et al., 1999). One direct set up of the dynamical representation from the atomic and vector representation is to use factor group analysis. A representation can be decomposed into the sum of its irreducible representations by the following formula

$$f_{\alpha} = \frac{1}{g} \sum_{G} \chi^{(\alpha)}(G) * \chi^{(\Gamma_{DG})}(G)$$

where  $f \alpha \Box$  is the appearance frequency of the irreducible representation  $\alpha \Box$ , g is the order of the symmetry group; the sum is over all symmetry operations G.

The Raman  $\Gamma_R$  and infrared active  $\Gamma_{IR}$  vibrations transform according to the representation of the second rank tensor and the vector representation, respectively (Damnjanović et al., 1983)

$$\Gamma_{R}$$
 =  $[\Gamma_{vec} \otimes \Gamma_{vec}]$  =  $A_{1g} \oplus E_{1g} \oplus E_{2g} (\oplus A_{2g})$   
 $\Gamma_{IR}$  =  $\Gamma_{vec}$  =  $A_{2u} \oplus E_{1u}$ 

According to the symmetries of Raman-active modes (Pelletier, 1999) for the armchair carbon nanotube with the chair vector (n1, n2), the point group for this kind nanotube belongs to Dnh when n is even and its Raman-active modes are denoted by A1g + E1g + E2g. Three flavors of modes are longitudinal, transversal radial (orthogonal to tube surface) and transversal axial (parallel to tube surface). Satio et al. (Satio et al., 1998) pointed out that the low frequency A1g mode is a radial breathing mode and two high frequency is belong to Eg modes, E1g and E2g. E mode has the same displacement pattern with additional standing wave on the circumference.

#### 2.3 Projection operators

The zigzag single-walled carbon nanotubes (SWCNTs) with (3,0), (4,0), and (5,0) structure were built using the tool in HyperChem7.0. The symmetries of the nanotube are D3d, D4d, and D5d respectively. Four different systems were studied in this work as follow: (1) gasphase SWCNT, (2) SWCNT with 23 water molecules in the  $a \times b \times c$  box, (3) SWCNT with 23 methanol molecules in the  $a \times b \times c$  box, and (4) SWCNT with mixed solvent of water and methanol molecules in the  $a \times b \times c$  box. Energy minima of systems (2) – (4) were carried out by Metropolis Monte carlo (MC) calculation which generate random configurations in regions of space that make the important contributions to the calculation of thermodynamic averages. Then the ab initio and semiemperical with AM1 were used to optimize the structure of the nanotubes. All the normal mode frequencies and IR intensity were calculated using the optimized structures.

To find a function or the displacement pattern of eigenvectors transforming as a particular irreducible representation, the projection operators in group theory have been applied. Consider an arbitrary function F. This function can, in general, be expanded into several irreducible representations  $F = \sum_{\alpha} \sum_{n} c_{\alpha}^{n} \zeta_{\alpha}^{n}$  where  $\alpha$  labels the irreducible representations,

 $c_{\alpha}^{n}$  are the coefficients of the expansion, and the  $\zeta_{\alpha}^{n}$  are functions transforming according to the representation  $\alpha$ . A projection operator defined by  $P_{l(n)}^{(\beta)} = \frac{d_{(\beta)}}{g} \sum_{G} D_{ln}^{(\beta)}(G) * (G)$  applied

to F picks out the symmetry adapted function  $\zeta_l^{(\beta)}$ . In equation  $d_\beta$  is the degeneracy of the irreducible representation  $\beta$ , g the order of the symmetry group, G are the symmetry operations, and  $D_{ln}^{(\beta)} \ln i$  is the lnth element of the representation matrix  $D^{(\beta)}$ . From a given function, and its irreducible presentation, functions can be generating if that function has a "component" or a "non-zero projection" along the irreducible presentation of interest. This explains the name of "projector". As an example, if there is an orthonormal set Li of the function  $\phi_1^{i_1}, \phi_2^{i_2}, ..., \phi_{l_{Li}}^{i_{Li}}$  which is used to form the  $i^{th}$  irreducible representation of a group by order h, for each operator, R, in the group, by definition we can have:

$$\mathbf{R}\phi^{i}_{t} = \Sigma s\phi^{i}_{s} \Gamma(R)^{i}_{st} \tag{1}$$

By producting (1) in  $[\Gamma(R)^{i_{s't'}}]^*$  and summing all over the symmetrical functions in the group we will have:

$$\sum_{R} [\Gamma(R)^{i}_{s't'}]^* R \varphi^{i}_{t} = \sum_{R} \sum_{S} \Gamma \varphi^{i}_{S} \Gamma(R)^{i}_{St} \Gamma(R)^{i}_{s't'}]^*$$
(2)

Considering  $\varphi_s^i$  are functions independent from R, the right side of (2) can be written as:  $\Sigma_s \varphi_s^i \Sigma_R \Gamma(R)^i_{st} [\Gamma(R)^i_{s't'}]^*$  So we have a series of Li terms and each of them are equal to a production of  $\varphi_s^i$  and a coefficient. These coefficients are following the orthogonality rule:

$$\Sigma_{R}\Gamma(R)^{i}_{st} \left[\Gamma(R)^{i}_{s't'}\right]^{*} = h/(L_{i}L_{j})^{1/2} \delta_{ij} \delta_{ss'} \delta_{tt'}$$
(3)

By use of the eq.(3), the eq.(2) is simplified as follows:

$$\sum_{R} \Gamma(R) i_{s't'} R \phi^{i}_{t} = (h/L_{i}) \phi^{i}_{s'} \delta_{ii} \delta_{tt'}$$

$$\tag{4}$$

Now, by introducing

$$P^{j}_{s't'} = L_{j}/h \sum_{R} \Gamma(R)^{i}_{st} \left[ \Gamma(R)^{i}_{s't'} \right]^{*} R$$

$$(5)$$

The eq.(4) gives the following form:

$$P^{j}_{s't'}\phi^{i}_{t} = \phi^{i}_{s'} \delta_{ij} \delta_{tt'}$$

$$\tag{6}$$

The  $P_{s't'}$  is call projection operator. The application of this operator on each  $\varphi_t^i$  is non-zero only when this function or some of its terms is a function of  $\varphi_{s'}^i$ . One of the most important application of this operator is projecting function  $\varphi_t^i$  from any function  $\varphi_t^i$ . In other words

$$P^{j}_{\mathbf{t''t}}\phi^{i}_{t} = \phi^{i}_{t'} \,\delta_{ij} \,\delta_{tt'} \tag{7}$$

By use of the projection operator on the base of  $L_j$  diagonal elements of a matrix, we can have some  $\varphi^{i_t}$  functions, which are the bases for the  $j^{th}$  irreducible presentation (Wilson, et al., 1955)

#### 2.4 The relation between projection and transfer operators

Assume that  $\Gamma_k$  (p)  $_{ij}$  is the  $ij_{th}$  element of the matrix which shows the  $p^{th}$  operator (Op) in k the  $k_{th}$  irreducible presentation. By this assumption the operator  $O_{k,ij}$  is defined as follows

$$O_{k,ij} = L_k / h \Sigma_p \Gamma_k (p)^*_{ij} Op$$
 (8)

Where h is group order and  $L_k$  is the presentation dimension. If i = j these operators called Projection operators,  $P_{k,ii}$ , in other words:

$$P_{k,ij} = O_{k,ii} \tag{9}$$

The non-diagonalized operators are called Transfer operators or shift operators,

$$T_{k,ij} = O_{k,ij}, i \neq j$$
 (10)

In one-dimensional presentations  $P_{k,ij}$  and  $O_{k,ii}$  are the same and we have no  $T_{k,ij}$ . With use of the above definitions, making the irreducible basis becomes possible in the following way:

At first the point group of the molecule is determined. Then the character of the system ( $\Gamma_{angles}$  or  $\Gamma_{bonding}$ ) is calculated. By use of the standard reduce formulation these characters can be reduced to give the irreducible presentations:

$$n_{\Gamma} = 1/h \Sigma_g n_g \chi_R \chi_{\Gamma}$$
 (11)

Where h is the order of the group, ng is the number of the symmetry operation in the class of g,  $\chi_R$  is the character of reducible presentation and  $\chi_\Gamma$  is the character of irreducible presentation for the symmetric operations of class g. In this part there is a note about the reducing the  $C_{\infty v}$  and  $D_{\infty h}$  point groups. The method of reduce is a different from the normal method of reduce. For more information see from the references (Cotton, 1971; Schafer & Cyvrin, 1971; Strommen & Lippincott, 1972; Alvarino, 1978; Flurry, 1979; Strommen, 1979). At the next step, the interested function is written by use of the projection operator. A set of the results gives the internal coordinate system for a given point group. There are several examples to illustrate this procedure in Table 1. The geometry and electrical properties of nanotube are very sensitive to dielectric constants. The normal modes also will be changed in the high dielectric constants. With the calculation of the normal modes using the U Matrix it is possible to get the F Matrix from the multiplication of frequency to the U Matrix. Solving the determination of F Matrix versus dielectric can be useful for understanding of the electrical behavior of nanotubes in the quantitative structure activity relationship (QSAR) studies. With use of the resulting coordinate system, the U Matrix (UMAT) can be written easily. These are matrices which perform the linear transformations on the internal coordinates sets (Alvarino & Chammoro, 1980).

#### 2.5 Linear combination of primitive's harmonic vibrations and UMAT

The molecules and their internal coordinates of D4d have been given in Fig. 2. By following the above steps a complete set of the linear combinations and their normalization coefficients are achieved. The irreducible representations of the symmetry group are given by A and B. These data are given in Table 1. We use the application of projection operator method in finding the coordinate system, and by using it the U matrix is written and finally the frequencies and distributions of peak position are achieved. The (3, 0), (4, 0), (5, 0) zigzag nanotubes were investigated. They have 66, 138, and 174 normal modes, respectively.

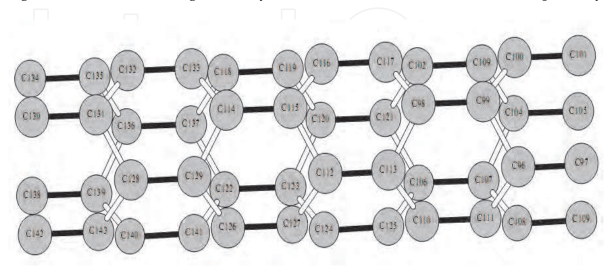


Fig. 2. The structure of (4, 0) nanotube in  $D_{4d}$  point group (Lee et al., 2009)

Ć,	H	£2	п.	и,	irreduced presents- tion	Primis 1198s	Linear comminents of primitives based on their symmetric $A_1, A_2, B_1, B_2$
r,	Test.	-	-	(Meg.)	3A, B2	Sales Control	$\hat{S}_{1}(A_{1}) = \frac{1}{\sqrt{2}}(\hat{R}_{1} - \hat{R}_{2}), \hat{S}_{2}(\hat{R}_{3}) = \frac{1}{\sqrt{3}}(\hat{R}_{1} - \hat{R}_{2}), \hat{S}_{2}(A_{1}) = \hat{R}_{3}.$
T <sub>2</sub>	3	-	-	3	2.4. B)	3	$S_1(A_1) = \frac{1}{\sqrt{2}} (B_1 - B_2), S_2(B_2) = \frac{1}{\sqrt{2}} (B_2 - B_2), S_2(A_1) = B_2$
Fg	3	-	-	3	24, 8 <sub>2</sub>	3	$S_1(A_1) = \frac{1}{\sqrt{2}} (R_{14} - R_{15}), S_2(R_2) = \frac{1}{\sqrt{2}} (R_{14} - R_{15}), S_2(A_1) = R_{25}$
T.	7	-	1	3	24 <sub>1</sub>	3	$E_{10}(A_1) = \frac{1}{\sqrt{3}}(B_{20} + B_{20}), S_{10}(B_2) = \frac{1}{\sqrt{2}}(B_{20} - B_{20}), S_{10}(A_1) = S_{10}$
f,	- 100		-		24 <sub>1</sub>		$R_{13}(A_1) = \frac{1}{\sqrt{2}} \left( R_{22} + R_{23} \right),  R_{24}(B_2) = \frac{1}{\sqrt{2}} \left( R_{23} - R_{24} \right),  R_{23}(A_1) = R_{23}$
Ţ,	· feet.	+	+	2140	24	T.	$S_{1g}(A_1) = \frac{1}{\sqrt{2}}(B_{2g} + B_{3g}), S_{1g}(B_{2}) = \frac{1}{\sqrt{2}}(B_{2g} - B_{3g}), S_{2g}(A_1) = B_{4g}$
T,	18	2	5)	2	THE SECOND SECON	18	$\begin{split} &S_{12}(A_1) = \frac{1}{2} \left( R_1 + R_2 + R_1 + R_2 \right), S_{22}(A_2) = \frac{1}{2} \left( R_1 + R_2 - R_1 + R_2 \right), \\ &S_{22}^* \left( A_1 \right) = \frac{1}{2} \left( R_1 + R_2 - R_1 - R_2 \right), S_{22}(A_2) = \frac{1}{2} \left( R_3 - R_1 + R_2 - R_2 \right), \\ &S_{22}(R_1) = \frac{1}{\sqrt{2}} \left( R_4 + R_{32} \right), S_{22}(A_3) = \frac{1}{\sqrt{2}} \left( R_4 - R_{14} \right), S_{22}(A_1) = \frac{1}{\sqrt{2}} \left( R_{41} + R_{32} \right), \\ &S_{22}(B_1) = \frac{1}{\sqrt{2}} \left( R_{41} - R_{22} \right), S_{23}(A_1) = \frac{1}{\sqrt{2}} \left( R_{21} - R_{22} \right), S_{22}(B_2) = \frac{1}{\sqrt{2}} \left( R_{21} - R_{32} \right), \\ &S_{22}(A_1) = \frac{1}{2} \left( R_{22} - R_{31} + R_{31} + R_{32} \right), S_{22}(B_1) = \frac{1}{2} \left( R_{22} - R_{32} - R_{32} - R_{32} \right), \\ &S_{21}^* \left( R_1 \right) = \frac{1}{2} \left( R_{22} - R_{31} + R_{31} + R_{32} \right), S_{22}(B_1) = \frac{1}{2} \left( R_{22} - R_{32} - R_{32} - R_{32} \right), \\ &S_{21}^* \left( R_1 \right) = \frac{1}{2} \left( R_{32} - R_{31} - R_{32} - R_{32} \right), S_{32}(A_2) = \frac{1}{2} \left( R_{22} - R_{31} - R_{32} - R_{32} \right). \end{split}$
F,	14	2	40	3	14, 16, 16,	14	$\begin{split} S_{11}(A_1) &= \frac{1}{2} \left( R_{11} - R_1 - R_1 - R_2 - R_3 \right) S_{22}(A_2) = \frac{1}{2} \left( R_{11} + R_2 - R_4 - R_2 \right), \\ S_{11}^{**}(A_2) &= \frac{1}{2} \left( R_1 + R_2 - R_{11} - R_3 \right), S_{22}(B_1) = \frac{1}{2} \left( R_1 - R_{11} - R_2 - R_3 \right), \\ S_{12}(A_1) &= \frac{1}{\sqrt{2}} \left( R_4 - R_{211} \right), S_{22}(B_2) = \frac{1}{\sqrt{2}} \left( R_4 - R_{22} \right), S_{32}(A_1) = \frac{1}{\sqrt{2}} \left( R_{43} - R_{24} \right), \\ S_{22}(B_2) &= \frac{1}{\sqrt{2}} \left( R_{21} - R_{21} \right), S_{32}(A_1) = \frac{1}{\sqrt{2}} \left( R_{21} - R_{22} \right), S_{32}(B_2) = \frac{1}{\sqrt{2}} \left( R_{21} - R_{22} \right), \\ S_{22}(A_1) &= \frac{1}{2} \left( R_{22} + R_{24} - R_{23} + R_{23} - R_{23} \right), S_{22}(B_1) = \frac{1}{2} \left( R_{33} - R_{31} + R_{12} - R_{32} \right), \\ S_{21}^{**}(B_1) &= \frac{1}{2} \left( R_{31} + R_{34} - R_{34} - R_{33} - R_{34} \right), S_{22}(A_2) = \frac{1}{2} \left( R_{33} - R_{31} + R_{32} - R_{34} \right). \end{split}$

