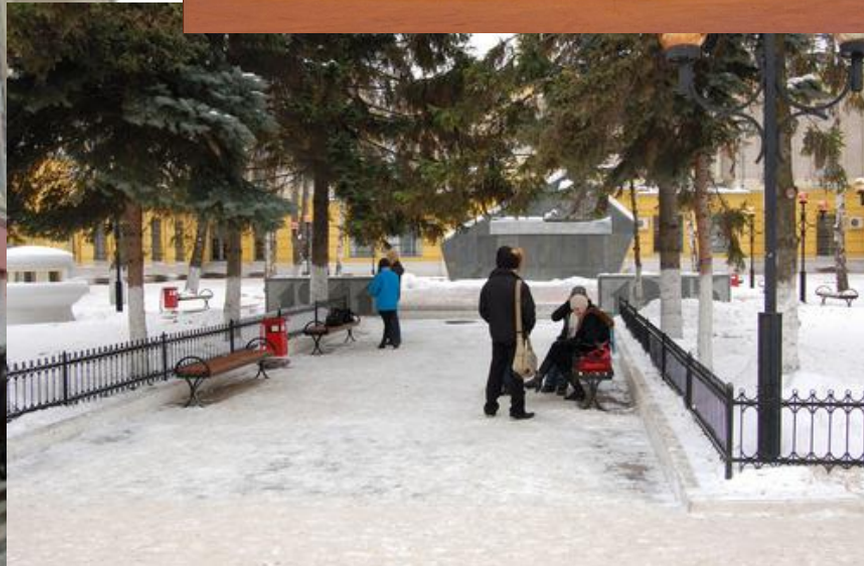


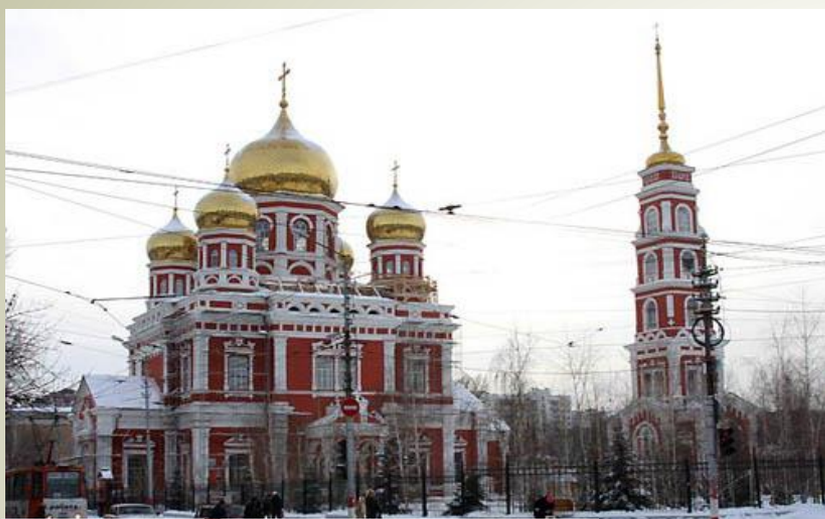
The development of methods of manipulation of behavior and properties of the molecules on the graphene

SCC DFTB MD, TBMD, REBO, AMBER, MARTINI and PM(3,6,7), ..
Developed Open-source software (OSS) "KVAZAR"

Prof. Dr. Olga E. Glukhova,
Head of Chair of Radiotechnology and Electrodynamics,
Head of Department of mathematical modeling of
Institute of nanostructures and biosystems
National Saratov State University, Russia
glukhovaoe@info.sgu.ru

Saratov State University, www.sgu.ru





**Radiotechnology and
Electrodynamics**

Modeling of nanodevices,
based on carbon nanoclusters,
nanoelectronics,
nanobiosystems mechanics,
molecular electronics,
mathematical modeling of
physical processes



Professor Glukhova

Theoretical and
applied
electrodynamics of
micro- and
extremely high wave
frequencies



Professor Davidovich

Radiotechnical research
methods of superconductors
as the objects for recording,
storage and processing of
information and
optimization of
transformator chains for
powerful impulse generators



Professor Saliy

3D displays,
mathematical
logic methods,
mathematical
modeling in
biology and
medicine



Professor Ten



Department of
Computer Simulations

High-Performance
Computing Division

Parallel Computing
Algorithms
Supercomputers
maintenance
Databases
construction and
maintenance

Mathematical
Modeling Division

FEM Modeling :
Biomechanics;
Construction
mechanics;
Solid structures
mechanics;
Structural mechanics;
Composite mechanics

Mechanics of
Nanostructures :
Mechanical properties
of nanostructures;
Mechanical properties
of bionanoobjects;
Multiscale modeling

Nanoelectronics:
Electronic
structure;
Emission
properties;
Electronic
conductivity



MECHANICAL PROPERTIES OF GRAPHENE

The local stress field of the atomic grid of nanostructures: original method

Olga Glukhova and Michael Slepchenkov // *Nanoscale*, 2012, 4, 3335–3344

It is proposed to carry out the calculation of the local stress field according to the following algorithm.

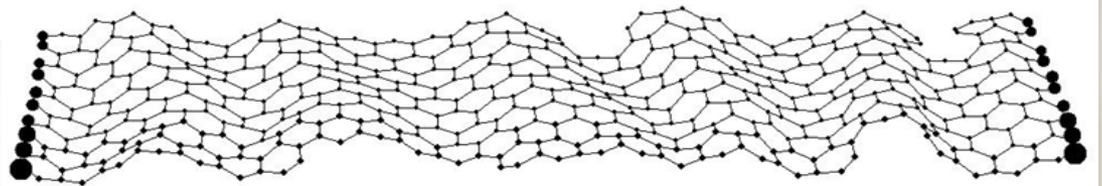
(1) Optimization of the initial structure by means of the quantum-chemical method.

(2) Calculation of distribution of the bulk energy density per atom using the empirical method.

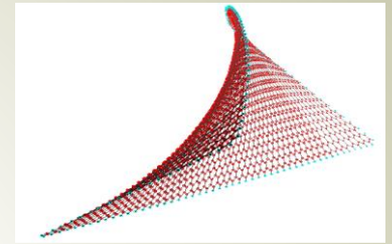
(3) Search of the atomic configuration of the nanostructure subjected to the external influence as a result of the energy minimization for coordinates, using the quantum-chemical method.

(4) Calculation of the distribution of the bulk energy density per atom in the structure subjected to the external influence, using the empirical method.

(5) Calculation of the local stress in the atomic grid according to the difference between the values of the bulk density of energy for the atoms of the structure subjected to the external influence, and the initial structure.



MECHANICAL PROPERTIES OF GRAPHENE



Study of deformations and elastic properties of nanoparticles and nanoribbons was implemented on the following algorithm

- 1) Optimization of atomic structure by entire system energy minimization on atomic coordinates (the atomic structure obtained from previous optimization);
- 2) Tension or compression of the atomic network of nanoribbon and reoptimization of atomic structure with fixed atoms on the nanoribbon ends;
- 3) Calculation the Young's pseudo-modulus for the elastic tension of nanoribbon on 1% on formula:

$$Y_p = \frac{F}{D} \frac{L}{\Delta L}$$

where a deformation force is given by $F = \frac{2\Delta E}{\Delta L}$. Here ΔE is the strain energy, namely, the total energy at a given axial strain minus the total.