r,	12	000		19	5.4, 3.4, 3.8, 3.8,	14	$\begin{split} S_{43}(A_1) &= \frac{1}{2} \left( B_{21} + R_{13} + B_{13} + R_{13} \right), \ S_{44}(A_2) = \frac{1}{2} \left( B_{21} - R_{23} + R_{13} - R_{13} \right), \\ S_{44}^+(A_2) &= \frac{1}{2} \left( B_{23} + B_{13} - R_{23} - R_{13} \right), \ S_{43}(B_2) = \frac{1}{2} \left( R_{13} - R_{21} + R_{13} - R_{23} \right), \\ S_{44}(A_1) &= \frac{1}{\sqrt{2}} \left( R_{24} + B_{23} \right), \ S_{47}(B_2) = \frac{1}{\sqrt{2}} \left( R_{12} - R_{23} \right), \ S_{43}(A_2) = \frac{1}{\sqrt{2}} \left( R_{44} + R_{21} \right), \\ S_{44}(B_2) &= \frac{1}{\sqrt{2}} \left( R_{44} - R_{21} \right), \ S_{53}(A_2) = \frac{1}{\sqrt{2}} \left( R_{43} + R_{24} \right), \ S_{53}(B_2) = \frac{1}{\sqrt{2}} \left( R_{31} - R_{43} - R_{43} \right), \\ S_{53}(A_3) &= \frac{1}{2} \left( R_{30} + R_{41} + R_{41} + R_{42} + R_{43} \right), \ S_{54}(A_2) = \frac{1}{2} \left( R_{30} - R_{41} + R_{41} - R_{43} \right), \\ S_{55}^+(B_1) &= \frac{1}{2} \left( R_{41} + R_{41} - R_{42} - R_{43} \right), \ S_{54}(A_2) = \frac{1}{2} \left( R_{50} - R_{41} - R_{43} - R_{43} \right). \end{split}$
Γ19	19		(8)	12	5.4, 3.4, 3.8, 3.8,	14	$\begin{split} S_{23}(A_1) &= \frac{1}{2} \left( R_{13} + R_{13} + R_{13} + R_{12} \right), \ S_{23}(A_2) = \frac{1}{2} \left( R_{12} - R_{13} + R_{13} - R_{12} \right), \\ S_{23}^+(A_2) &= \frac{1}{2} \left( R_{12} + R_{12} - R_{13} - R_{13} \right), \ S_{23}(B_1) = \frac{1}{2} \left( R_{13} - R_{13} + R_{12} - R_{13} \right), \\ S_{23}(A_1) &= \frac{1}{\sqrt{2}} \left( R_{23} + R_{23} \right), \ S_{23}(B_2) = \frac{1}{\sqrt{2}} \left( R_{13} - R_{23} \right), \ S_{23}(A_1) = \frac{1}{\sqrt{2}} \left( R_{31} + R_{43} \right), \\ S_{31}(B_2) &= \frac{1}{\sqrt{2}} \left( R_{31} - R_{43} \right), \ S_{32}(A_1) = \frac{1}{\sqrt{2}} \left( R_{32} + R_{33} \right), \ S_{33}(B_2) = \frac{1}{\sqrt{2}} \left( R_{32} - R_{33} \right), \\ S_{34}(A_1) &= \frac{1}{2} \left( R_{34} + R_{34} + R_{33} + R_{34} \right), \ S_{33}(A_2) = \frac{1}{2} \left( R_{31} - R_{32} - R_{33} \right), \\ S_{34}^+(B_1) &= \frac{1}{2} \left( R_{34} + R_{34} - R_{34} - R_{34} \right), \ S_{33}(A_2) = \frac{1}{2} \left( R_{41} + R_{44} - R_{47} - R_{13} \right). \end{split}$

Table 1. The combination and their normalization coefficients of (3,0) nanotube in D3d point group (Lee et al., 2009)

The character of the system assigned by  $\Gamma$  was calculated from the character tables and the UMAT are written from the application of projection operator. Vibrational Calculation was carried out by the MOLVIB algorithms and by Hyper Chem. Calculation and a few sets of calculation were performed. Molecular motions can be assigned by the potential energy distribution (PED) analysis among internal coordinates by the method of the projection operator. There are good agreements between the most cases.

#### 2.6 Normal mode dependence on dielectric

As can be inferred from Table 1 and the Fig. 3, 4, and 5, there are good agreements between the semi and Monte Carlo and even ab initio calculation. In Table 1 the various of intensity and frequency and potential energy from different methods are shown versus the inverse dielectric for some normal modes. From Fig. 3 we have two maximum for both of energy and frequency in the dielectric between 77.40 up to 70.42 and also the third maximum is located in the 61.76. This region range is considered to be the unstable geometry of nanotubes which are very sensitive to dielectric. After these range the frequency, intensity and energy goes toward a stable geometry which are not sensitive to dielectric. The same results are obtained in the insets Fig. B and C of Fig. 3 for D3d of normal mode 61 and 66 respectively. In the Fig 4, similar to normal mode 1 and 131 and 138 are shown with A, B, and C respectively for nano tube (4 0) in D4d point group, a common general behavior is observed in this nanotube as same as (3 0) nanotube, only with a shift in data, this shift is due to the difference between the geometrical structures of two nanotubes.

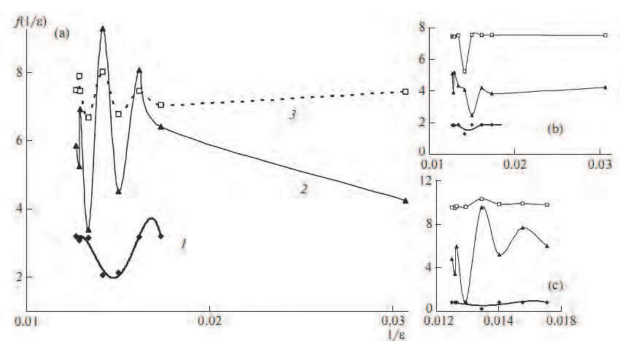


Fig. 3. The natural logarithms of the potential energy (1), intensity (2), and frequency (3) of three normal modes (a) 1, (b) 61, (c) 66 versus inverse of dielectric constant for nanotube (3, 0) with D3d point group by AM1 calculation (Lee et al., 2009)

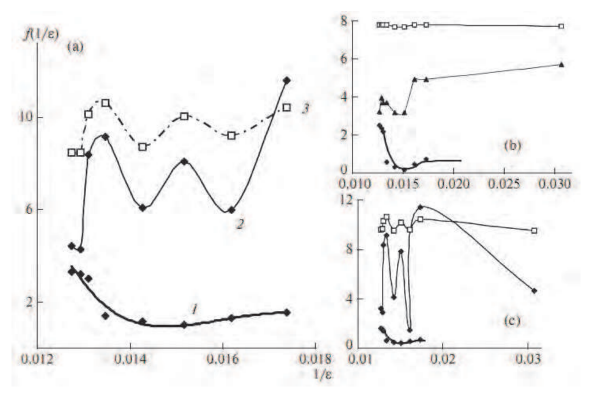


Fig. 4. The logarithms of the potential energy (1), intensity (2), and frequency (3) of three normal modes (a) 1, (b) 131, (c) 138 versus inverse of dielectric constant for nanotube (4, 0) with D4d point group by AM1 calculation (Lee et al., 2009)

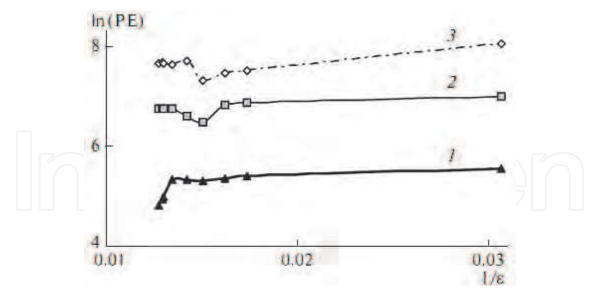


Fig. 5. The logarithms of the potential energy (PE) of three different zigzag nanotubes (1) D3d, (2) D4d, (3) D5d versus inverse of dielectric constant by MC simulations (Lee et al., 2009)

#### 2.7 Potential energy dependence on the dielectrics of zigzag nanotubes

In Fig. 3 three line of potential energy for three nanotubes are shown by the Monte Carlo calculation versus dielectric constants. For D3d symmetric nanotube, the logarithm of potential energy increases as the dielectric constant reduces from 78.39 up to 76.10 while the D4d and D5d symmetries show an unchanged potential energy in this region. Beyond this point, the potential energy of D4d and D5d nanotubes drop and rise again to be in a new equilibrium, whereas the potential energy of the D3d mostly constant as the dielectric constant decreases. There are some changing in the energy in variable with the dielectrics above 60, and by decreasing the dielectrics the energy of three nanotubes goes toward constant variables. Similar trends between three figures and there are very good agreement with *ab initio* calculation in the Table 1.

## 3. Stability of SWCNTs: Solvents and temperature effects by molecular dynamics simulation and quantum mechanics calculations

Structural properties of solvents such as water, methanol, and ethanol surrounding single-walled carbon nanotube (SWCNT) and mixtures of them as well have an effects on the relative energies and dipole moment values. Because some of the physicochemical parameters are elated to structural properties of SWCNT, the different force fields can be examined to determine energy and other types of geometrical parameters, on the particular SWCNT. Because of the differences among force fields, the energy of a molecule calculated using two different force fields will not be the same. The structure of SWCNT as well as its dipole moments and relative energies has been studied by molecular dynamics simulation and quantum mechanics calculations (Monajjemi et al., 2010). The term "Ab Initio" is given to computations which are derived directly from theoretical principles, with no inclusion of experimental data. The most common type of ab initio calculation is called a Hartree-Fock (HF) calculation, in which the primary approximation is called the central field approximation. A method, which avoids making the HF mistakes in the first place, is called

Quantum Monte Carlo (QMC). There are several favors of QMC variational, diffusion, and Green's functions. These methods work with an explicitly correlated wave function and evaluate integrals numerically using a Monte Carlo integration. These calculations can be very time consuming, but they are probably the most accurate methods known today. In general, *ab initio* calculations give very good qualitative results and can give increasingly accurate quantitative results as the molecules in question become smaller (Monajjemi et al., 2008a). In general, there are three steps in carrying out any quantum mechanical calculation. First, prepare a molecule with an appropriate starting geometry. Second, choose a calculation method and its associated options. Third, choose the type of calculation with the relevant options and finally, analyze the results. We will give a short detail of computational method in the following section.

#### 3.1 Molecular mechanics (Monte Carlo simulation)

The Metropolis implementation of the Monte Carlo algorithm has been developed by studying the equilibrium thermodynamics of many-body systems. Choosing small trial moves, the trajectories obtained applying this algorithmagree with those obtained by Langevin's dynamics (Tiana et al., 2007). This is understandable because the Monte Carlo simulations always detect the so-called "important phase space" regions which are of low energy (Liu & Monson, 2005). Because of imperfections of the force field, this lowest energy basin usually does not correspond to the native state in most cases, so the rank of native structure in those decoys produced by the force field itself is poor. In density function theory the exact exchange (HF) for a single determination is replaced by a more general expression of the exchange correlation functional, which can include terms accounting for both exchange energy and the electron correlation, which is omitted from HartreeFock theory:

$$E_{ks}=v++1/2+E_{\gamma(\rho)}+E_{C(\rho)}$$

where  $E_{\chi(p)}$  is the exchange function and  $E_{C(p)}$  is the correlation functional. The correlation function of Lee et al. includes both local and nonlocal terms (Lee et al., 1988).

#### 3.2 Langevin dynamics (LD) simulation

The Langevin equation is a stochastic differential equation in which two force terms have been added to Newton's second law to approximate the effects of neglected degrees of freedom (Wang & Skeel, 2003). These simulations can be much faster than molecular dynamics. The molecular dynamics method is useful for calculating the time-dependent properties of an isolated molecule. However, more often, one is interested in the properties of a molecule that is interacting with other molecules.

### 3.3 Effect of differenct solvents of temperatures of SWCNTusing molecular dynamics simulation and quantum mechanics calculations

Difference in force field is illustrated by comparing the energy calculated by using force fields, MM+, Amber, and Bio+. The quantum mechanics (QM) calculations were carried out with the GAUSSIAN98 program based on HF/3-21G level. In the Gaussian program a simple approximation is used in which the volume of the solute is used to compute the radius of a cavity which forms the hypothetical surface of the molecule (Witanowski et al., 2002; Mora-Diez et al., 2006). The structures in gas phase and different solvent media such as

water, methanol, ethanol, and mixtures of them have been compared. The structure of SWCNT as well as its dipole moments and relative energies has been studied by molecular dynamics simulation and quantum mechanics calculations within the Onsager self-consistent reaction field (SCRF) model using a Hartree-Fock method (HF) at the HF/3-21G level and the structural stability of considered nanotube in different solvent media and temperature (between 309K and 327K) have been compared and analyzed.