- 4) Calculation the Young's modulus for the elastic tension of nanoribbon on 1% on formula:

$$Y = \frac{F}{S} \frac{L}{\Delta L}$$

The bulk energy density w_i of the atom i was calculated by the formula:

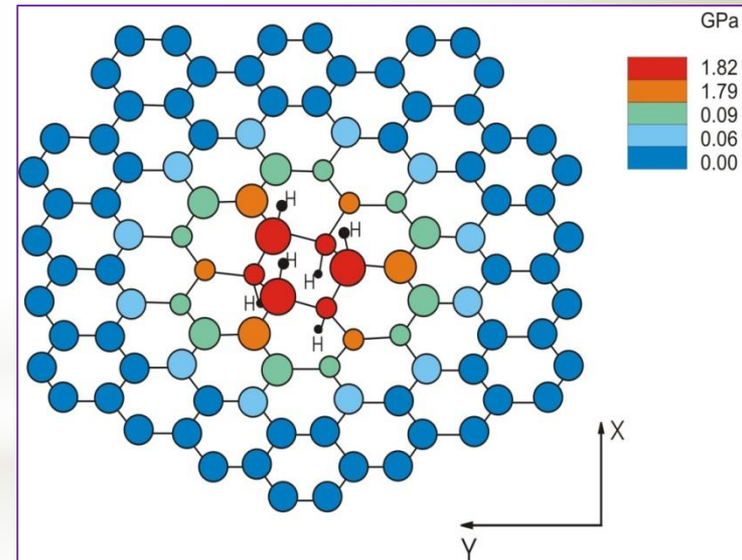
$$w_i = \left(\sum_{j(\neq i)} (V_R(r_{ij}) - B_{ij}V_A(r_{ij})) + \sum_{j(\neq i)} \left(\sum_{k(\neq ij)} \left(\sum_{l(\neq ij,k)} V_{\text{tros}}(\omega_{ijkl}) \right) \right) + \sum_{j(\neq i)} V_{\text{vdW}}(r_{ij}) \right) / V_i$$

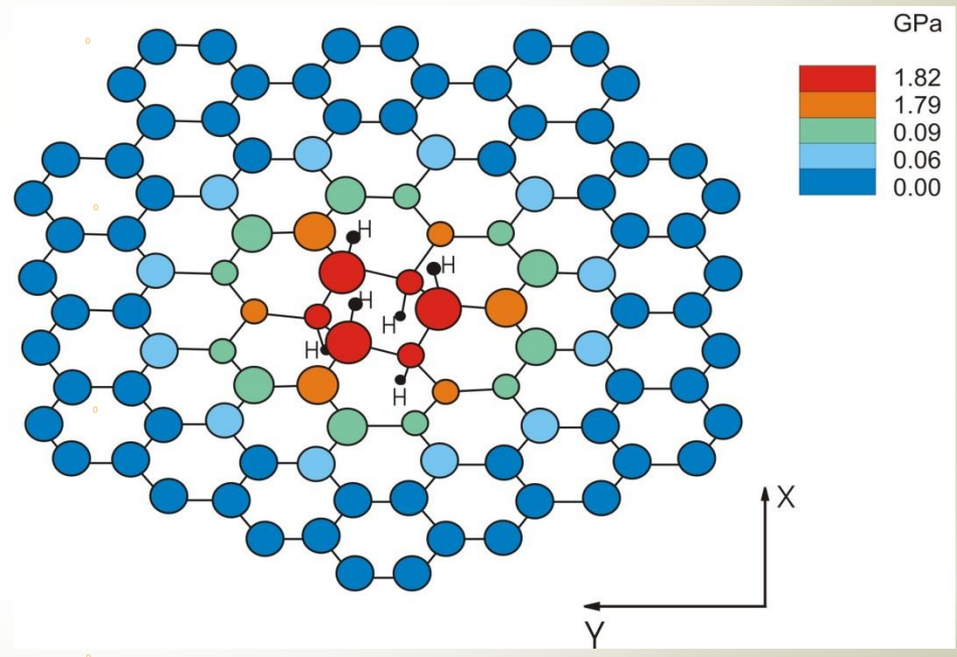
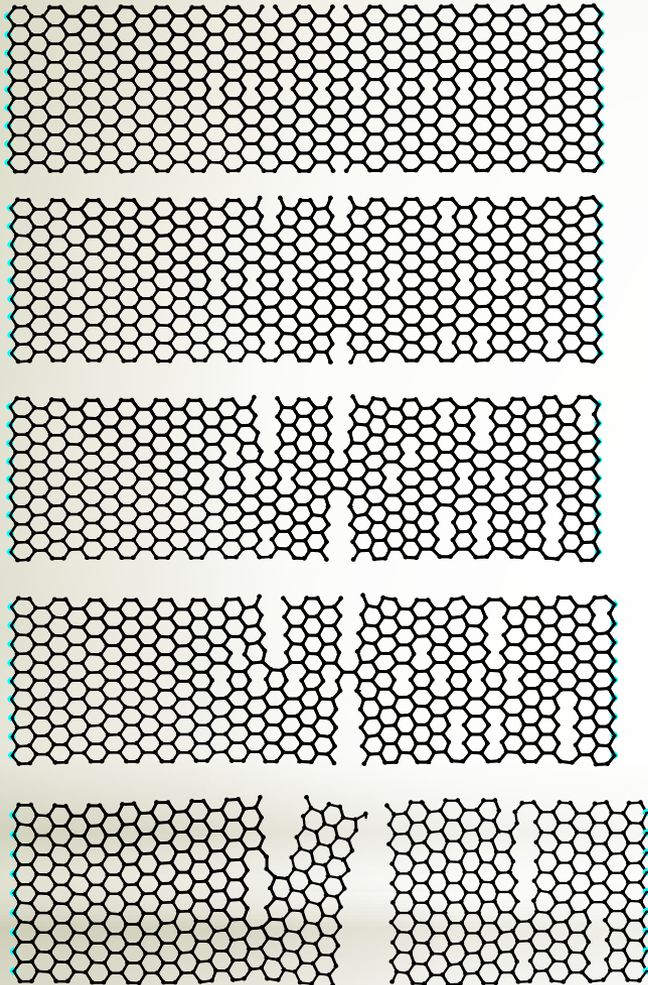
$$V_i = \frac{4}{3} \pi r_0^3$$

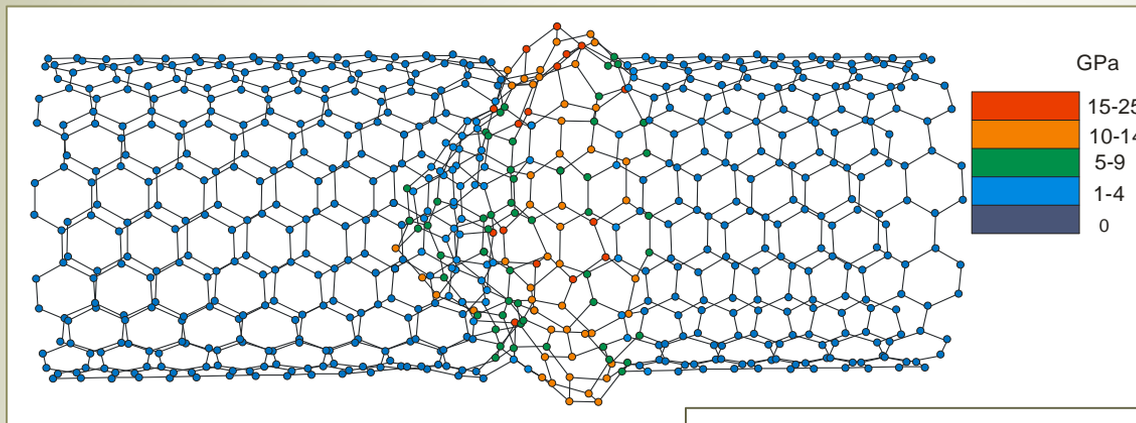
The stress of the atomic grid near the atom with number i is calculated as:

$$\sigma_i = |w_i - w_i^0|$$

where w_i^0 is the bulk energy density of the i^{th} atom of the graphene sheet which is in equilibrium; w_i is the bulk energy density of an atom of the graphene sheet subjected to the external influence (deformation, defect formation, *etc.*). The value of w_i^0 in the centre of the graphene sheet is equal to -58.60 GPa. At the edges of the graphene sheet the bulk energy density is higher since the atoms of the edges have only two links with other carbon atoms. It is equal to -41.54 GPa on an armchair edge and on the zigzag edge is equal to -40.64 GPa. It is suggested that without an external influence the stress equals to zero for the atoms in the centre and at the edges of the graphene sheet.



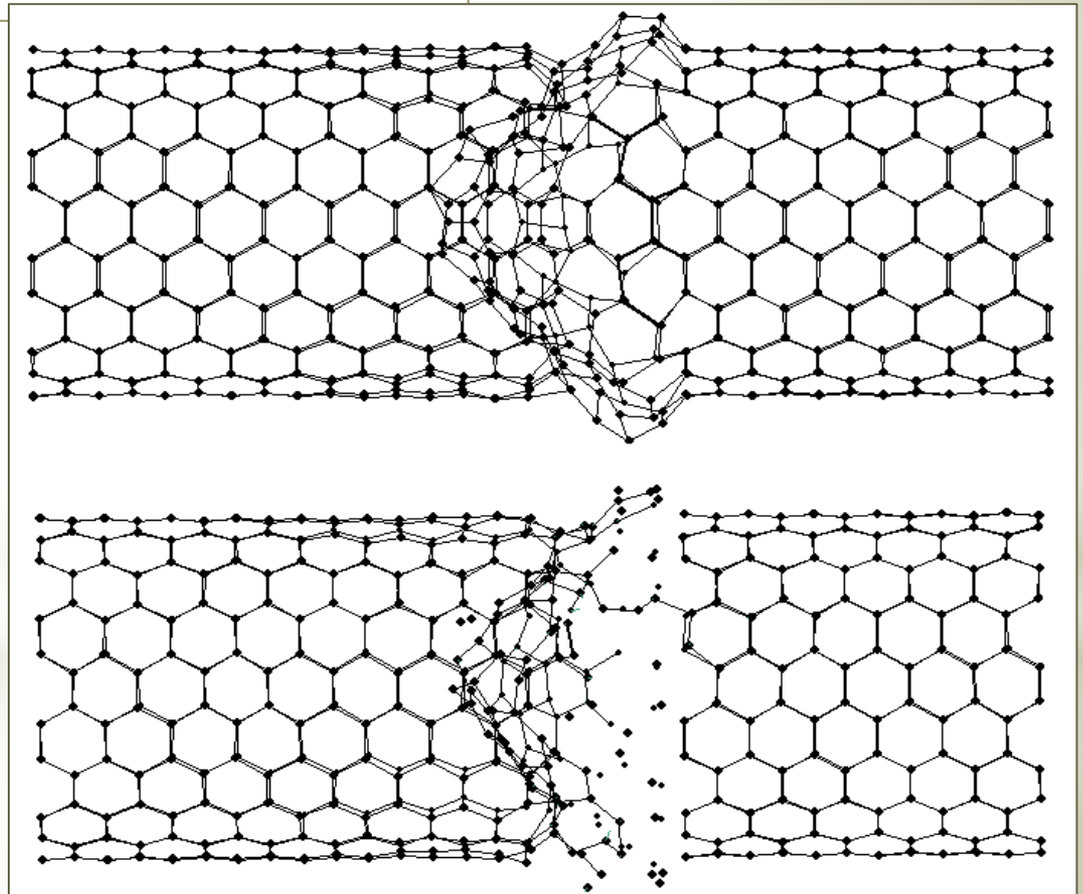


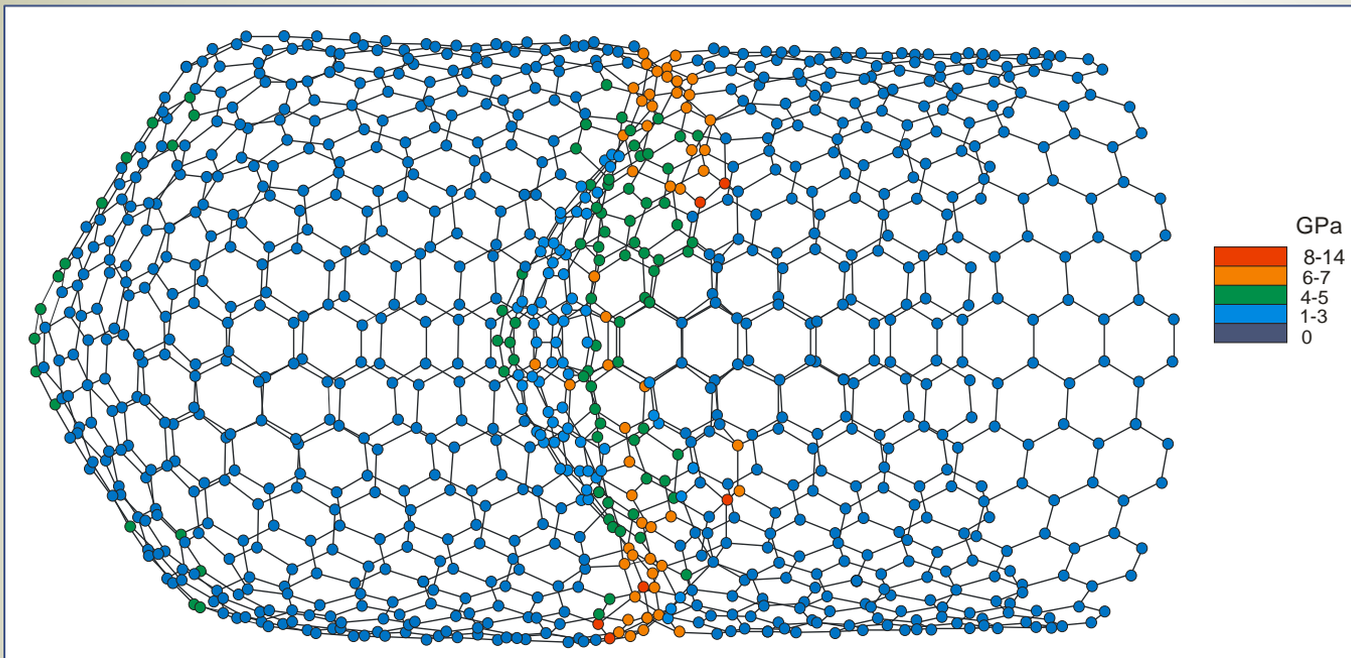


Destruction of the structure of bamboo-like CNT during the increase of the temperature

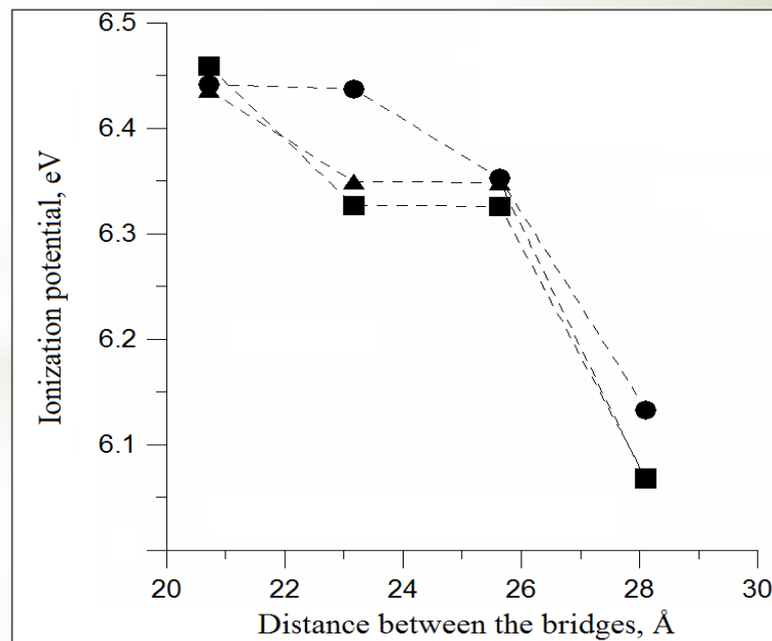
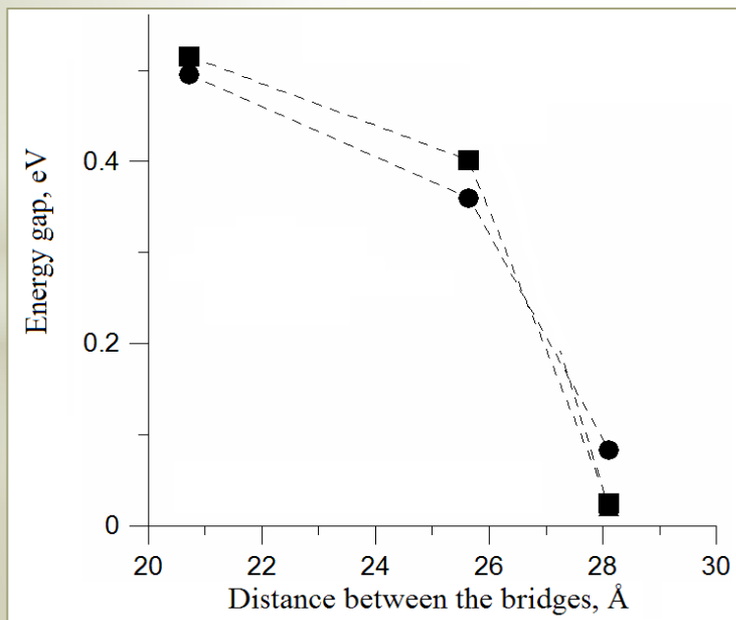
O.E. Glukhova, I.V. Kirillova, A.S. Kolesnikova, E.L. Kossovich, G.N. Ten
 // Proc. of SPIE. 2012. Vol. 8233. P. 82331E-1-82331E-7.

O.E. Glukhova, A.S. Kolesnikova, M.M. Slepchenkov Stability of the thin partitioned carbon nanotubes // Journal of Molecular Modeling 2013. Volume 19. Issue 10. P. 4369-4375.