Since the influence between a molecule in solution and its medium can describe most simply by using Onsager model, in this model we have assumed that the solute is placed in a spherical cavity inside the solvent. The latter is described as a homogeneous, polarizable medium of dielectric constant. We started our studies with HF/3-21G gas phase geometry and water, methanol and ethanol surrounding SWCNT and mixtures of them as well. The results obtained from Onsager model calculations are illustrated using the energy difference between these conformers which are quite sensitive to the polarity of the surrounding solvent. The solvent effect has been calculated using SCRF model. According to this method, the total energy of solute and solvent, which depends on the dielectric constant  $\varepsilon$  has been listed in Table 2.

	Dielectric constant					Tempera	iture (K)				
Medium	$\Delta E  (\mathrm{Kcal/mol})$	309	311	313	315	317	319	321	323	325	327
Gas phase	1	-0.1075	-0.1227	-0.0916	-0.1878	-0.1287	-0.1575	( = i)	-0.0511	0	-0.1721
Water	78.39	-0.1929	-0.1965	-0.1573	0	-	-0.2629	-0.2316	-0.2517		
Methanol	32.63	-0.0743	0	-	-0.0653	-0.1936	-0.0545	-0.1942	-0.0056	-0.0615	-0.0770
Ethanol	24.55	0	26	-0.0129	-0.0120	-0.0382	-0.0764	-0.0901		-0.0918	
Water-Methanol	70.763	-0.0029		-0.0790	-0.0846	_	-0.0918	-0.0483	-0.0510	0	-0.0485
Water-Ethanol	69.416	-0.0840	-0.0852	-0.0442	-0.0215	20	-0.0276	-0.0890	0	-0.0144	-0.0610

Table 2. Theoretical relative energies at different temperature and dielectric constant

These energies have been compared with the gas phase total energy CNT at the HF/3-21G level of theory and different solvents, and the graph of energy values versus dielectric constant of different solvents has been displayed at considered temperatures in Fig. 6.

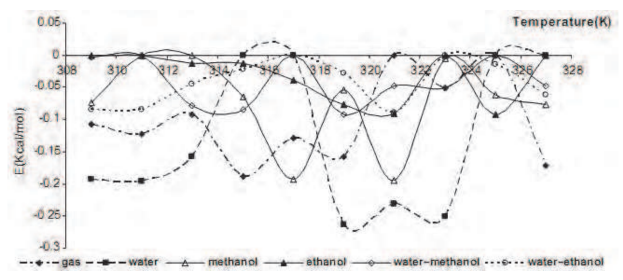


Fig. 6. The relative energy values at different temperatures in different solvents.

Since the solute dipole moment induces a dipole moment in opposite direction in the surrounding medium, polarization of the medium in turn polarizes the charge distribution in the solvent. The dipole moment value of SWCNT in different solvent media and at different temperatures has been reported in Table 3.

Medium	Temperature (K)											
Dielectric constant	309	311	313	315	317	319	321	323	325	327		
Gas phase	14.8595	14.9783	15.8972	1.4569	12.7230	0.9106	N-4	15.1528	1.5045	0.7366		
Water	1.3302	1.8910	1.0015	7.0286	_	0.7134	1.6209	0.9655	-	_		
Methanol	1.4854	22.4244		9.5556	5.1790	9.4344	7.7342	21.3318	7.4751	7.2150		
Ethanol	0.7029		2.4926	11.8624	14.9195	0.9389	0.9646	-	0.4796	9 <u>2. 33</u>		
Water-Methanol	12.4020	46.8625	4.4462	4.2048	-	4.8710	5.7352	5.9246	3.8414	6.1855		
Water-Ethanol	0.8771	5.9677	6.3670	6.6015	-	3.8895	4.4362	6.4769	4.9592	5.8514		

Table 3. Theoretical dipole moment values at different temperatures

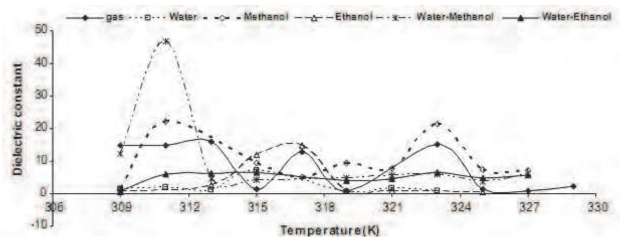


Fig. 7. The dipole moment values at different temperatures

One much more practical approach consists of calculating the molecular volume as defined through the contour of constant electron density, equating this (nonspherical) molecular volume to the radius of an ideally spherical cavity, and adding a constant increment for the closest possible approach of solvent molecules. This latter approach was used in Gaussian when the volume keyword was being used. In this work, we studied the structural properties of water, methanol, and ethanol surrounding SWCNT and mixtures of them as well as using molecular dynamics simulations. We used different force fields for determination of energy and other types of geometrical parameters, on the particular SWCNT. Because of the differences among force fields, the energy of a molecule calculated using two different force fields will not be the same. So, it is not reasonable to compare the energy of one molecule calculated with a particular force field with the energy of another molecule calculated using a different force field. In this study difference in force field illustrated by comparing the energy calculated by using force fields, MM+, AMBER, and BIO+. Theoretical energy values using difference force fields which are the combination of attraction van der Waals forces due to dipole-dipole interactions and empirical repulsive forces due to Pauli repulsion have been demonstrated in Table 4 and Fig. 8.

Medium	Dielectric constant	The Mr. S.	E (kcal/mol	LVA SILIA
Gas phase	1	487.9812	382.4433	1628.176
Water	78.39	419.9189	316.7664	1560.24
Methanol	32.63	535.3614	431.2506	1533.685
Ethanol	24.55	468.4228	359.4595	1601.808
Water-Methanol	70.763	413.1738	309.1905	1484.31
Water-Ethanol	69.416	637.6032	539.6583	1476.198

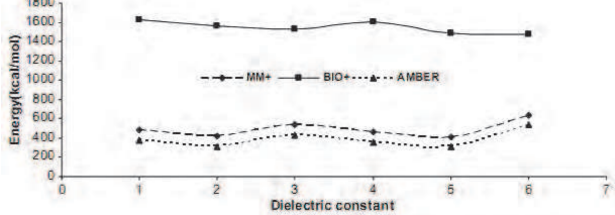


Fig. 8. The energy values using different force fields

The result of the calculated dipole moment, quadrupole moment, octapole moment, and hexadecapole moment values of SWCNT has been reported in Table 5, and optimized structures of nanotube in different media are shown in Fig. 9.

		Di	pole mom	first	Qua	drupole mo	mart		00	apole mome	it					Hexadecapole	moment				
		x	¥	2	XX	YY	ZZ	XXX	YYY	222	XYY	XXY	XXXX	YYYY	2222	XXXY	XXXZ	YYYX	YYYZ	ZZZX	
Solvair	Tamp		TOT		XY	XZ	YZ	XXZ	XZZ	YZZ	YYZ	XYZ	222Y	XXYY	XX22	YYZZ	XXYZ	YYXZ	ZZXY		ΔĐ
SWCNT <sub>III</sub> - (4.7)	809	-14.6440	0.1798 14.8595	-2.5154		-856.2825 -18.7461	1100000	-1865.5072 -163.5864	8.5450 -38.7868	-28.9528 3.5868	-80.9815 -5.9538	8:0217 -2:0505		U. 200 march	-15289.2158 -19607.8861	77	5000 CO	81.5565 -28.5092	6.0818 -1.7187	-117.1608	-0.10781
	811	14,7152	9.1097 14.9788	1.8418	-807.8251 -91.6888	-650.1956 -16.8081			8.8112	7.6879 5.8740	88.5805 8.4026	182,9625 17,0082	-84,7751		-15186.5588 -19811.4622				-18,4200 -67,6681	-108.4675	-0.12279
	818	15.4846	-2.4142 15.8972	2.6682	-818.8514 22.6096				-12.1769 68.7774	16.5687 -7.0598	82:9814 7:5674	-198.6747 -12.5116	-6.1266		-15085.1868 -19681.8544		-1609.4717 194.6596		25.23.40 52.2818	-107.2728	-0.00162
	815	0.4058	-0.9088 1.4569	1.0256	-712,1719 -1,4415		-655,5195	78.0014	1.6508 -15.4389	14/0220	27.5257 2.5424	-88.2198 6.0805	5.1805		-15084.1688 -19162.4177	3.30	-957.0449 -79.8518		-14.0118 -6.8946	-24.9192	-0.18788
	317)	18,7077	0.5180	-0.3458	-797,7998 -0,5071	W-33	57.	1995,0914 -51,5899	1.8741 81.7008	-8.1095 -0.5721	18:4802 1.8640	40,7788	0.9670		-15081.0940 -10870.8155		530,3633 71,3800	-98.1915 -87.6927	10,8582 -25,4519	184.9009	-0.12871
	319	-0.8855	-0.8186 0.9106	0.1865	-715.8088 6.2185	-660.1265 0.7884		+16.8860 4.8022	-5.4686 1.9000	-1.7150 -2.2088	-17:5566 1:5087	+26.1266 -9.8140	17.7446		-15140.8960 -19185.9741		-16,7640 -82,0774		-58.8547 20.6524	9.4804	-0.15756
	821	41	4	2	_	20	<u>~</u>	20	2	_	4	20	<u>~</u>	-	2	<u>a</u>	-	-2	_	_	25
	928	-15.0621	-1.6222 15.1528	-0.8827	-807,7058 +28,7476			-1869.0078 -20.9444		2.4810 -4.1928	-66.1265 0.2815		-40.8885		-14980.8568 -19849.2798			-914.8518 -118.8090	1,9922 -68,6385	-17.4442	-0.05117
	825	-0.1757	0.7800 1.5045	1.2987	-780.6618 2.4855	1.30		-81.5621 104.6224	8.8981 14.7922	22.7699 -1.6996	-91.7619 1.4874	52.8000 -15.8525	11.8987		-15183.8142 -10508.0024	2430	-488.4887 219.0057		15.1284 87.9482	52.2549	0
	827	0.0947	-0.6480 0.7866	-0.8874	-718.5698 6.1065			44,8072	-8.8020 29.1555	-0.8140 -2.4015	-21:7776 -7:6069	-71.8008 80.6841	28.2809	The State of	-15042.5782 -10183.7106	TO SERVICE STATE OF THE PARTY O	486.8275 195.8418	79.5713 28.7602	-11.8200 14.5437	18,5049	-0.19484
	829	-1.5989	1.0704	-1.1918	-782.6384 11.4800				12.0277 84.0020	-29,5590 5,8512	-58:9188 8:1985		-51,2774		-15283.1608 -19522.6786		265.8962 -451.6274		-44.7107 106.8356	55,0741	-9.17912