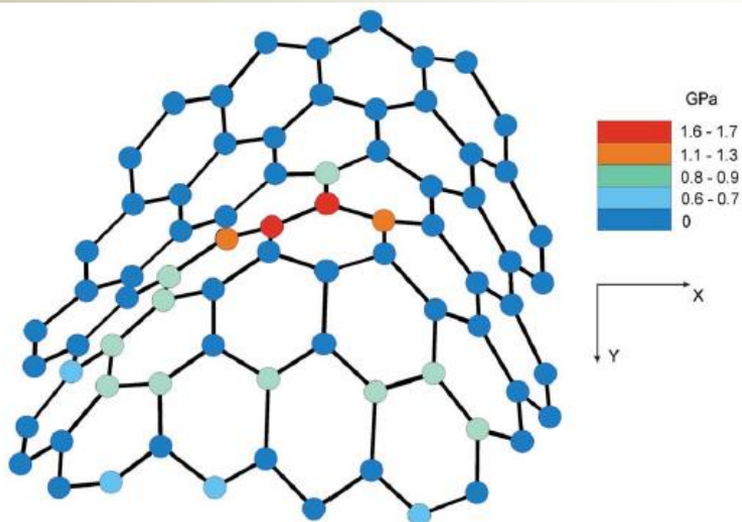




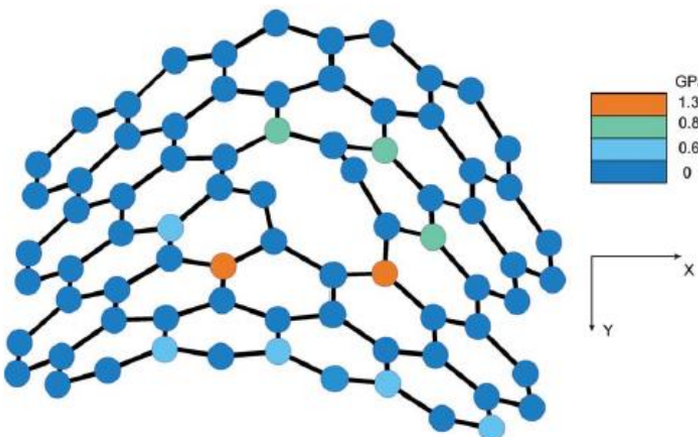
O.E. Glukhova, A.S.
 Kolesnikova, M.M.
 Slepchenkov Stability of
 the thin partitioned
 carbon nanotubes //
 Journal of Molecular
 Modeling 2013. Volume
 19. Issue 10. P. 4369-
 4375.



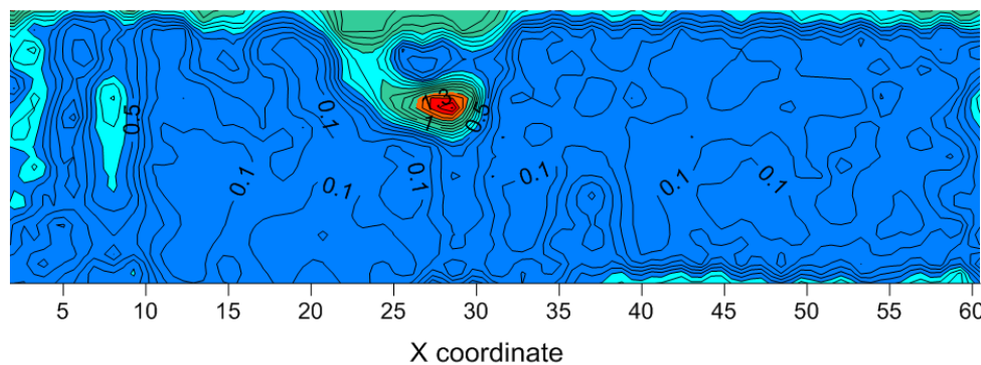
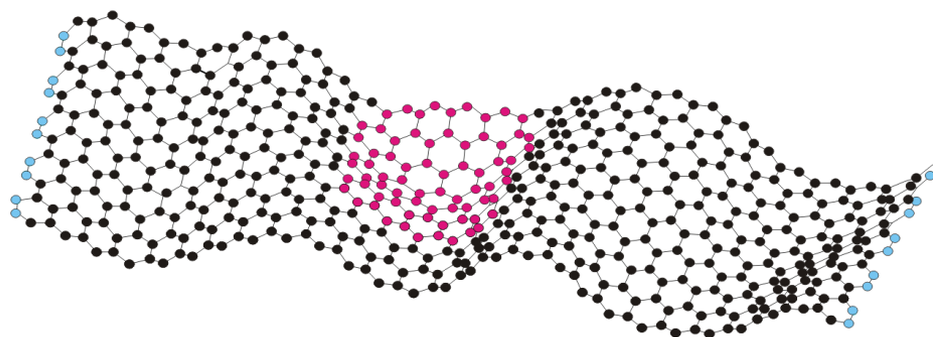
Prediction of the defects appearance



(a)

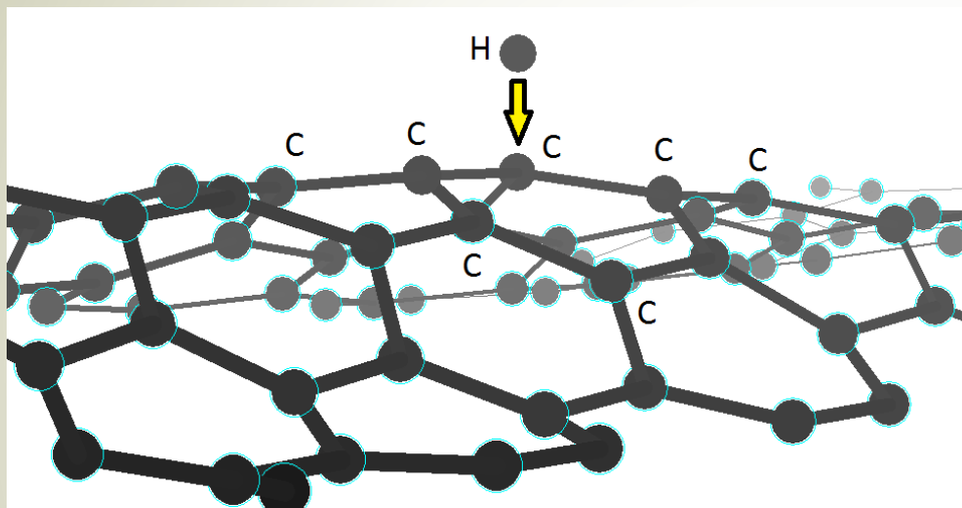


(b)

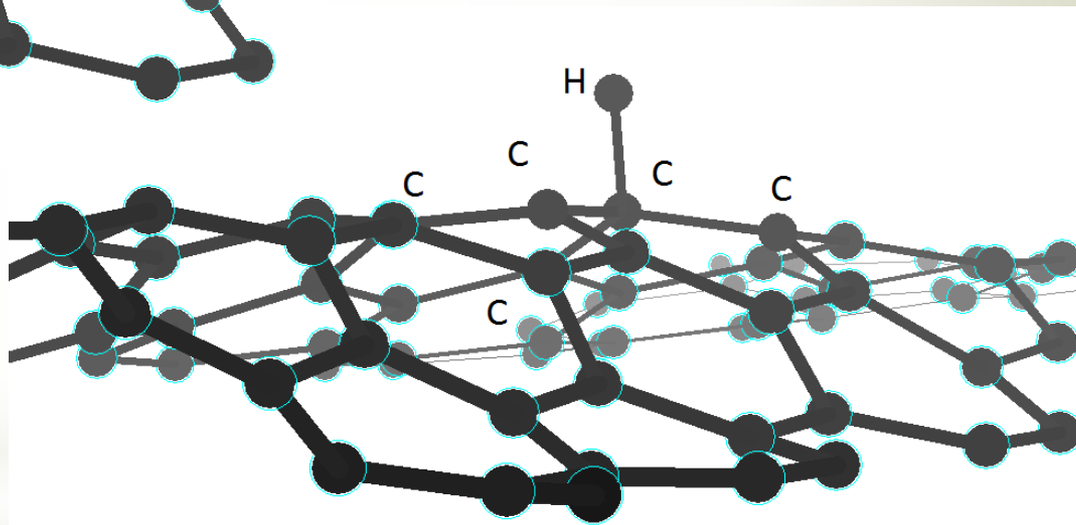


Defects of the C–C bond are observed only between atoms with a local stress value of about 1.8 GPa. One of the most stressed sections of the atomic grid containing a defect is presented in Fig. 4. From the Figure it is clearly seen that after C–C-bond breaking the atomic grid reconstructs and the stress decreases. The enthalpy of the defect formation equals $163.5 \text{ kcal mol}^{-1}$.

The influence of a curvature on the properties of nanostructures



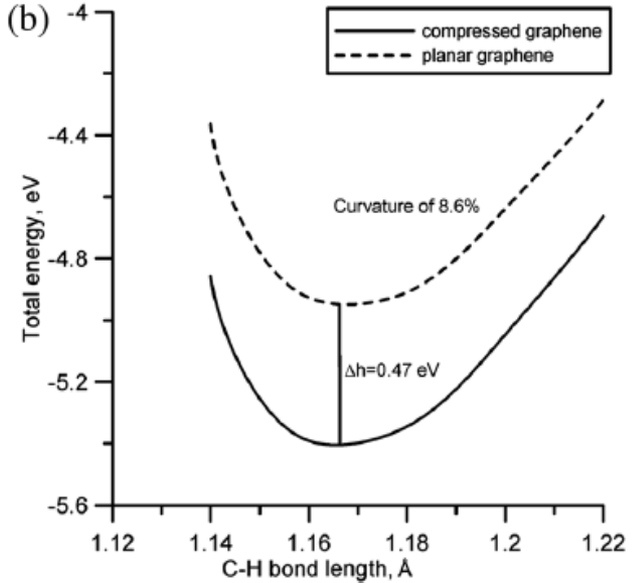
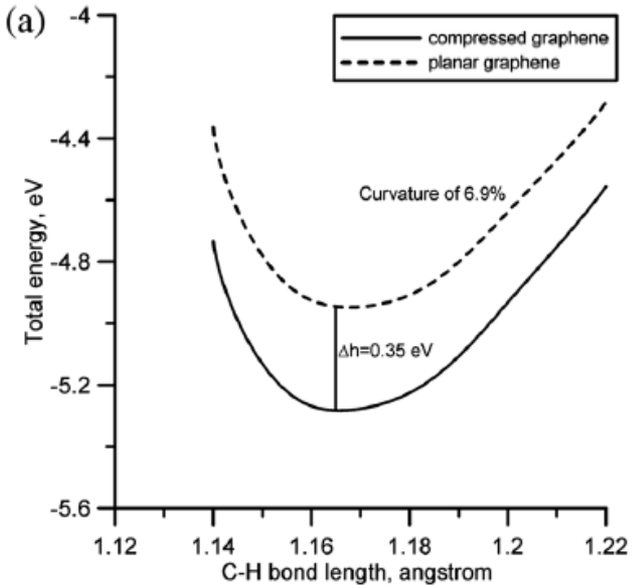
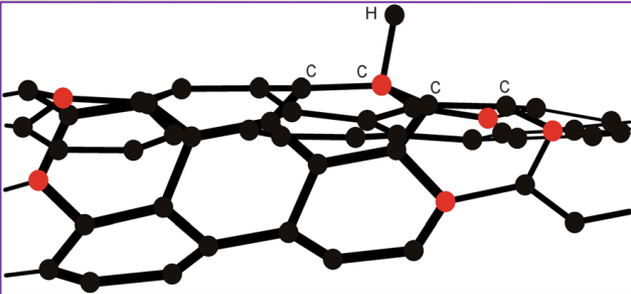
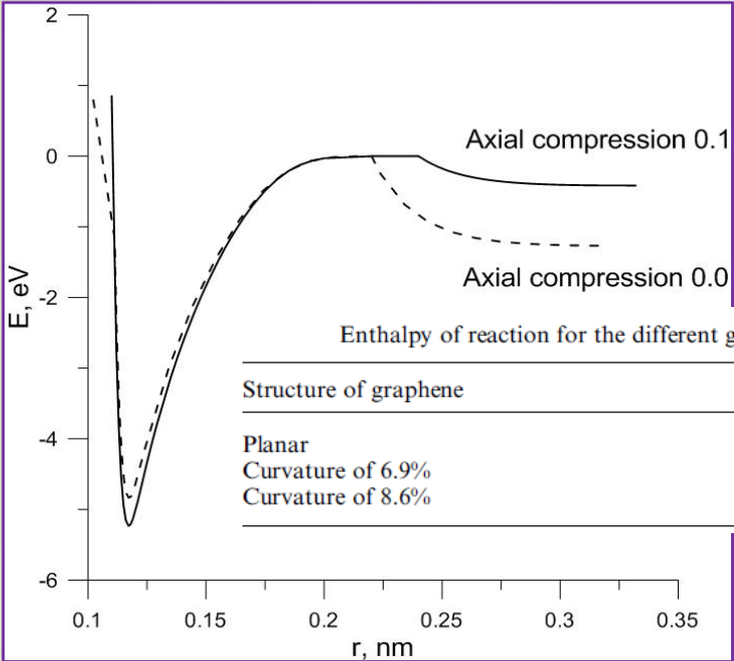
The absorption of H-atom
on the atomic network



O.E. Glukhova, I.V. Kirillova, M.M. Slepchenkov The curvature influence of the graphene nanoribbon on its sensory properties // Proc. of SPIE. 2012. Vol. 8233. P. 82331B-1-82331B-6.

Olga E. Glukhova, Michael M. Slepchenkov Influence of the curvature of deformed graphene nanoribbons on their electronic and adsorptive properties: theoretical investigation based on the analysis of the local stress field for an atomic grid // Nanoscale 2012. Issue 11. Pages 3335-3344. DOI:10.1039/C2NR30477E.

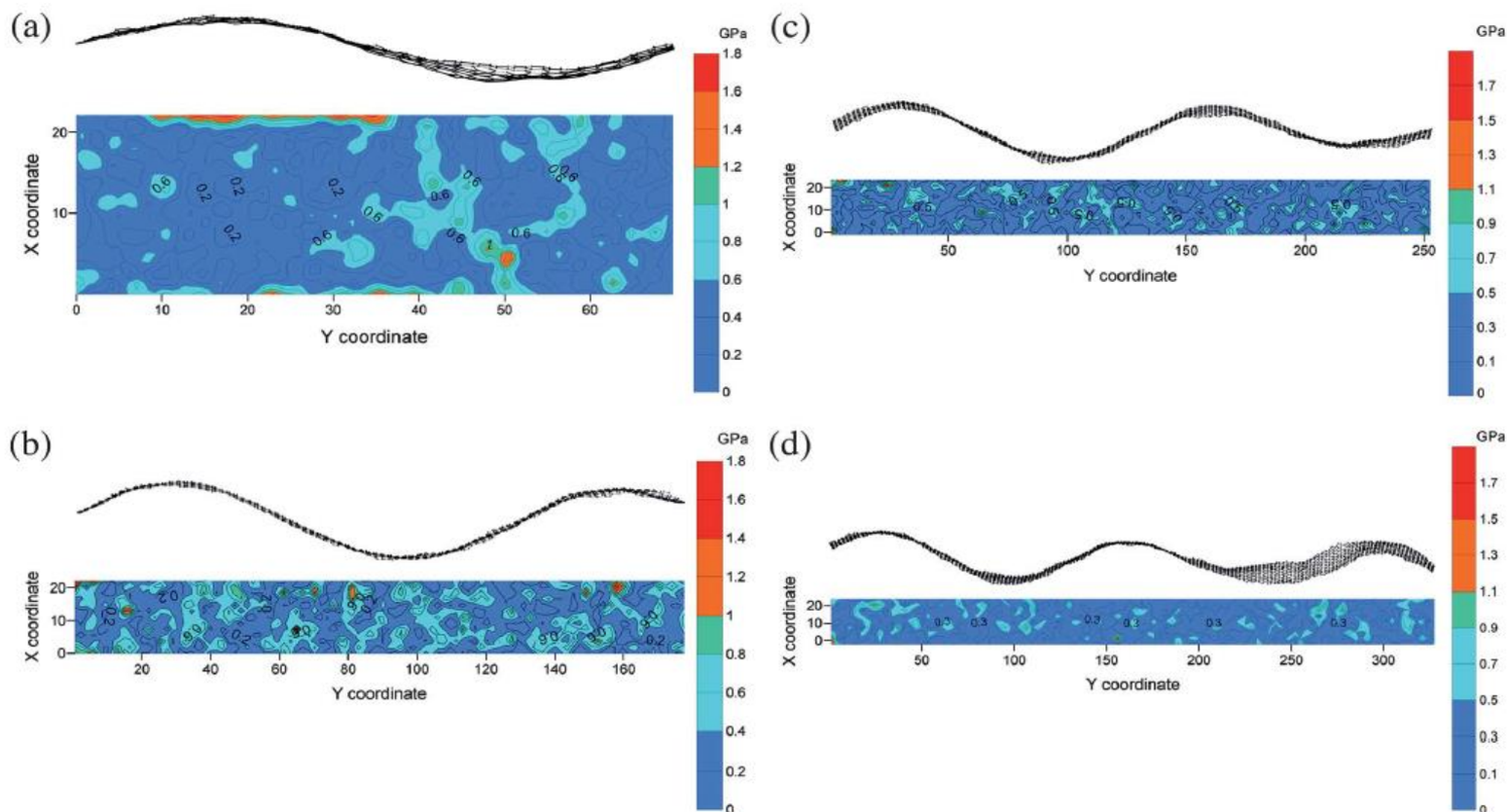
The total energy of the structure depends on the distance between the hydrogen atom and the carbon atom. (The dashed line is the interaction of the hydrogen atom with planer graphene nanoribbon; the solid line is the interaction of the hydrogen atom from wave-like graphene nanoribbon)



The dependence of the chemical C–H interaction energy on the length of the C–H bond for the planar and compressed graphene nanoribbon: (a) with curvature of 6.9%; (b) with curvature of 8.6%.