		Di	pole mom	ant	Qua	efrupole mo	meent		Oc	tapole mome	nt				- 1	Hexadecapole	moment				
		x	Y	Z	xx	YY	ZZ	XXX	YYY	ZZZ	XYY	XXY	XXXX	YYYY	ZZZZ	XXXY	XXXZ	YYYX	YYYZ	ZZZX	
Solvent	Temp	E	TOT		XY	XZ	YZ	XXZ	XZZ	YZZ	YYZ	XYZ	ZZZY	XXYY	XXZZ	YYZZ	XXYZ	YYXZ	ZZXY		ΔE
SWCNT <sub>He</sub>	300	0.0078	0.2025	-0.0500	-701.6414	-659,6880	-650.2722	98.1872	-1,7948	-12.0079	20.6872	18.0168	*****	-15454.1671	-15158/1740	400.6782	186,5951	48.5804	20.8502	84,2050	-0.10205
(,7)-water	811	0.6781	1.8802 -1.6782	0.5587	6.2040		1.2202 -660.8602	-00.7160 78.6571	0.7986	9.2518 -8.2561	1.1876	14.2028 -114.5140		-10780.6881 -16816.6978			44.4400 -887.0681	-51.0879 -69.7868	88.5765 26.7012	-61,9885	-0.196564
			1,8910		7:4520	-8.8901	0.8200	40.5644	-5.6268	-8.7160	-1.7549	-19.8898		-10899.1198			188.7880	-1.4452	80.2818		
	818	-0.0178	-0.1928 1.0015	-0.8526	-710.8840 -2.8107		-860.9079 -0.4097	-110.0898 -27.0978	-6.0947 -16.9197	-8.0210 7.0109	1.8566	-27:0916 18:9556		-15541.5077 2 -19694.5970			-416.0805 8.8751	-61.2680 -4.6660	-97.1668 -8.9262	-64,7470	-0.157321
	815	2,0644	-1.9958	6.0525	-919.8898			288.6782		45.8260		-169.0800		-15854.1849				-26.8119		-115.8540	0
	1228		7.0286		-0.7842	-2.1383	1.7635	481.3963	65.4141	-8.6692	9.2140	-2.1691	-8.779	-20571.4992	-20208.1909	-4986.6548	141.2782	85.4590	41.1084		
	817	-	-	-	-0	-	_	-	_	-	-	-01	_	_	_	_	-	_	_	-	
	819	-0.8182	0.6148	0.1881	-717:1820 5:8925		-658.0192 0.9784	-47.0089 11.1880	13.0846	6.1816 1.0844	-19.9581 -1.1981	24.6970 15.6969	-14 898	-15548.7987 8 -19602.2749				66.9447 2.8647	15.9164	-56.9492	-0.262943
	321	-1.5387		-0.1057	-791.8000		1000		-6.9968	3.8848	-61:9817	-98.4512		-15812.4488				15000	-7.2658	8.2568	-9.231687
			1.6200		-8.7784	3.6885	-2.7907	8.6626	38.8813	-4.1044	-9,7555	-15.1862	-15.895	-20032.9216	-19247.4581	-5149.7668	-215.4586	28,8882	-29,2197		
	828	-0.0868	0.4220	0.8658	-721.0088 2.8852		-668.4265 -2.4879	5.0558 64.6885	2:1894 -10:6017	1.8781 -5.5754	9.1011	87.4186 18.8921		-15586,2084 -19698,1419			-516,1989 -86,2298	-10:8218 -44:8580	-84.8078 -17.7865	-99.5529	-0.951758
	825	-	-	_			_		-	_	_		-	_		_	-	_		-	_
			-		-	-	-	-34		-	-	-	-	-		-		-	-		
	329		-	_	_	-	_	-40	_	1	-	4	-	_	_	_	-	_	_	_	-
																					_
		X	y Y	Z	XX	trupole noon	ZZ	XXX	YYY	epole momen	XYY	XXY	XXXX	YYYY	2222	XXXY	XXXZ	YYYX	YYYZ	ZZZX	
Solvan	Temp		TOT		XY	XZ	YZ	XXZ	XZZ	YZZ	YYZ	XYZ	ZZZY	XXYY	XXZZ	YYZZ	XXYZ	VYXZ	ZZXY	LLLA	ΔE
SWCXT <sub>IIP</sub>	800	0.2801		-0.6917	-720.7922	Weekly .	-650,7962	49.1538	8.1861	2,8580	4.7784	3.8844	Associate 1	-15479.6684	SACILIANS.	850.4418	616.7408	0.900+	-16.4335	68.7184	0
(,7)-erhanol			0.7020		11.1676	6.0212	-1.2966	-48.9874	8.0104	7:6095	-1.8645	-17.0071		-10523.0452			-108.8600	5.4027	7,0261		
	811	-	-	-	-	-	-24	-	0.770	<b>177</b>	-	-	-	-	10.000	-84	-	25	.=	10	-
	318	n anew		En anes	-718.0002		-861.7489	191.5845	15,0467	-2.6289	6,4948	79.7805		-15381,7243	-11000 0000	71,5196	-818.9898	95,1178	- TO 100	-53.9676	20.010000
	918	43481	2.4926	-0.0018	1.8857	-4.7548	-7.8847	-77.6895	28.5458	-2.4518	2.4626	64.2695		-19581.6480				13.0305	-4.2161		70012992
	815	-10.8367	-2.8081	-4.0597	-775.4044	-657,7152	-658 2211	-986.0580	-11.5718	-88.1805	-65.8448	-246.7706		-15487.7086	-14898,7005	-2298.7486	-2482.9815	-225.0128	-64.8804	-195,6884	-0.012007
			11.8624			-80,0028	-5.9464	-287, 4888		-15,2682		-83.1000		-19909.6824							
	317	-14.8319	0.5870	-1,5997	-649.1182 3.0209	-854.8977 5.8111	-659.1986 -4.4158		-8.0525 -46.0965	-6.8110 5.7715	-88.9254 -0.2718	34.5488		-15468.8268 -10295.7204				20.6689	-80.1784 0.7480	19:2782	-0.038208
	819	-0.0060	0.7655	0.5850	-724.1507	-660.2018	-061.8000	18.2858	18.9486	8.0818	-8.1416	1.4580		-15504.7425	-15141,2187	266.6062	-284.8215	62.2686	3.0252	-98.5704	-0.076462
			0.0889		5.2801	-4.1268	-0,6848	89.5798	11,4550	8.8070		-19.4796		-19869,2985				-48.6580	68.0648		
	321	-0.4662	0.8892	0.0987	-712.4428 -7.4881	-662.8886 8.0080	-655.6262 0.7290	-28.7600 1.1267	4.5630	2.8793	+04.1119 -8.1459	-17.5582		-15545.0286 -10968.5852			108.7805 49.8215	-27.5819 24.5080	5.4363	26.7263	-0.09019
	823	-	-	-	-	-	T-87	-	-	77	-	-	-	-	-	-	-	15	-	177	-
			-		- T	<del>-1</del> 8	T82	<del>-3</del> 8	-	77	- <del></del>	- T	<u> </u>	S 77	-76	TEX	5-22	855	· 53		
	325	-0.0367	-0.1718 0.4796	0.4452	-714.7845 -5.6081	-661.9806 10.1184	-856.1977 0.4297	-5.2028 25.0012	-1.0884 -21.0880	-4:9482 -1:0686	11:8696 8:0058	-82,7107 8,3842		-15890.1851 -19909.9894			770.0961 37,7004	-79.5779 . 25.6834	32.1014 20.1106	68:2614	-0.09184
	827	-	0.400	_	-0.0001	-		-		-13000	-		3.0000	-			-	-0.0834	_	_	-
			-		-	-	-30	-	7	36 37	-	-	-	-	7	-30	-		-70		
		X	pole mom	ant Z	Qua	YY	mad 22	XXX	YYY	rapole mome	XYY	XXY	XXXX	YYYY	2222	Hexadecape	NKXZ	YYYX	YYYZ	ZZZX	-
Solvent	Temp		TOT		XY	XZ	YZ	XXZ	XZZ	YZZ	YYZ	XYZ	222Y	XXYY	XX22	YYZZ	XXYZ	YYXZ	ZZXY		45
SWCNT <sub>III</sub>	809	0.0008	-0.8428	1 0000	-788.4178	-ees once		-04 7057	-7.4988	17,2280	88.8576	-46.9181		-15591.168		0 56.187	7 768.018	9,7876	21.6808	NA DED	7 -0.0743
(4.7)-mathana		0.000	1.4854	1.4000	-2,7156				-38.7405	-9.4087				-10001.108 05 -10002.474					-58.9477	41,000	- Lateral
	811	-22.1691	-8.0275 22.4244	-1.4898	-848.0728 -16.5498		-659.4412 1,4492	-1803.7568 -108.5824		-5.1855 -6.7140	-89.9411 -7.1575	-287,1919 28,9161		-15718.504 16 -20006.050				71.8516	4,7828 -118,9810	88.261	0
	818	-	-	-	-20.0405	-	-	-110.0024		_	-1.401-	_	_	-2.0.00.3001		0210.080	-	-	-	-	-
			-07		-	-	-	-	-	-	-0	-	-	-	-	-	-	-	-		
	815	-9.4004	0.6458	1.5890	6.2405			-792,1821 161,1667	-6.0824 -69.1918	12.0697 -7.1102	-56,2600 12,1004			-15549.140 96 -20050.955	8 -15105.105 9 -10875.090		8 2746.0020 1 844.1687	0 101.860f 7 121.8207			0.0858
	817	-4.8984	1.6719	0.2802				-424.0047			-106.7746			-15407.574							5 -0.1986
			61700			-6.8753				-0.8222	7.1808			50 -19966.174							
	819	-0.0188	9.8794	-1.4596	768,9798 19,1250			-807.1521 -07.8407		-4.4180 0.5194	-108.8988 -7.2787			-15546.687 8 -20156.618				249.7531 -20.9878		-20.001	4 -0.0545
	821	7.5748		-1.5470				648.0518		-22.1678				-15516.582						186.989	4 -0.1942
			7.7842		-8.7412	14.1488	-4.4997	-118.0825	54.6421	-8.7788	-5.1815	39.0484	-0.949	06 -19654.850	8 -20027.701	5 -5749.824	9 -418.887	7 80.5004	52,1812		
	323	91.9919	1.0895	0.9666			-653.0300 -4.4284	1778.9615 36.7607	-12.7977 98.1184	9.1955	109.5071			-15535.256 86 -20304.190						-120.820	a -0.0056
	825	-710492	-1.9855	1.6621				-544.6619		2.9082		-108.5488		-15597.055						94.952	9 -0.0615
			7.4751		-15.5308				-22,6981	-7.8816	5.8047			2 -20164.085							
	327	-7,0087		0.7735				-715.1820		2.0462				-15647.811							0 -0.070
			72150		16:7621	-7.6146	-1.8681	97.4825	-14:4641	-0.0950	4.9120	-16.8116	-16.48	19 -20224.822	+19477.820	+ -6140.792	8 -182/0418	44.3731	6.2946		