Geometrical characteristics of the curved armchair graphene nanoribbons compressed up to 98% of initial length

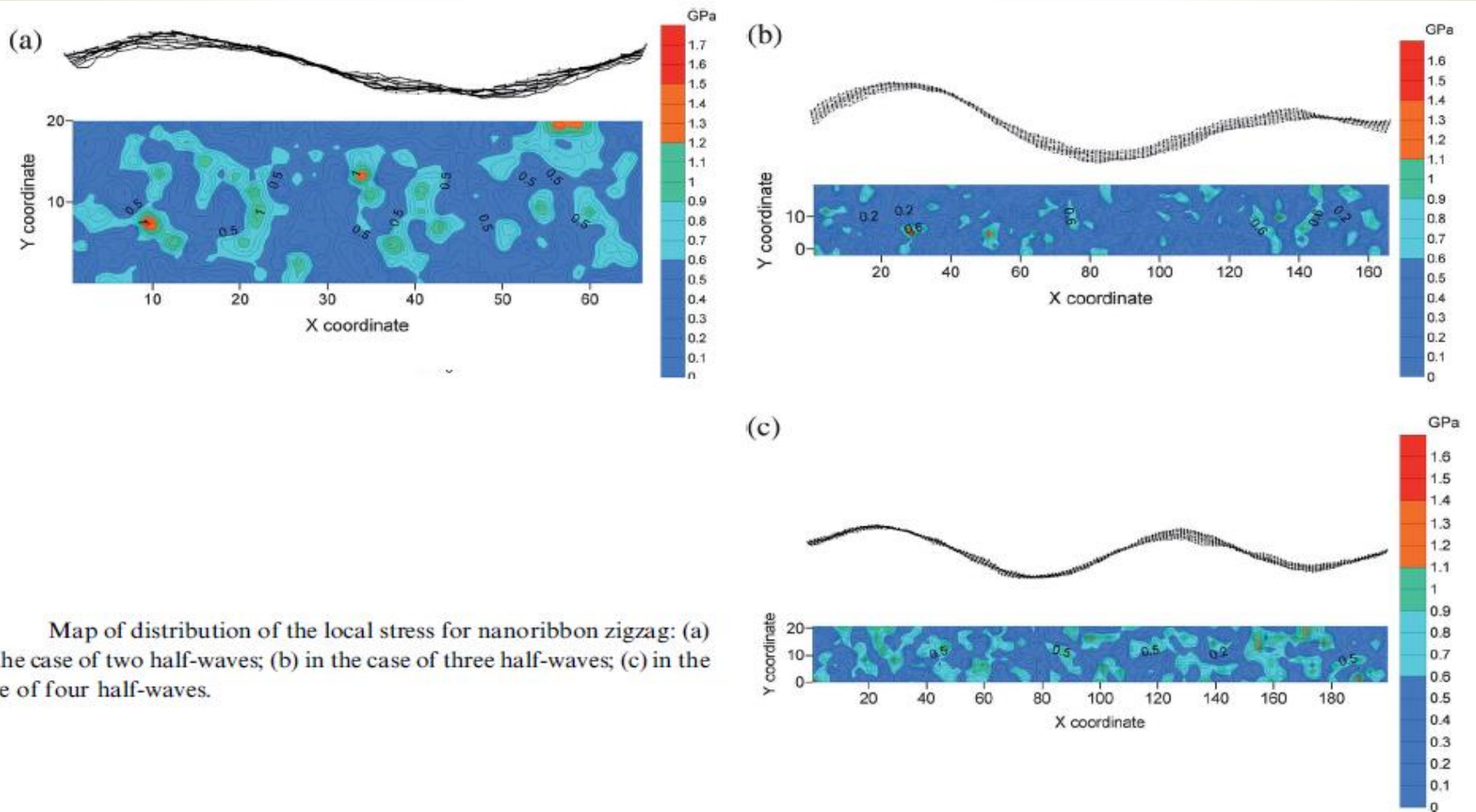
Number of half-waves	Number of atoms in structure	Length of nanoribbon/Å	Length of half-wave/Å	Amplitude of half-wave/Å	Number of hexagons in half-wave	Width of nanoribbon/Å
2	646	71.0	35.5	2.2	9	22.4
3	1634	181.7	60.5	5.3	14	22.4
4	2318	258.4	64.6	5.65	15	22.4
5	3002	335.12	66.2	5.4	15	22.4



Map of distribution of the local stress for the nanoribbon armchair: (a) in the case of two half-waves; (b) in the case of three half-waves; (c) in the case of four half-waves; (d) in the case of five half-waves.

Geometrical characteristics of the curved zigzag graphene nanoribbons compressed up to 98% of the initial length

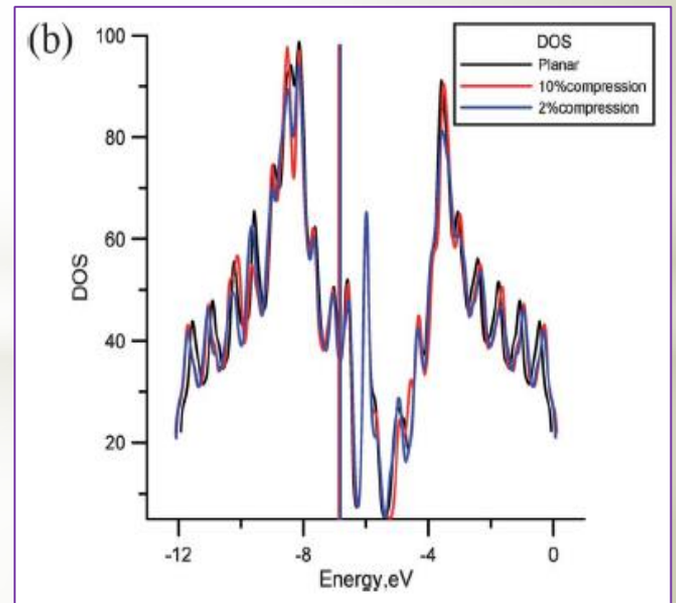
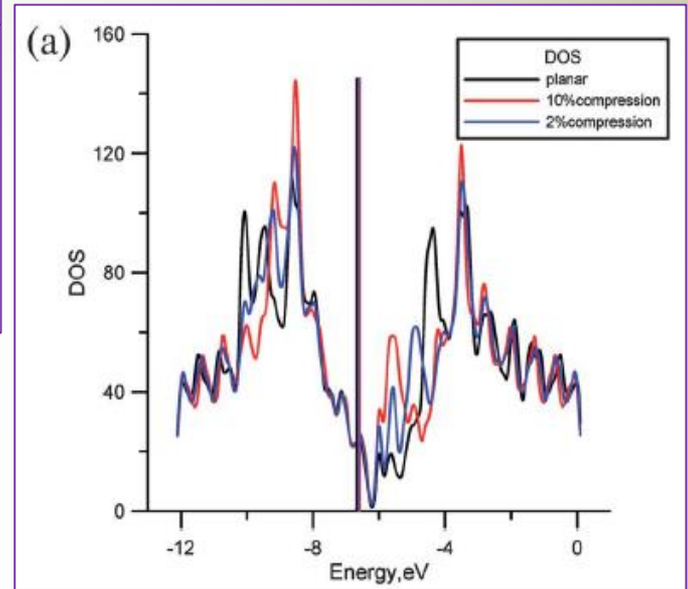
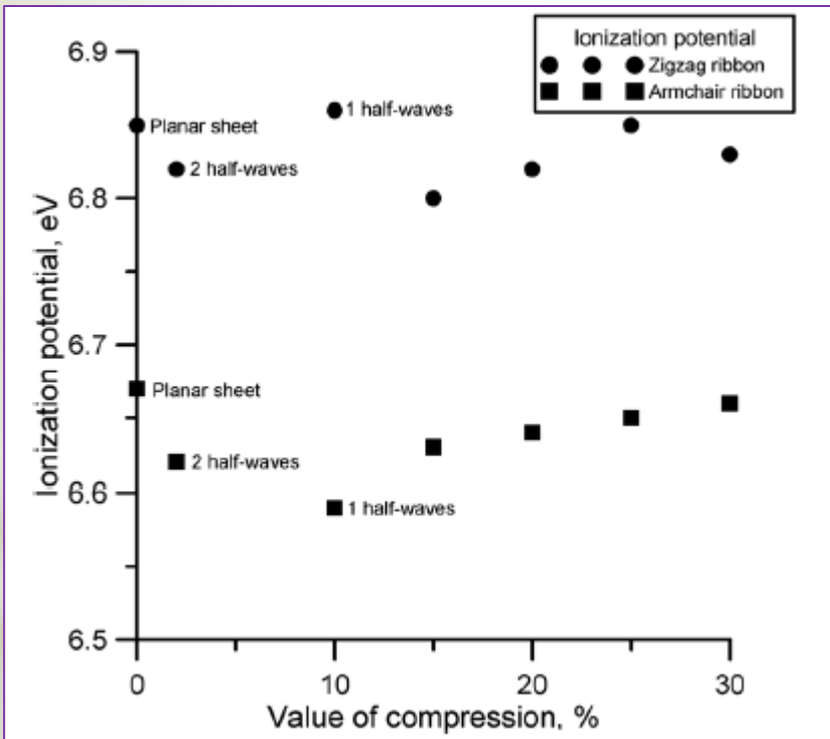
Number of half-waves	Number of atoms in structure	Length of nanoribbon/Å	Length of half-wave/Å	Amplitude of half-wave/Å	Number of hexagons in half-wave	Width of nanoribbon/Å
2	550	65	32.5	2.8	12	19.88
3	1390	165.18	55.06	5.4	20	
4	1670	198.7	49.6	5.6	20	



Map of distribution of the local stress for nanoribbon zigzag: (a) in the case of two half-waves; (b) in the case of three half-waves; (c) in the case of four half-waves.

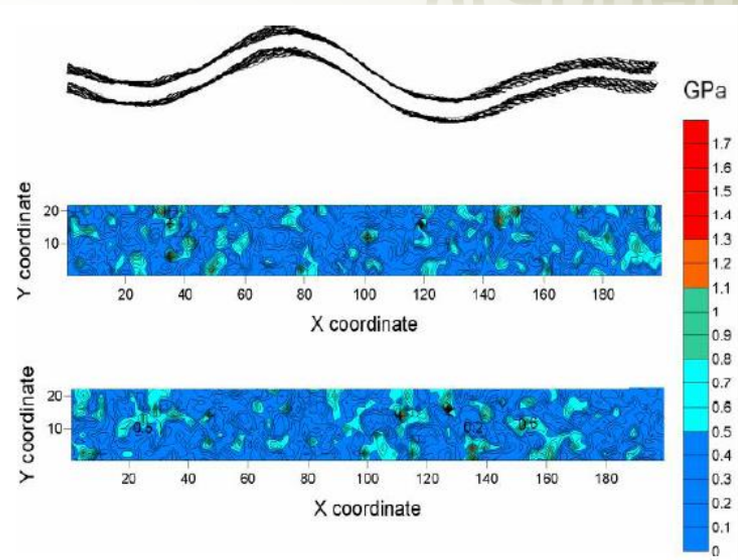
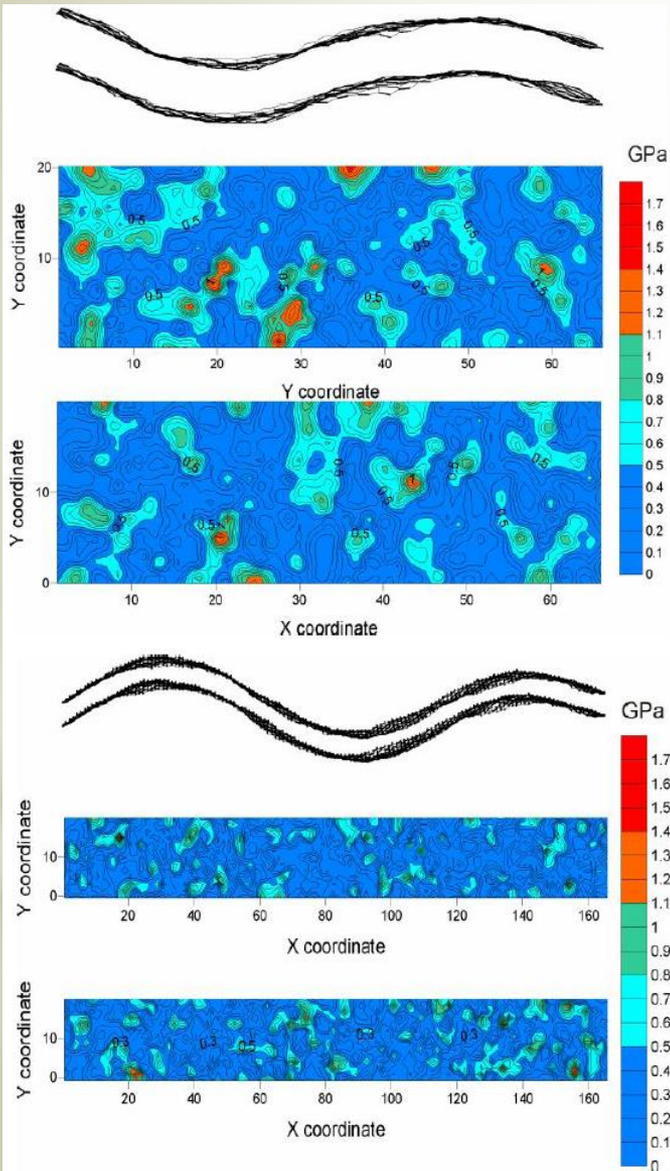
Some parameters of the electronic structure of nanoribbons

Number of half-waves	Length of half-wave/Å	IP/eV	E_{gap}/eV
<i>Armchair ribbon of width 22.4 Å</i>			
2	71.0	6.63 (6.65)	0.04 (0.03)
3	181.7	6.50 (6.53)	0.04 (0.03)
4	258.4	6.44 (6.47)	0.02 (0.01)
5	335.12	6.41 (6.44)	0.04 (0.02)
<i>Zigzag ribbon of width 19.88 Å</i>			
2	65	6.82 (6.84)	0.04 (0.02)
3	165.18	6.79 (6.81)	0.01 (0.01)
4	198.7	6.80 (6.81)	0.01 (0.01)



O.E. Glukhova, A.S.Kolesnikova Mechanical and emission properties of thinnest stable bamboolike nanotubes // Journal of Physics: Conference Series 393 (2012) 012027.