		Dip	sole mome	nt	Qua	irupole nior	uart		Qet	spole monest	ij.				- 1	Hexadempole	moment				
		X	Y	Z	xx	YY	22	XXX	YYY	222	XYY	XXY	XXXX	YYYY	2222	XXXY	XXXZ	YYYX	YYYZ	ZZZX	
alvent	Temp		TOT		XY	XZ	YZ	XXZ	XZZ	YZZ	YYZ	XYZ	ZZZY	XXYY	XXZZ	YYZZ	XXYZ	YYXZ	ZZXY		ΔE
WENT	300	0.6751	0.4929	-0.2656	-712.5204	-655.2920	-663.5865	88:9541	4.7006	-3.7045	28.8280	-4.0988		-15425,2285	-14705.8716	-322.8457	-885,2783	-91.6718	7.9264	-2.6228	-0.08409
(4,7)-water =			0.8771		-5.5040	-5.4217	1,2912	-2.5657	-94.9551	8.8902	4.0472	-19.9889	18.0864	-19563.9745	-19988.9281	-5024.1802	129,1484	-68.4288	-24.6220		
ethanol	311	-5.5825	-2.0802	0.0805	-782.2581	-661.7991	-664.7478	-455, 1862	-7.5924	15.8594	84.4629	-116.0547		-16609.5859	-15464.7467	-649.8821	2086,5891	-60,5729	0.8470	821.7151	-0.08522
			5.0677		-10.1584	24,7078	4.5989	122.0901	-94.0679	-16.4499	3.4878	21.2429	75.1825	-10746.7585	-10088.7274	-5854.2888	208.8526	21.0294	-163.9256		
	313	-5.7577	-1.9182	1:9257	-729.6999	-861.8050	-606.4156	-470.8752	-11.8652	23.0506	-9,4557	-102.0868		-16386.2652	-15508.7958	-88.8670	1097,9675	-8.2373	24,1864	154.0812	-0.04428
			6.3570		-2.8586	12.8961	5.8830	182,2235	-51,1778	-19.9148	0.8418	27.6542	31:2730	-19652.0260	-19877,7251	-6897.7896	296.6608	86.0241	-140.9214		
	815	-6.0967	-15304	2,0100	-789.6855	-659:4816	-667,9510	-456.6356	-18.9449	33.0751	18.8187	-84.3076		-16745/0814	-15508.9051	156,6532	884.4741	84,1865	1.8435	124.5123	-0.02155
			6,6015		0.4228	10.4290	3.6778	180.5086	-74.0004	-18.6272	-5.2588	47.5489	35,8160	-10587.8082	-19918.5928	-5876.6258	271.6607	19,4347	-126.8068		
	317	-	12	-	-	-	-22	-	22	-	12	-	-20	-	22	- 225	-	46	-	-	
			7		-	-	-	-	-	-	7 <del>10</del>	-	-	-	100	-	-	-01	-0)		
	310	3.6530	-0.5571	-1.2138	-741.7181	-658,7982	-609:0092	880.4041	-12.0154	-81.1418	14,1088	1.7784		-16762.4308	-15507,5685	482.0524	989.6860	41.7507	-24.9775	177,1792	-0.02761
			3.8805		7.8811	10.2308	+2.1702	-120.2621	28.0542	-8.8930	4.9285	9.7397	-18.1290	-10467.3868	-20018.8812	-5368.4028	-136,5317	-81.1998	104.7985		
	321	-4.2787	-1.1785	0.1629	-784.0286	-658.9000	-666,2975	-858,0007	-6.9988	5.5957	-11.8927	-58:6552		-16826,9490	-15529.4892	-1114.0878	2005.0098	-158 1294	18.1706	828,0898	-0.08901
			4.4862		-16.3384	25.8618	4.8309	28.4876	-28.4655	-17.0712	3.9308	85,2408	11.8985	-19450.6580	-19865,0505	-5406,7469	225,7249	24,2015	-159,0748		
	328	-6.4297	0.2887	0.7430	-746.4179	-661.1947	-668.6051	-535,8458	2,7830	16.7089	-18.4007	88.4890	******	-16628.9472	-16714.0099	-1198.8638	\$51,6483	-208.3875	16:0704	89.6261	0
			6.4769		-17.1796	7.9488	4.8720	84.8085	-51.1856	-15.8860	6.1600	85.7176		-19489.8441			295,9076		-189.6450		
	325	-9,4107	1,7180	-1,4580	+148.2502	-657.5881	-008,5018	-390,0851	25.7112	-15.4201	-26.6555	77,1864		+16554.2850	-15714.6875	885,0122	+1184.6577	109.8558	82.8184	-186,7683	-0.01448
	- BOK	- ALACANA	4,0592			-18.6525	4.4219	-128.8076	-25.6500	18.1474	-4.6996	26.9011	47.8881	-19208,1286			855,0758	-62.0532	178.0200	1500 (100)	
	207	5.7009	0.9564	0.0078	-740.9578	-661.7147	-064.2185	480.0848	7,2989	12.6085	28.5207	72.6647		-18690.0002	-15000.1424	-059.2665	-1600.9506	-14E 1430	-42.4670	-280.6280	-0.05101
<u> </u>	-		5:8514 Olpole mar		-18.8708	-20,7771 adrupole m	+6.5813	111.0608	87.0460	11:9591 tapole mouse	1:8006 nt	87 8983	-79.1748	-19411.3850	-19505.5488			-26.2798	-122/2988		v.
			· anti-		-18.8708	-90,7771	+6.5813		87.0460	11.9591		AT MARK	-79.1748	~19411.1850) YYYY	-19505.5488 2222	-8452,6271		-26.2798 YYYX	-120.2988 YYYZ	ZZZX	V
Solvent	Taur	X	Olpole mon	nent	-18:8708 Qu	-20,7771 adrupole m	-6.5811 ouseut	111.0008	87.0460 Oc	11.9591 tapole nimos	in the second					-5452.5271 Hexadeospole	i monient			ZZZX	ΔE
Solvent SWCNT <sub>112</sub>	Taup	X	Dipole mor	nent Z	-18.8708 Qu XX	-20.7771 adrupole m VY X2	-6.5811 onsent 22 Y2	111.0608 XXX	97.0460 Oc VVV X2Z	11.9591 tapole minus	XYY	XXY	XXXX	YYYY	2222	-5452.5271 Hexadeospoi NXXV VY2Z	XXXZ XXXZ	YYYX	YYYZ	discount of the same of the sa	ΔE
-	Taur	X	Dipole mor Y TOT	2 4 3 7 2	-18.8708 Qu XX XX	-20.7771 adrupole m VV XZ 5 -658.063	-6.5811 onsent 22 Y2 8 -656.6401	XXX XXX XXZ 1 -1159.8462	97.0460 VVV X2Z 14.7805	11.9591 tapole number 222 Y22	XYY YYZ	XXY	XXXX	YYYY XXYY -15635,9311	2222 XX22	-5452.5271 Hexadeospoi NXXV VY2Z	XXXZ XXYZ XXYZ	YYYX	YYYZ ZZXY	discount of the same of the sa	no de la constanta
SWCNTing	Taur	-11.388	TOT 12.4020	2 2 2 4171	-18.8708	-20.7771 adrupole m VV XZ 5 -658.000 6 #2.516	-6.5813 ousset 22 V2 8 -656.6492 0 -4.880	XXX XXZ 1 -1159.8462 380.9982	97.0460 999 922 14.7805 -86.8268	11.9591 tapole number 222 Y22 41.5695	XYY  YYZ  -60.4705	XXV XYZ 217.5594 -72.4616	XXXX ZZZV	VYYY  XXYY  -15635,0319 2 -20204,2667	2222 XX22 -14508.0011 -10726.7141	-5452,5271  Hera-lexapole  NXXV	XXXZ XXYZ XXYZ	YYYX YYXZ 105.1808 58.5011	YYY2 ZZXY -88/2100 78/8472	discount of the same of the sa	-0.00206
SWCNT <sub>112</sub> (4.7)-water	Tour	-11.388	TOT 12.4020	2 4.171 1 -48.025	-18.8708 Qu XX XY XY -701.4888-020.068	-20.7771 adrupole m VV XZ 5 -658.000 6 #2.516	HI J811  misert  ZZ  YZ  8 -656,0-001  0 -4,080  0 -881,0651	XXX XXZ 2 -1159.8462 480.9982 5 621.0819	97.0460 999 922 14.7805 -86.8268	11.9591 tapole nimos 222 Y22 41.5695 1.2714	NYY VYZ -60.4705 8.5941	XXV XYZ 217.5504 -72.4616 6.3188	XXXX ZZZV	YYYY XXYY -15635,9311	2222 XX22 -14508:0011 -10726:7141 -28056:2845	-5452,5271  Hexadeospole  NXXV	8 MONIONI S 8 M M M M M M M M M M M M M M M M M M M	YYYX YYXZ 105.1808 58.5911	YYY2 ZZXY -88/2100 78/8472	847.8424	-0.00206
SWCNT <sub>112</sub>	Tour	-11.385 -8.776	Pipole more  Y  TOT  12.4020 14.4020 14.4030	2 4170 1 -46.029	-18.8708 Qu XX XY XY -701.4888-020.068	-20.7771  adrupole m  VV  X2  5 -658.600  6 42.516  8 -687.581  2 -176.759	V2  V2  V2  8 -656,6401 0 -881,6656 6 -2,7680	XXX XXZ 2 -1159.8402 3 680.9982 5 621.0810 1 -657.4988	97.0460 VVV NZZ 14.7805 -36.8268 -1.8798	11.9591 222 Y22 41.5695 1.9714 -1.433.9166	NYV VYZ -60.4705 8.5941	XXV XYZ 217.5504 -72.4616 6.3168 -87.7188	XXXX ZZZV 9.095	YYYY  XXYY  -15635,0319 2 -20004,2667 -15710,0006	2222 XX22 -14508-0011 -10726-7141 -28056-2841 -22405-5008	-5450.5271  Heradeoquil  XXXV	8 monisuit 8XXZ XXYZ 8524.4047 -8014.6728 288.8880	YYYX YYXZ 105.1808 88.5911 110.0702	VVVZ ZZXY -88.2100 78.8479 16.5485	847.8424 -5186.7589	-0.00206 -86.7480
SWCNT <sub>112</sub> (4.7)-water	Tour 800	-11.385 -8.776	TOT 12.6020 440.6030 46.8020	2 4175 1 -48.025	Qu XX XY XY -101.458 22.978 88 -620.068 2.640 66 -749.118	-20.7771  adrupole m  VV  X2  5 -658.600  6 42.516  8 -687.581  2 -176.759	+6.5811  COLUMN T	XXX XXZ 1 -1150.8402 480.9982 6 021.0819 1 -657.4983 896.8204	87.0460 VVV XZZ 14.7805 -86.8268 -1.8798 -870.6486 -18.5840	11.9591 tapele mon.e 222 Y22 41.593 1.2714 -1480.9166 -89.4802	XYV YYZ -60,4705 8,5941 125,4008 -16,5880	XXV XYZ 217.5504 -72.4616 6.3168 -87.7188 -82.1804	XXXX ZZZV 9.0950 - 182.618	YYYY  8XYY  -15635,0315 2 -20004,2667 -15710,3006 7 -19483,7004	2222 XXZZ -14508-9011 -19726-7141 -28056-2845 -22405-5098	-5452.5271  Hexadeospoli  XXXY  YYZZ  1075.8405  -5022.0658  820.4061  -5825.4211  758.0188	8 monisort XXXZ XXXZ 8524.4047 -880.4878 -8014.6728 288.8800 -856.6283	YYYX YYXZ 105.1808 88.5911 110.0702 -200.8256	VVV2 ZZXY -88.2100 78.8472 16.5485 -84.6808	847.8424 -5186.7589	-0.00206 -86.7480
SWCNT <sub>112</sub> (4.7)-water	Tour 800	-11.388 -8.779	TOT  \$5 2.5890 12.4020 44 -0.6030 45.8622	2 4.171 2 4.171 1 -46.028	Qu XX XY XY -101.458 22.978 88 -620.068 2.640 66 -749.118	-20.7771  edrupole m  YY  XZ  5 -658.000  6 42.516  8 -687.581  2 -176.759  9 -662.220  6 -10.602	+6.5811  22  V2  8 -656,4402  0 -4.860  0 -881,0656  8 -2.7082  5 -661,8855  6 .5861	XXX XXZ 1 -1158.8492 621.0810 621.0810 -057.4988 396.3294 181.0510	87.0460 VVV XZZ 14.7805 -86.8268 -1.8798 -870.6486 -18.5840	11.9591  zapris minuse  zzz  Yzz  41.5995 1.9714 -1.483.9165 -90.4892 -8.4895	XYV YYZ -60,4705 8,5941 125,4008 -16,5880 56,0750	XXY XYZ 217.5504 -72.4616 6.3168 -81.7180 -92.1804 -85.2807	XXXX ZZZV 9.095 -182.618	YYYY  XXYY  -1688 9319  2-20204 2667  -18710 3000  1-10488 70046  -16025 7088  2-20870 5692	2222 XXZZ -14508-9011 -19726-7141 -28056-2845 -22405-5098	-5452.5271  Hawadeospoli  5XXX  VYZZ  1075.8405  -5022.0058  820.4061  -5805.8211  758.0188  -5008.5564	8 moniest XXXZ XXVZ 8024-4047 -8014-6728 283-8800 -806-6283 410-3642	VVVX VVXZ 105.1808 58.5911 110.0792 -200.8276 45.7953	YYYZ ZZXY -88.2100 78.8472 16.5485 -84.6808 65.1172	847.8424 -5186.7589 -85.0982	-0.00206 -86.7480 -0.07908
SWCNT <sub>112</sub>	Tour 800 - 811	-11.388 -8.779	TOT 12.4020 46.8020 2.4020 46.8020 4.4080	2 4.172 2 4.172 1 -48.025 5 1.785	Qu XX XY XY 4 -701.488 22.078 2.040 6 -740.118 8.028	-20.7771  adrupole m  YY  XZ  5 -658.000  6 42.516  8 -687.581  2 -176.759  9 -692.200  6 -10.602  2 -682.009	-6.5811  ZZ  YZ  3 -656.44901  0 -4.0800  0 -881.0651  8 -0.7081  5 -0.01.8855  6.580  8 -0.00.2584	XXX XXZ 1 -1150 8402 880 9982 6021 0810 1 -057 4983 890 8204 1 81 0510 1 298 1491	87.0460 YYY XZZ 14.7805 -36.8268 -1.8798 -879.6485 -18.5840 -5.6580 2.5880	11.9591 ZZZ YZZ 41.5935 1.9714 -1480.9186 -89.4892 -8.8670	XYY  YYZ  -60,4705 8,5941 125,4008 -16,5880 58,0750 22,5080	XXY XYZ 217,5504 -72,4616 6,3168 -87,7188 -92,1804 -50,2857 -1,5866	XXXX ZZZV 9.095 -362.618 56.887	YYYY  XXYY  -1668.0311  -20204.2667  -16710.8000  -16087.05092  -16087.05092	ZZZZ XXZZ -14508-0011 -19726-7141 -28056-2840 -14802-9408 -14802-828	-5452.5271  Hawadeospoli  5XXX  VYZZ  1075.8405  -5022.0058  820.4061  -5805.8211  758.0188  -5008.5564	8XXZ XXXZ 8XXZ 8024 4047 -80146728 285 8800 -856 6283 410 3042 -865 9110	YYYX YYXZ 105.1808 58.5911 110.0792 -299.8216 45.7958 -108.0026	VYYZ ZZXY -88/2100 78/8472 16/5485 -84/6808 65/1172 16/2-6425	847.8424 -5186.7589 -85.0982	-0.00206 -86.7480 -0.07908
SWCNT <sub>112</sub> (4.7)-water	Tour 800 - 811	-11.888 -8.770 8.051	TOT 12.4020 44.4020 45.8020 46.8020 44.400	2 4.172 2 4.172 1 -48.025 5 1.785	-18.8708 Qu XX XY XY -791.458 -92.978 88 -929.968 -2.5407 86 -749.178 -8.628 -93.758 -756.758	-20.7771  adrupole m  YY  XZ  5 -658.000  6 42.516  8 -687.581  2 -176.759  9 -692.200  6 -10.602  2 -682.009	-6.5811  ZZ  YZ  3 -656.44901  0 -4.0800  0 -881.0651  8 -0.7081  5 -0.01.8855  6.580  8 -0.00.2584	XXX XXZ 1 -1150 8402 880 9982 6021 0810 1 -057 4983 890 8204 1 81 0510 1 298 1491	87.0460 YYY XZZ 14.7805 -36.8268 -1.8798 -879.6485 -18.5840 -5.6580 2.5880	11.9591  tapele monte  ZZZ  YZZ  41.5995 1.2714 -1480.0166 -89.4802 -8.5670 -14.7818	XYY  YYZ  -60,4705 8,5941  125,4008 -16,5880 56,0750 22,5080 43,8880	XXY XYZ 217,5504 -72,4616 6,3168 -87,7188 -92,1804 -50,2857 -1,5866	XXXX ZZZV 9.095 -362.618	YYYY  XXYY  -1668.0311  -20204.2667  -16710.8000  -16087.05092  -16087.05092	ZZZZ XXZZ -14508-0011 -19726-7141 -28056-2840 -14802-9408 -14802-828	-5452.5271  Helta-leogoli  SXXY  YYZZ  1075.8405  -5022.0658  820.4061  758.0688  -5098.5564  598.7184	8XXZ XXXZ 8XXZ 8024 4047 -80146728 285 8800 -856 6283 410 3042 -865 9110	YYYX YYXZ 105.1808 88.5911 110.0702 -209.8256 45.7958 -108.0026 79.8187	VYYZ ZZXY -88/21/00 78/3472 16/5485 -84/86/8 65/1172 16/26/25 0.7087	847.8424 -5186.7589 -85.0982	-0.00296
SWCNT <sub>112</sub> (4.7)-water	Tour 809 811 818 815	-11.888 -8.770 8.051	TOT 12.4020 44.4020 45.8020 46.8020 44.400	2 4.172 2 4.172 1 -48.025 5 1.785	-18.8708 Qu XX XY XY -791.458 -92.978 88 -929.968 -2.5407 86 -749.178 -8.628 -93.758 -756.758	-20.7771  adrupole m  YY  XZ  5 -658.000  6 42.516  8 -687.581  2 -176.759  9 -692.200  6 -10.602  2 -682.009	-6.5811  ZZ  YZ  3 -656.44901  0 -4.0800  0 -881.0651  6 -601.8855  6 .5805  5 -601.8858	XXX XXZ 1 -1150 8402 880 9982 6021 0810 1 -057 4983 890 8204 1 81 0510 1 298 1491	87.0460 YYY XZZ 14.7805 -36.8268 -1.8798 -879.6485 -18.5840 -5.6580 2.5880	11.9591  tapele monte  ZZZ  YZZ  41.5995 1.2714 -1480.0166 -89.4802 -8.5670 -14.7818	XYY  YYZ  -60,4705 8,5941  125,4008 -16,5880 56,0750 22,5080 43,8880	XXY XYZ 217,5504 -72,4616 6,3168 -87,7188 -92,1804 -50,2857 -1,5866	XXXX ZZZV 9.095 -362.618	YYYY  XXYY  -1668.0311  -20204.2667  -16710.8000  -16087.05092  -16087.05092	ZZZZ XXZZ -14508-0011 -19726-7141 -28056-2840 -14802-9408 -14802-828	-5452.5271  Helta-leogoli  SXXY  YYZZ  1075.8405  -5022.0658  820.4061  758.0688  -5098.5564  598.7184	8XXZ XXXZ 8XXZ 8024 4047 -80146728 285 8800 -856 6283 410 3042 -865 9110	YYYX YYXZ 105.1808 88.5911 110.0702 -209.8256 45.7958 -108.0026 79.8187	VYYZ ZZXY -88/21/00 78/3472 16/5485 -84/86/8 65/1172 16/26/25 0.7087	847.8424 -5186.7589 -85.0982	-0.00206 -86.7480 -0.07908
SWCNT <sub>112</sub>	Tour 809 811 818 815	-11.888 -8.775 -8.000	TOT 12.4020 44.4020 45.8020 46.8020 44.400	2 4171 1 -40.025 5 1.781	-18.8708 Qu XX XY XY -791.458 -92.978 88 -929.968 -2.5407 86 -749.178 -8.628 -93.758 -756.758	-90.7771 sdrupole m VY X2 5 -658.000 6 42.516 8 -687.591 2 -176.759 0 -662.220 6 -10.602 2 -682.009 7 -7.918	-5.5811  DESCRIPTION OF STREET OF ST	XXX XXZ -1150.8402 -880.9982 -621.0810 -655.4988 -891.401 -181.0810 -181.0810 -181.0810 -181.0810	87.0450 VVV XZZ 14.7805 -36.8268 -1.8798 -871.6485 -1.85810 -5.65810 28.2824	11.0501  ZZZ  YZZ  41.5005 1.2714 -1.450.056 -80.4802 -8.5485 -8.8670 -14.7818 -0.8805	XYY  YYZ  -60,4705 8,5941  125,4008 -16,5880 56,0750 22,5080 43,8880	XXY XYZ 217,5504 -72,4616 6,3158 -87,7189 -82,1804 -85,2857 -1,5896 -88,3850	XXXX ZZZV 9.0950 -382.618 56.8871	YVVV  XXVV  -15683.0818 2 -20204.28676 -18710.3000 7 -19488.7004 -18025.7085 -180870.5090 -180870.5090 -180870.5090	ZZZZ XXZZ -14508-0011 -19726-7141 -28056-2840 -14802-9408 -14802-828	-5452.5071  NXXV  YYZZ  1075.8405  -5022.0658  820.4061  -5825.2018  508.7184  -508.8518	8XXZ XXXZ 8XXZ 8024 4047 -80146728 285 8800 -856 6283 410 3042 -865 9110	YYYX YYXZ 105.1808 58.5911 110.0702 -200.8216 45.7958 -108.0026 70.8187 -84.1165	VYYZ ZZXY -88/21/00 78/3472 16/5485 -84/86/8 65/1172 16/26/25 0.7087	841.8424 -5185.7580 -65.0982 -40.8671	-0.00206 -36.7480 -0.07906 -0.08466
SWCNT <sub>112</sub>	Tour 300 811 813 815 815	-11.888 -8.775 -8.000	V TOT 12.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020 14.4020	2 2 4171 1 4 40.025	-18.8708 Qu XX XY 4 -701.488 22.078 88 -620.068 2.5410 8.628 6.580 0 -786.758	-20.7771  adrupole m  VV  XZ  5 -658.0006 6 42.516 8 -687.581 2 -176.759 0 -682.200 6 -10.602 2 -682.000 7 -7.918 8 -662.458	-5.5811  miser  ZZ  YZ  8 -656.5400  0 -881.0656  8 -2.7086  5 -661.8855  6 -668.885  6 -669.2584  8 -660.2584  0 -657.0560  0 -657.0560	XXX XXZ 1 - 1159.8402 1 - 657.4983 1 298.1401 1 18.5808 2 11.5808	87.0450 VVV XZZ 14.7805 -36.8268 -1.8798 -870.6485 -5.6580 25.2824 -7.2400	11.0501  ZZZ  VZZ  41.5005 1.0714 -14802 -8.4802 -8.8670 -14.7818 -0.8805	XYY YYZ -60,4705 8,5941 125,4008 -16,5880 56,0750 22,5080 43,8880 16,9016	XXV XVZ 217.5504 -72.4616 6.3158 -87.7189 -82.1804 -85.2857 -1.5806 -88.8850	55 55 55 55 55 55 55 55 55 55 55 55 55	YYYY  XXYY  -15031,0810  -20201,2007  -15710,8000  -16007,0509  -15007,0509  -15007,0509  -15007,0509  -15007,0509	ZZZZ  XXZZ  -14508-0011 -10726-7141 -20206-2940 -22405-5008 -14602-2881 -14602-2881 -14602-2881 -14602-2881	-5452.5071  NXXV  YYZZ  1075.8405  -5022.0658  820.4061  -5825.2018  508.7184  -508.8518	8 monissit XXXZ XXVZ 8524-4047 -850-48728 -8014-6728 -865-6918 -865-6918 -885-7629 -1298-7610	YYYX YYXZ 105.1808 58.5911 110.0702 -200.8216 45.7958 -108.0026 70.8187 -84.1165	YYYZ ZZXY -882100 78.8470 16.5485 -84.6818 65.1172 162.6425 9.7087 128.7279	841.8424 -5185.7580 -65.0982 -40.8671	-0.00206 -36.7480 -0.07906 -0.08406
SWCNT <sub>112</sub>	Tour 300 811 813 815 815	-11.888 -8.779 8.900 8.700	TOT  \$5 2.5890 12.4020 14 -0.609 40.8621 44462 16 0.4080 42046 42046	2 2 4171 1 -48,009 5 1,784 1 5 1,017	Qu XX XY XY 14-15.0 XX XY 14-701.458 22.058 88 -020.068 2.540 86 -749.178 8.028 6.580 -746.518 19.185	-20.7771  adrupole m  VY  XZ  5 -658.000 6 42.516 8 -687.581 2 -176.759 0 -682.200 6 -10.602 2 -682.000 7 -7.918 8 -662.458 1 -15.478	-5.5811  22  V2  8 -656.6402  0 -4.860  0 -881.0555  6.561.885  5 -651.885  5 -652.584  3 -2945  0 -657.854  1 0.9445	XXX XXZ 1 - 1159.8402 1 - 657.4983 1 298.1401 1 18.5808 2 11.5808	87,0460 VVV XZZ 14,7805 -36,8208 -1,8708 -870,6485 -18,5840 -5,6550 25,2824 -7,2400 26,0887	11.9501  222  Y22  41.5005 1.2714 -1480.0166 -89.4802 -8.8486 -8.8670 -14.7818 -0.8805 -5.7600	58 8010 15,2807	XXV XVZ 217.5504 -72.4616 6.3158 -87.7189 -82.1804 -85.2857 -1.5806 -88.8850	55 55 55 55 55 55 55 55 55 55 55 55 55	YYYY  XXYY  -15031,0810  -20201,2007  -15710,8000  -16007,0509  -15007,0509  -15007,0509  -15007,0509  -15007,0509	2222 XX22 -14598-9011 -28256-7143 -28256-2845 -14802-9455 -14802-853 -14802-853 -14802-853 -14802-853 -14802-853 -14802-853 -14802-853 -14802-853 -14802-853	-5452.5071  Mexadenapoli  5XXV  VV2Z  1055.8405  -5002.0658  820.4061  580.4061  580.8554  -5008.5554  -5008.8518  -5008.4805	8 monissit XXXZ XXVZ 8524-4047 -850-4878 -8014-6728 -805-6248 410-0442 -865-9110 -885-7629 -1298-7610 -97-2160	VVVX VVXZ 105.1808 88.5011 110.0702 -200.8216 45.7958 -108.0026 79.8187 -84.1165	YYYZ ZZXY -88.2100 78.8472 16.5485 -84.6608 65.1172 162.6225 9.7987 128.7279 -24.8207 171.1821	847.8424 -6186.7580 -65.0982 -40.8671 -71.3886	-0.00296 -86.7489 -0.07908 -0.08498 -0.09182
SWCNT <sub>112</sub>	Tour 309 811 818 815 817 819	-11.888 -8.779 8.900 8.700	TOT  \$5 2.5890 12.4020 14 -0.609 40.8621 44462 16 0.4080 42046 42046	2 4.171 2 4.171 1 -40.025 5 1.784 5 1.217 7 -2.151	-18.8708 Qu XX XY -701.458 92.978 92.9600 96749.178 9.029 11748.6111 19.185 16748.188	-20.7771  adrupole m  VY  XZ  5 -658.000 6 42.516 8 -687.581 2 -176.759 0 -682.200 6 -10.602 2 -682.000 7 -7.918 8 -662.458 1 -15.478	-5.5811  22  Y2  8 -656,5401  0 -4.880  0 -881,055  6 -661,255  5 -661,855  5 -661,855  1 0,5461  0 -657,854  1 0,5441  2 -660,4124	XXX XXZ -1150.8402 880.9982 621.0510 -657.4088 890.3294 181.0510 1296.1401 118.5898 -471.15090 -471.15090 -246.2160	87.0460 YYY XZZ 14.7805 -86.8268 -1.8708 -870.6485 -18.5840 -5.6580 25.3892 -7.2400 26.0887 -16.5062	11.0501  222  Y22  41.5005  1.0714  -1.480.0166  -8.4805  -8.8670  -14.7818  -8.8650  -5.7600  -5.6816	50 4705 8 5041 125 4008 -16 5880 22 5080 15 988 -17 5880 -18 5880 -18 5880 -18 580 -18 580 -18 580 -18 580 -18 580 -18 580 -18 580	XXY XYZ 217.5504 -72.4616 6.8155 -87.7189 -92.1804 -85.2857 -1.5896 -88.8850 -2.0844 -67.9627	NXXX 222V 9.005 -002.618 56.087 -003.610 8.000	YYYY  XXYY  -15085.0815 2 -202014.2007 -18710.8000 7 -10488.7004 -15007.6002 -15007.6021 -15007.6021 -15007.6021 -15007.6021	2222 XX22 -14508-3011 -10708-7191 -28056-2891 -14020-9468 -10925-2887 -14802-8728 -14802-8728 -14802-9191 -1457-0671	-5452,5071  NXXV  YYZZ  1075,8405  -5022,0058  820,4001  758,0384  508,7184  -5068,8518  1599,8402  1599,8402  -5081,8805  -1084,1805  -1084,1805	8 moniest XXXZ XXYZ 8524 4047 -880 4878 -801 4878 280 880 280 880 -805 9110 887 7629 -1298 7610 07 2160	VVVX VVXZ 105.1808 58.5911 110.0792 -290.8216 45.7958 -108.0026 79.8187 -84.1165 	VVVZ  ZZXY  -88.2100  78.8472  16.5485  -84.6808  65.1172  162.6925  9.7087  128.7279  -24.8207  171.1821  -28.9555	847.8424 -6186.7580 -65.0982 -40.8671 -71.3886	-0.00296 -86.7489 -0.07908 -0.08498 -0.09182
SWCNT <sub>112</sub> (4.7)-water	Tour 309 811 818 815 817 819	-11.886 -8.775 -9.951 -0.007	Y TOT TOT 45 2.580 14 -0.020 14 -0.020 14 -0.020 14 -0.021 14 -0.021 14 -0.027 14 -0.027 14 -0.027 15 -0.027 16 -0.027	2 2 4171 2 2 4171 1 4 -40.005 3 1.784 3 1.074 5 1.207	-18.8708 Qu XX XY -701.458 92.978 92.9600 96749.178 9.029 11748.6111 19.185 16748.188	-90,7771  VY  X2  5 -68,000  6 42,516  6 42,516  6 10,602,200  6 -10,602,200  7 -70,18  8 -602,000  1 -15,478  8 -602,553  8 -15,478	-6.5811  22  Y2  3 -656,5492  0 -4.860  0 -881,0651  5 -6.1865  5 -6.363  3 -2946  0 -657,0542  1 0.9441  2 -660,4124  6 -2.0273	XXX XXZ -1150.8402 880.9982 621.0810 -057.4988 899.8294 181.0810 1296.1401 118.5898 -271.1509 128.4000 -444.2160 8 -140.6488	87.0460 YYY XZZ 14.7805 -86.8268 -1.8708 -870.6485 -18.5840 -5.6580 25.3892 -7.2400 26.0887 -16.5062	11.9501  ZZZ  VZZ  41.5005 1.2714 -1.431.0166 -89.4802 -8.4805 -1.4.7818 -0.8805 -5.7600 -5.6816 -15.0540	50 4705 8 5041 125 4008 -16 5880 22 5080 15 988 -17 5880 -18 5880 -18 5880 -18 580 -18 580 -18 580 -18 580 -18 580 -18 580 -18 580	XXY XYZ 217.5504 -72.4616 -6.3108 -97.71804 -85.2857 -1.5896 -88.8850 -2.0844 -57.9627 -120.9415 -66.8855	XXXX ZZZV 9,005 -02,418 66,487 -48,037 -14,798	YYYY  XXYY  -15685.0311 2 -20204.2000 -18710.8000 7 -19488.7000 -18067.0200 -18067.0200 -18067.0200 -18067.0200 -18067.0200 -18067.0200 -18067.0200 -18067.0200 -18067.0200 -18067.0200	2222 XX2Z -14508-0011 -10726-7142 -28256-2845 -28455-5096 -14502-9465 -14862-8721 -10745-6229 -14582-0101 -14457-6671 -16860-1006	-5452.5071  NXXV  YYZZ  1075.8405  -5022.0058  820.4051  758.0388  -508.5084  508.7184  -508.8518  1599.8402  -508.1805  -1084.1805  -1084.1854  -5107.8802	8 monusit  XXXZ  XXVZ  8524 4047  880 4878  -8014 6728  283 5800  -805 6100  387 7029  -1208 7610  07 2160  -1152 8088  -186 1022	VVVX  VVXZ  105.1808 58.5911 110.0792 -200.8216 45.7958 -108.0026 70.8187 -84.1105 -107.2057 -181.0272 -105.8807 -188.8515	22XY -88.2100 78.8472 16.5472 16.5472 16.5425 0.7087 172.7279 24.8207 171.1821 -28.0555 -184.0081	847.8424 -5186.7589 -65.0982 -40.8671 -71.8886 -88.2708	-0.00296 -36.7489 -0.07906 -0.08496 -0.09082
SWCNT <sub>112</sub>	Tour 800 811 818 815 817 810 821	-11.886 -8.775 -9.951 -0.007	TOT TOT TOT 10.400 10.4	Z 2 4171 1 -48.029 6 1 1 5 1 1 5 1 1 1 1 1 1 1 1 1 1 1 1 1	-18.8708  Qu XX  XY  14791.438  22.978  88929.968  2.640  86748.738  6.585  11745.611  19.185  -748.808  -14.808	-20,7771  VY  X2  5 -688,600  42,516  5 -688,600  6 -22,516  5 -602,002  7 -7018  8 -602,003  8 -602,003  1 -16,205  6 -16,205  1 -16,205  5 -16,205  8 -602,003  8 -602,003  8 -602,003  8 -602,003	-6.5811  22  V2  8 -656,0402  0 -4.0800  0 -881,0651  6 -601,8851  5 -601,8851  5 -601,8851  1 -004,258  3 -204  1 -004,412  1 -004,412  2 -001,242  2 -001,242  2 -001,242  2 -001,242  2 -001,242	XXX  XXZ  -1150.8692  621.0810  621.0810  635.2041  181.2508  471.15098  471.15098  1 284.2051  1 244.2051  1 -140.4488  -585.2023	87.0460 YYY KZZ 14.7805 -86.8208 -1.8798 -870.6485 -5.6840 -5.6840 -5.6840 -5.6840 -7.2400 26.2824 -7.2400 -7.2400 -86.4025 -86.4025 -86.4025 -86.4025	12.9501  ZZZ  VZZ  41.5005 1.2714 -1.431.0166 -99.4802 -8.4865 -1.4.7818 -0.8805 -5.7600 -5.6816 -15.0540 -11.6777	XYV  VYZ  -60,4705 8,5041 125,6080 60,5080 61,	XXY XYZ 217.5504 -72.4616 -6.3108 -97.71804 -85.2857 -1.5896 -88.8850 -2.0844 -57.9627 -120.9415 -66.8855	XXXX 222V 0.005 -02.618 56.651 48.051 -12.706	XXYY -16685.0311 -20204.2667 -16710.3006 7 -19488.7004 -16055.7065 -16067.6270 -20208.1004 -15060.7850 -16162.1668 8 -20416.8786	2222 XX22 -14508-0011 -10708-7147 -28256-2845-5008 -10808-0465 -10808-2851 -14808-2851 -14808-2851 -14808-0455 -14884-0455 -14884-0455 -14884-0455	-5452.5071  Hexa-decapoli  5XXY  YYZZ  1075.8405  -5022.0658  820.4081  -5805.1221  158.0088  -5008.5564  1509.8402  -5088.184  -5088.8518  1509.8402  1509.8402  1509.8402  1509.8402  1509.8402  1509.8402  1509.8402  1509.8402  1509.8402	8 WORLD STATE OF THE STATE OF T	YYYX YYXZ 105.1805 56.5011 110.0725 -106.0025 70.5187 -84.1105 -106.0025 -106.0025 207.0856 207.0856	22XY -88.2100 78.8472 16.5472 16.5472 16.5425 0.7087 172.7279 24.8207 171.1821 -28.0555 -184.0081	847.8424 -5186.7589 -65.0982 -40.8671 -71.8886 -88.2708	-0.00296 -36.7489 -0.07906 -0.08496 -0.09082
SWCNT <sub>112</sub>	Tour 800 811 813 815 815 817 828	-11.889 -3.770 -3.951 -4.716 -5.687	TOT TOT 12 -0.007 14 -0.003 15 -0.003 16 -0.003 17 -0.003 18 -0.00	Z 2 4171 1 -48.029 6 1 1 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	-18.8708 Q0 XX XY 4 -701.488 22.078 8 -629.068 2.640 60 -789.128 6.585 71 -745.6111 19.18.58 -14.870 55 -749.580	-20,7771  VY  X2  5 -68,8000  42,516  6 42,516  6 42,516  7 -70,86  6 -10,700  7 -70,86  8 -602,200  7 -70,86  8 -602,500  8 -10,700  8 -10,700  8 -10,700  8 -10,700  1 -10,700	-5.5811  mused  ZZ  YZ  8 -656,8400  0 -881,0656  8 -27,0806  5 -661,8851  5 -661,8851  6 -662,053  0 -657,8544  1 -064,412,6  6 -2,073  2 -661,2402  7 -7,8408	XXX  XXZ  -1150.8692  621.0810 -657.4888 396.3294 128.13510 1298.1491 118.3698 471.1500 1-244.2690 1-240.6488 1-385.2628 1-385.2628 1-385.2628	87.0460 YYY XZZ 14.7805 -80.8208 -1.8708 -870.6485 -1.8708 -5.6580 23.2824 -7.2400 26.8802 -36.4025 -5.6700 -27.4808	12.9501  ZZZ  VZZ  41.5905 1.2714 -2.480.0165 -9.4802 -8.8605 -14.7818 -0.8805 -5.7600 -5.6816 -15.2540 -11.9677 -4.4880	XYY  YYZ  -00 4705 85041 105 4000 50 0750 50 0750 63 8800 15 2001 65 88 8010 15 2007 66 7106 11 3174	XXV XYZ 217.5004 -72.4016 -63.158 -67.7188 -68.3800 -28.8800 -72.0844 -67.0027 -70.00415 -70.00415 -70.00415 -70.00415 -70.00415	XXXX ZZZY 0.002 0.002 0.002 0.003 48.007 48.007 -14.700	XXYY -15635 0815 -20209 2667 -16710 3000 -16710 3000 -16625 7685 -20870 5690 -15005 7650 -20880 1694 -15005 7650 -20880 1694 -15005 7650 -20880 1694	2222 XX22 -14508-9011 -10708-7142 -22405-5098 -1020-9465- -1020-9465- -1020-9465- -10705-10705- -14882-9191 -19708-1106- -14882-9191 -14	-5452.5071  Hexade expol  SXXV  YYZZ  1975.8405  -5022.0658  820.4081  -5826.4081  1580.088  -5088.5564  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402  1590.8402	8 WORLD STATE OF THE STATE OF T	YYYX YYXZ 105.1808 56.5011 11100792 45.7063 45.7063 70.918 94.1165 107.2067	VYVZ ZZXY -88 2100 16 5487 -81 600 65 1172 160 6425 0 1087 -71 1821 -28 6555 -64 600 158 6765	367.8626 -5186.7580 -65.0982 -40.8671 -71.8886 -83.2708 05.1824	-0.00296 -36.7489 -0.07906 -0.08496 -0.09082
SWCNT <sub>112</sub> (4.7)-water	Tour 800 811 813 815 815 817 828	-11.889 -3.770 -3.951 -4.716 -5.687	TOT TOT 12 -0.007 14 -0.003 15 -0.003 16 -0.003 17 -0.003 18 -0.00	2 4171 2 4171 1 40.025 1 1.784	Qu XX XY 4 -701.488 22.978 88 -629.968 2.540 66 -749.178 8.628 61 -745.611 1745.611 1718.65 65 -748.168 -74.808 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268	-20,7771  VY  X2  5 -68,8000  42,516  6 42,516  6 42,516  7 -70,86  6 -10,700  7 -70,86  8 -602,200  7 -70,86  8 -602,500  8 -10,700  8 -10,700  8 -10,700  8 -10,700  1 -10,700	-5.5811  OUISM!  ZZ  VZ  8 -656.5400  0 -881.0558  -2.7081  5 -661.8855  6 .5860  2 -860.2584  1 -0.9441  2 -660.41246  6 -2.0273  2 -661.2207  7 -7.3408  8 -661.8015	XXX XXZ 2 -1150 8602 621 0810 1657 4083 396 3294 181 0510 129 1491 118 5609 1471 1509 128 4609 1471 1508 151 1510 158 1609 158 1609	87,0460 VVV XZZ 14,7805 -36,8208 -1,8798 -870,6485 -13,5840 -5,6580 25,2824 -7,2400 20,0887 -36,4002 -86,4002 -86,4002 -5,6700 -27,4808 -0,0721	12.9501  ZZZ  YZZ  41.5905 1.2714 -2.480.0165 -90.4802 -8.8505 -14.7818 -0.8805 -5.7600 -5.6825 -14.7828 -14.7828 -14.7828 -14.7828 -14.7828 -14.7828	XYY  YYZ  -0.14705 8.5041 103.0000 20.5000 24.8880 10.9016 -8.7017 -8.7017 11.3174 -98.3470	XXV XYZ 217.5504 6.3108	NXXX 222V 9.005 9.005 56.841 48.035 -14.708 -86.117	VYVV  XXYV  -15635.0315  -20204.2667  -15710.3000  -15025.7687  -20028.1694  -15050.7850  -15050	2222 XX22 -14508-9011 -10706-7141 -20056-20405-5098 -14000-9465 -10905-2687 -14802-8701 -10760-1100 -14805-6071 -14805-8005 -14885-80101 -14885-80101 -14885-80101 -14885-80101 -14885-80101 -14885-80101 -14885-80101 -14885-80101	-5452.5071  Mexade sapoli  5XXV  VVZZ  1075.8405  -5022.0658  820.4061  -5825.4271  758.0088  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564  -5008.5564	8 monunit XXXZ XXYZ 8524 4047 -850 4878 -801 4578 288 8800 -865 9110 687 7629 -1298 7610 07 2169 -1158 1088 -167 0487 -508 8721 -1407 0000	YYYX YYXZ 105.1808 56.5011 11100792 45.7063 45.7063 70.918 94.1165 107.2067	VYVZ ZZXY -88 2100 16 64672 16 64672 16 64672 16 7067 17 1821 24 8207 17 1821 15 6766 -17 7842	367.8626 -5186.7580 -65.0982 -40.8671 -71.8886 -83.2708 05.1824	-0.0206 -86.7489 -0.07908 -0.08466 -0.09182 -0.04833 -0.05100
SWCNT <sub>112</sub> (4.7)-water	Tour 800 811 813 815 815 817 828	-11.388 -8.776 -9.001 -5.007 -5.889	V TOT TOT \$6 2.580.0 \$12.400 \$4 4.0.600 \$48.800 \$2.580.0 \$4.0.600 \$4.000	2 4.75 2 4.75 3 4-40.025 5 1.784 7 -2.15(-1.77) 7 -2.15(-1.77) 8 0.742	Qu XX XY 4 -701.488 22.978 88 -629.968 2.540 66 -749.178 8.628 61 -745.611 1745.611 1718.65 65 -748.168 -74.808 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268 65 -749.268	- 20,7771  VV  X2  X3  5 - 688,000  6 22,516  6 22,516  6 - 100,002  7 - 7 018  8 - 100,002  1 1 - 15,478  8 - 100,250  5 - 100,002  1 2 - 100,002  2 - 100,002  2 - 100,002  2 - 100,002  2 - 100,002  2 - 100,002  2 - 100,002  3 - 10,0	-6.5811  22  Y2  3 -650,4401  0 -4.0800  0 -881,0558  6 -051,8555  5 -651,8551  1 0,0441  2 -650,4124  6 -2.0273  7 -7,840  8 -661,8501  5 -661,8501  5 -661,8501  5 -661,8501	XXX  XXZ  -1150.8462  530.9982  6021.9810  -051.4983  181.9510  198.4900  118.5998  471.1509  128.4900  1-444.2160  -144.2160  1-58.2083  -158.5028  -68.5028  -68.8866  -68.8866  -68.8866  -68.8866  -68.8866  -68.8866	37.0460 VVV XZZ 14.7805 -80.8208 -1.8788 -1.8580 -25.880 25.880 -7.2400 26.0887 -16.5002 -81.425 5.8700 -27.4808 -0.0721 -7.4980	11.9501  tapole manuse  ZZZ  YZZ  41.5005 1.2714 -1480.0166 -80.4802 -8.5486 -8.670 -14.7818 -0.8805 -5.6816 -15.040 -11.5077 -2.4880 14.72769 -20.1092	XYY  YYZ  -00 4705 85041 925,4006 60,0700 62,5880 68,8010 68,8010 11,2607 11,416877 -00,7700 11,4176 -08,1870 64,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252 66,7252	XXV XV2 217.5504 63.158 63	XXXX ZZZY 0.00526 -02.618 56.685 -14.708 -86.177 -9.589	VYVV  XXYY  -15035.0810 2 -202014.2007 -18710.8000 7 -10488.7004 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15005.0502 -15006.0502 -15006.0502 -15006.0502	2222 XX22 -14508-9017 -19726-71-91 -28256-2845-29265-5098-14892-91992-9886-11-91766-1107-110766-110768-10768-10768-10768-10768-70768	-5452.5071  NXXV  YYZZ  1075.8405  -5022.0058  820.4061  -580.5291  758.088  -508.5518  1599.3402  -508.154  -508.8518  1599.3402  -508.184  -5107.8802  1941.1480  -508.6996  -1888.6996  -1888.6996  -1888.6996  -1888.6996	8 monuelit  XXXZ  XXVZ  8524 4047  -850 4878  -8614 6728  288 5800  -865 6948  -100 462  -865 911  07 2160  -1152 8088  -106 1022  107 2087  -508 8721  -1207 5090  -1208 5051	YYYX  YYXZ  105.1809 58.5011 1100792 45.7056 -108.0026 -108.0026 -107.2057 -107.2057 -107.2057 -107.2056 181.5057 -188.2105 207.0056	VYYZ ZZXY -882200 78.6172 16.5485 -84.6088 65.1172 162.6625 27.827 24.8207 171.1871 -28.6555 -184.6081 -76.4001 158.6755 67.2078	367.8626 -5186.7580 -65.0982 -40.8671 -71.8886 -83.2708 05.1824	-0.00206 -00.7480 -0.07008 -0.08406 -0.09180 -0.08400

Table 5. The calculated dipole moment, quadrupole moment, octapole moment, and hexadecapole moment values of SWCNT

### 4. NMR and IR theoretical study on the interaction of doping metal with carbon nanotube (CNT)

Numerous electrical measurements on SWCNT ensembles have revealed that chemical doping by donors (Li, K, Cs, or Rb) or acceptors (Br<sub>2</sub>, I<sub>2</sub>, or acids) decreases the room temperature electrical resistance by up to two orders of magnitude at saturation doping (Kaaoui et al., 1999; Coluci et al. 2006). The important problem is metals passing through cells membrane. Because, there are barriers for them passing through protein canals in cells membrane. Additionally, upon interaction, changes in activity, stability, and solubility ions compatibility may occur in cells. A lot of studies are for replacing protein canals into cells membrane for passing proteins, drug and ions of metal. Therefore the presence of the SWCNT and its consequences to the biological activity of ions metal are of high impact in

the development of biosensors, immunoassays and drug delivery systems (Zhang et al., 2005; Ganjali et al., 2006). This work, we used armchair carbon nanotube (5, 5) and (6, 6). Indeed, vibrational frequencies of finite-length carbon nanotubes were recently examined (Tagmatarchris & Prato, 2004) and another result of 319.9 cm<sup>-1</sup> is consistent with oscillations along the radial directions (radial modes), although it cannot be assessed accurately due to the sensitivity to the number of rings (Yumura et al., 2005). We suggest that SWCNT intercalate into cells membrane replacing protein canals and are studying passing metal ions (Na, Mg, Al, and Si) in length of SWCNT by Quantum Mechanics (QM).

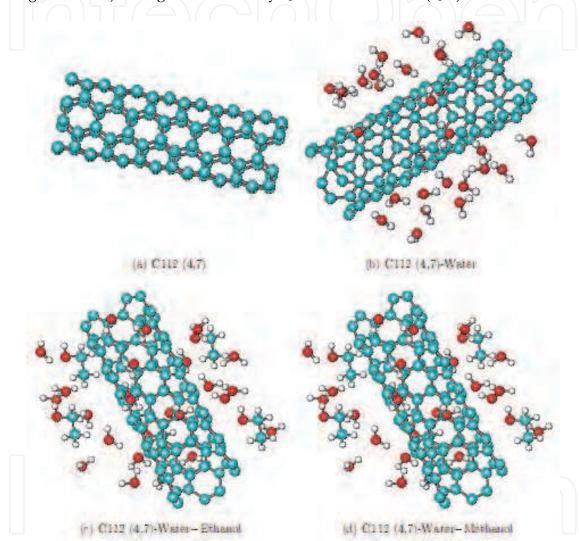


Fig. 9. Optimized structures of nanotube in different media

#### 4.1 Computational details

The geometry optimizations were performed using an all-electron linear combination of atomic orbitals Hatree–Fock (HF) and density functional theory (DFT) calculations using the Gaussian A7 package. SWCNTs (100–120) from kind of armchair carbon nanotubes (5, 5) and (6, 6) show in Fig. 10. We are interested in the structural features of single-walled carbon nanotube (SWCNT) in the ground state an atomic and amino acids (His and Ser). In HF theory the energy has from:

$$E_{ks}=v++1/2-1/2$$

where v is the nuclear repulsion energy,  $\rho$  is the density matrix,  $\langle hp \rangle$  is the one electron (kinetic plus potential energy).  $1/2 \langle P_j(\rho) \rangle$  is the classical coulomb repulsion of the electrons and  $-1/2 \langle P_k(\rho) \rangle$  is the exchange energy resulting from the quantum (fermions) nature of electrons.

In density function theory the exact exchange (HF) for a single determinant is replaced by a more general expression the exchange correlation functional, which can include terms accounting for both exchange energy and the electron correlation, which is omitted from Hartree–Fock theory:

$$E_{ks}=v++1/2+E_{\chi(\rho)}+E_{C(\rho)}$$

where,  $E_{\chi(\rho)}$  is the exchange function and  $E_{C(\rho)}$  is the correlation functional. The correlation function of Lee, Yang, and Parr is includes both local and non-local term (Kar et al., 2006). The optimizations of solids are carried out including exchange and correlation contributions using Becks three parameters hybrid and Lee–Yang–Parr (LYP) correlation [B3LYP]; including both local and non-local terms with the program Gaussian A7 package (Lee et al., 1988; Becke, 1993; Becke, 1997).

Compared to Raman spectroscopy, much less information about the vibration properties of carbon nanotubes can be gained from IR spectra. This limitation mainly results from the strong absorption of SWCNTs in the IR range. Accurate predictions of molecular response properties to external fields are of general significance in various areas of chemical physics. This especially refers to the second-order magnetic response properties (NMR), since the magnetic resonance based techniques have gained substantial importance in chemistry and biochemistry that NMR data shown with two parameters isotropic ( $\sigma_{iso}$ ) and an isotropic ( $\sigma_{aniso}$ ) shielding.

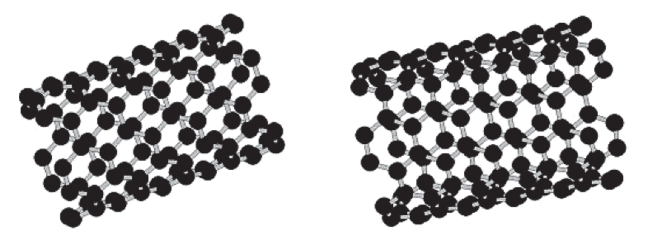


Fig. 10. The optimized configuration Side-view SWCNT:  $C_{100}$  (a) and  $C_{120}$  (b)

#### 4.2 Interaction of Na, Mg, Al, Si with Carbon Nanotube (CNT): NMR and IR Study

The B3LYP and HF by 6–31G and 6–31G\* calculaion for the molecular SWCNT models with Na, Mg, Al, and Si considered were validated by the calculated 13C and 1H NMR shifts and thermodynamic properties of an open-ended SWCNT (5, 5) and (6, 6) molecular systems (Monajjemi et al., 2009). The total energy (Etotal) of this interaction is listed in Table 6, which the Etotal increase to converge with an increasing carbon number.

Ene	rgy total		N	a		N	1g	I	XI	5	ŝi
(H	artree)	HF	B3LYP	HF	B3LYP	HF	B3LYP	HF	B3LYP	HF	B3LYP
C <sub>100</sub>	6-31G	-2241.98	-2256.53	-2401.83	-2416,63	-2438.87	-2453,67	-2480,80	-2495,72	-2527.27	-2542.30
(5.5)	6-31G*	-2269,16	-2284.21	-2430.75	-2446.16	-2468.15	-2483,58	-2510,66	-2526,16	-2557,67	-2573,29
C <sub>120</sub>	6-31G	-2990,32	-3009,391	-3150,15	-3169,61	-3187,08	-3206,55	-3229,18	-3248,72	-3275,58	-3295.37
(6,6)	6-31G*	-3026,37	-3046.26	-3188.02	-3208.34	-3225,34	-3245,70	-3267.95	-3288.37	-3315,00	-3335.57

Table 6. The total energy calculated in various basis set at HF & B3LYP for SWCNTs (5,5) and (6,6) with ions metal Na, Mg, Al and Si

In this study the metals on the center of a hexagon (HC) and muse are related to competitive interactions between ions metal and SWCNTs. The structural electronic and magnetic properties have been investigated. The most stable configuration for Si adsorbed on SWCNTs is also at the (HC) site at competitive another atoms of SWCNT because the electro negativity is the most great. The calculated amounts of Dipole, Quadrupole, Octapole, and Hexa-decapole moments at the HF and B3LYP levels in various basis set are given in Table 7. Hybridizing Coefficient is different in various methods and basis set.

Calculations of the NMR shifts with the magnetic field perturbation method of GIAO (gauge in dependent atomic orbital) incorporated with the program Gaussian A7 package. The results of the calculations for the carbon nearest neighbors' atoms in SWCNTs are presented in Table 7. The calculated magnetic shielding in Figs. 11, 12 was converted into  $\sigma_{iso}$ ,  $\sigma_{aniso}$  chemical shifts by 13C absolute shielding in SWCNT (5, 5). They are worth noting that the last approach leads to a substantial improvement in the calculated magnetic properties. Regarding the method for achievement of gauge invariance for the present case, at the B3LYP and HF levels on the other hand at the hybrid B3LYP level, GIAO is found to be slightly superior. The calculated infrared is for C100 at HF/6-31G with Na, Mg, Al, and Si. They showed in Table 2. The properties thermodynamic are decrease with increase electro negativity atoms.

	Cv <sub>th</sub> (cal/mol)	S <sub>th</sub> (cal/mol)	H <sub>th</sub> (cal/mol)	Gth (cal/mol)	Eth (cal/mol)
SWCNT <sub>100</sub> (5.5)	104.25	146.61	245,64	201.93	245.05
SWCNT <sub>100</sub> Na	103,13	139,32	262,12	220.60	261.68
SWCNT <sub>100</sub> -Mg	103,44	138.53	255,49	214.21	255.04
SWCNT <sub>100</sub> -Al	105,85	142.81	247.06	204.50	246,61
SWCNT <sub>100</sub> -Si	105,58	143,55	243,11	200,33	242.65

Table 7. Calculated thermal energy, thermal enthalpy, total enthalpy, thermal entropy, thermal Gibbs free energy, Gibbs free energy, and heat capacity by IR-HF/6-31G

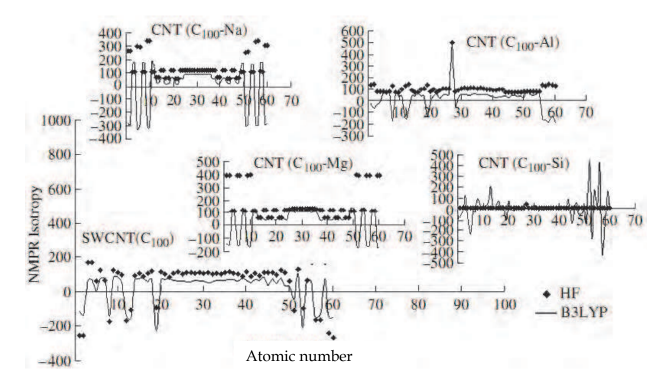


Fig. 11. NMR isotropy diagrams of SWCNT ( $C_{100}$ ) for HF/6–31G ( $\spadesuit$ ) and BLYP/6–31G ( $\neg$ ) method

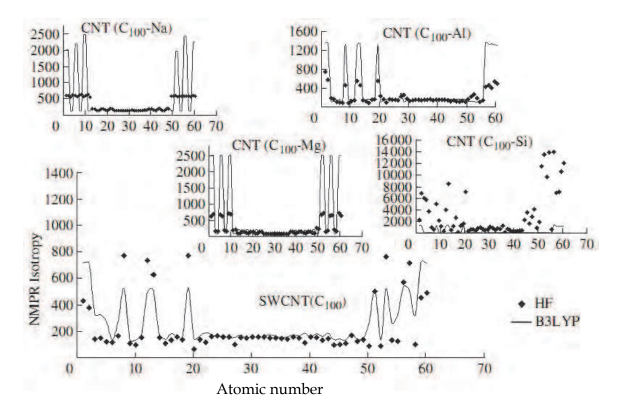


Fig. 12. NMR anisotropy diagrams of SWCNT (C100) for HF/6–31G ( $\spadesuit$ ) and BLYP/6–31G ( $\lnot$ ) method

#### 5. Conclusion

Carbon Nanotubes have been intensively studied due to their importance as building block in nanotechnology. The special geometry and unique properties of carbon Nanotube offer great potential applications, including Nanoelectronic devices, energy storage, gas sensing, chemical probe, electron transport, and biosensors, field emission display, etc. Such devices operate typically on the changes of electrical response characteristics of the Nanowire active component with the application of an externally applied mechanical stress or the adsorption of chemical or bio-molecule. For a better understanding of the physical and electronic properties of single-walled carbon Nanotubes (SWCNTs) at the Nano scale, a challenging task in theoretical calculation is needed in order to design the specific material properties because of the large size of the SWCNTs and their complicated and size dependent electronic structure. Modeling of functionalized Nanotubes and nanostructures for such technologies of SWCNTs can be greatly benefit from the first principles methods based on the density functional theory (DFT). The equilibrium position, adsorption energy, binding energy, charge transfer, and electronic band structures can be computed for different kinds of SWNTs. Effects of surrounding medium and intrinsic structural defects can also be taken into account. In this work we review some recent DFT investigation on the gas-sensing properties and the dielectric properties. Charge transfer and gas-induced charge fluctuation might significantly affect the transport properties of SWNTs. The size and chirality's of the carbon Nanotubes were typical determined from the SWCNT Raman energy spectra of a peak around 150-300 cm<sup>-1</sup>, due to the radial breathing mode. Besides, the geometry and electrical properties of Nanotube are very sensitive to dielectric constants which we can observe from the normal mode analysis. A calculation method for identifying the Raman modes of SWCNTs based on the symmetry of the vibration modes has been discussed. The Raman intensity of each vibration mode varies with polarization direction, and the relationship can be expressed as analytical functions. Each Raman active mode of SWCNT can be distinguished from the group theory principle.

In section 2, with the calculation of the normal modes using the U Matrix it is possible to get the F Matrix from the multiplication of frequency to the U Matrix. Solving the determination of F Matrix versus dielectric can be useful for understanding of the electrical behavior of nanotubes in the quantitative structure activity relationship studies. The geometry and electrical properties of nanotube are very sensitive to dielectric constants. The normal modes also will be changed in the high dielectric constants.

In section 3, *Ab initio* calculations were carried out with GAUSSIAN 98 program at the HF/3-21G level of theory to investigate the effects of polar solvents and different temperatures on the stability of SWCNT in various solvents. The results obtained from Onsager model calculations are illustrated using the energy difference between these conformers which are quite sensitive to the polarity of the surrounding solvent, that the water and methanol solvents can be suggested as the most compatible solvent for studying the structural properties of SWCNT. Also orientation of the water molecules at the CNT-water interface can be affected by the orientation of the water dipole moment. Moreover, among the energy values obtained from different MM+, AMBER, and BIO+ force fields, the AMBER force field is the most proper force field for studying SWCNT.

In section 4, A Quantum Mechanics (QM) is used for investigated the nature of metals transport and interaction with single-walled carbon nanotubes (SWCNTs) inter membranes. Metal species can be transported actively by a combination of SWCNT-membranes

conducting channels that have been used for bio-molecular and detection. Ab initio calculations using DFT/B3LYP and HF levels with 6-31G and 6-31G\* basis set of theory have allowed the determination of structure electronic, properties thermodynamic, magnetic properties for SWCNTs with Na, Mg, Al, and Si. NMR chemical shielding tensors in the methods framework makes it possible to study the chemical shift of specific group in carbon nanotubes in absence and presence metals. A comprehensive on effects of atoms on SWCNTs were revealed that it is on its electronic structure: 1) transfer of charge from the atom to the SWCNTs; 2) electrostatic interactions between the delocalized e electrons of the SWCNTs and atoms. The basis set used 6-31G and 6-31G\* that increasing electronegativity metals increased the total energy. The proportion SWCNTs were changed by them. The results are presented for T = 310 K, the temperature of human's body. In fact, it was determined that SWCNT blocked potassium channels in a dose-dependent manner. Fullerenes were discovered to be less effective channel Blockers than CNT. The mechanism was solely dependent on the size and shape of the nano-particles. They also concluded that electrochemical interactions are between CNT and the ion channels.

#### 6. Acknowledgment

The work has been supported by Thailand Research Fund (TRF), Thailand Center of Excellence in Physics (ThEP), Center for Innovation in Chemistry (PERCH-CIC), and the National Research University Project under Thailand's Office of the Higher Education Commission, Thailand for financial support.

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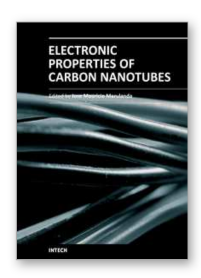
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#### **Electronic Properties of Carbon Nanotubes**

Edited by Prof. Jose Mauricio Marulanda

ISBN 978-953-307-499-3 Hard cover, 680 pages Publisher InTech Published online 27, July, 2011 Published in print edition July, 2011

Carbon nanotubes (CNTs), discovered in 1991, have been a subject of intensive research for a wide range of applications. These one-dimensional (1D) graphene sheets rolled into a tubular form have been the target of many researchers around the world. This book concentrates on the semiconductor physics of carbon nanotubes, it brings unique insight into the phenomena encountered in the electronic structure when operating with carbon nanotubes. This book also presents to reader useful information on the fabrication and applications of these outstanding materials. The main objective of this book is to give in-depth understanding of the physics and electronic structure of carbon nanotubes. Readers of this book should have a strong background on physical electronics and semiconductor device physics. This book first discusses fabrication techniques followed by an analysis on the physical properties of carbon nanotubes, including density of states and electronic structures. Ultimately, the book pursues a significant amount of work in the industry applications of carbon nanotubes.

#### How to reference

In order to correctly reference this scholarly work, feel free to copy and paste the following:

Majid Monajjemi and Vannajan Sanghiran Lee (2011). Quantum Calculation in the Prediction of the Properties of Single-Walled Carbon Nanotubes (SWNTs) and Nanotube Bundles, Electronic Properties of Carbon Nanotubes, Prof. Jose Mauricio Marulanda (Ed.), ISBN: 978-953-307-499-3, InTech, Available from: http://www.intechopen.com/books/electronic-properties-of-carbon-nanotubes/quantum-calculation-in-the-prediction-of-the-properties-of-single-walled-carbon-nanotubes-swnts-and-



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