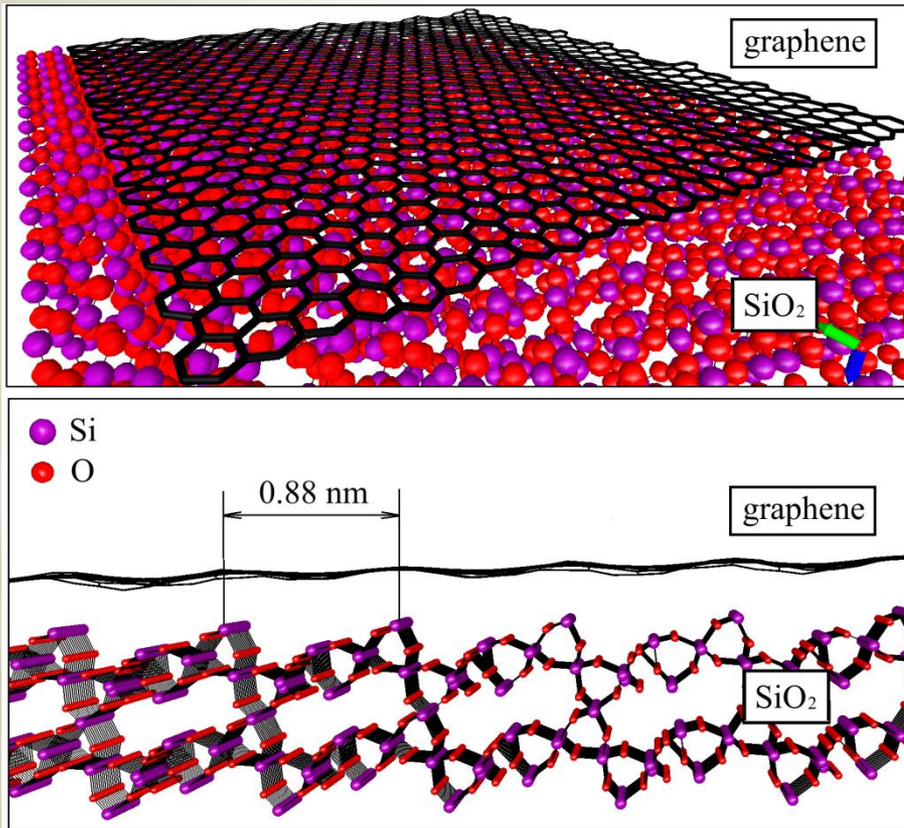
* The compression process of bi-layer graphene



Geometrical characteristics of the curved zigzag bi-layer graphene nanoribbons compressed up to 98% of the initial length

Num ber of half- wave s	Num- ber of atoms in struc- ture	Length of nanorib- bon, Å	Leng- th of half- wav- e, Å	Ampli- tude of half- wave, Å	Num- ber of hex- agons in half- wave	Width of nanorib- bon, Å
2	1100	65	32.3	3.1	13	
3	2780	165.18	55.4	5.48	20	19.88
4	3340	198.7	49.8	5.55	20	

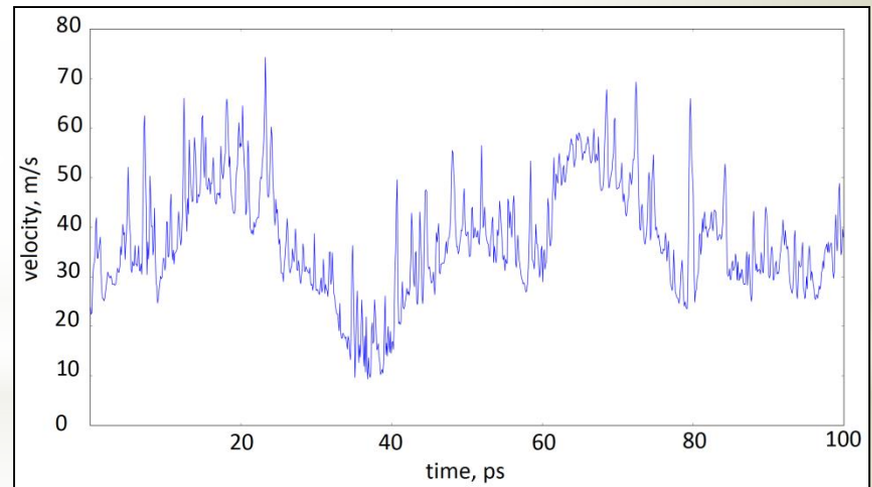
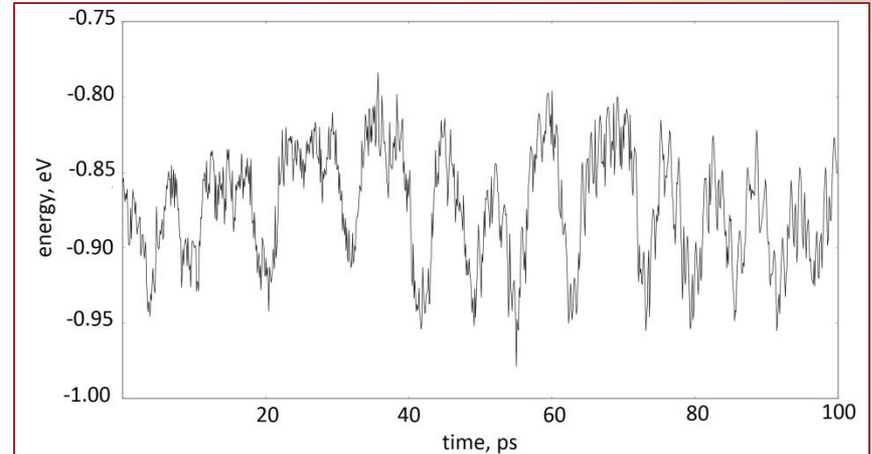
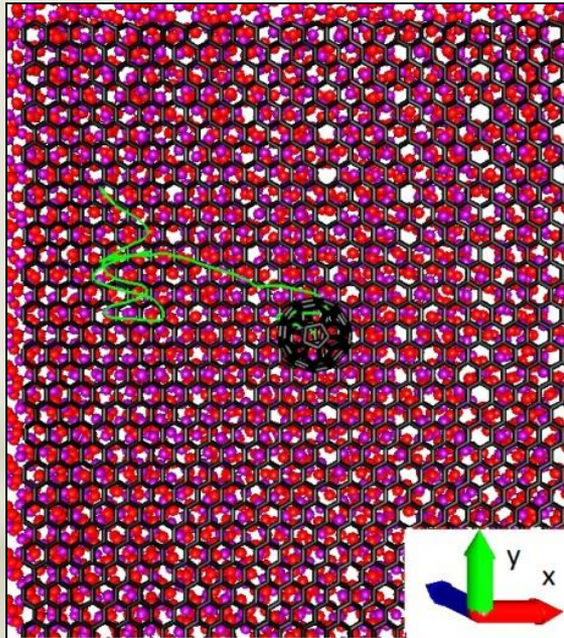
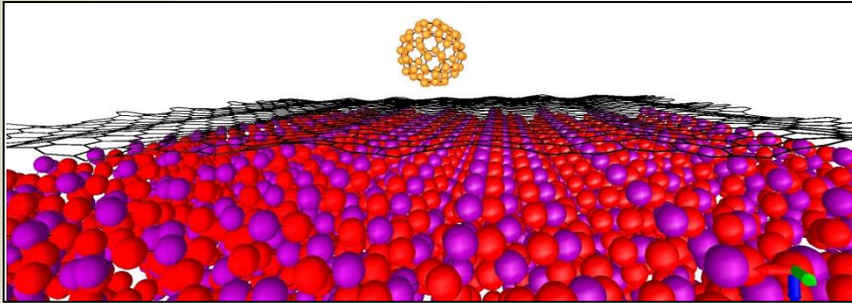
* The control of movement of C60 on rippled graphene located on substrate SiO₂



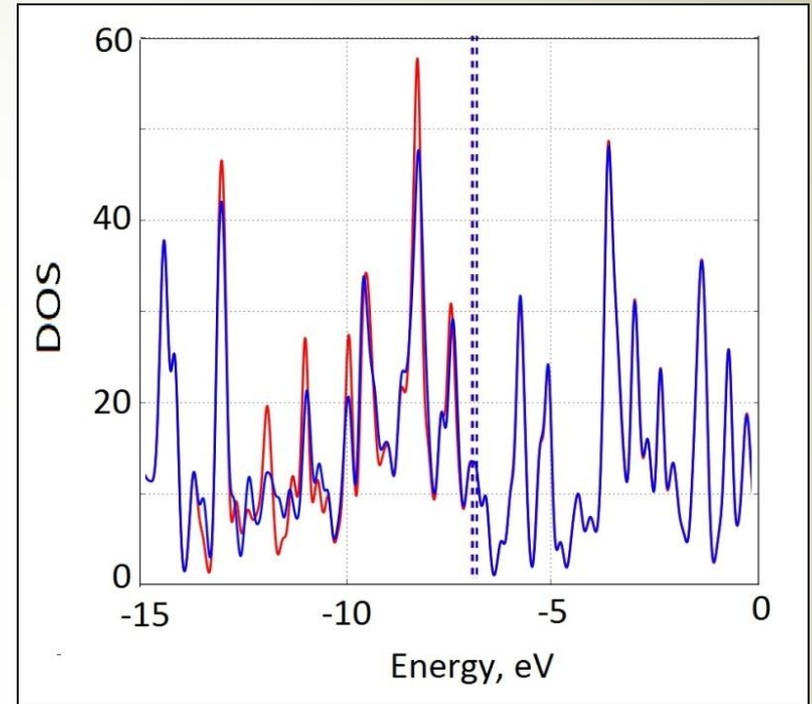
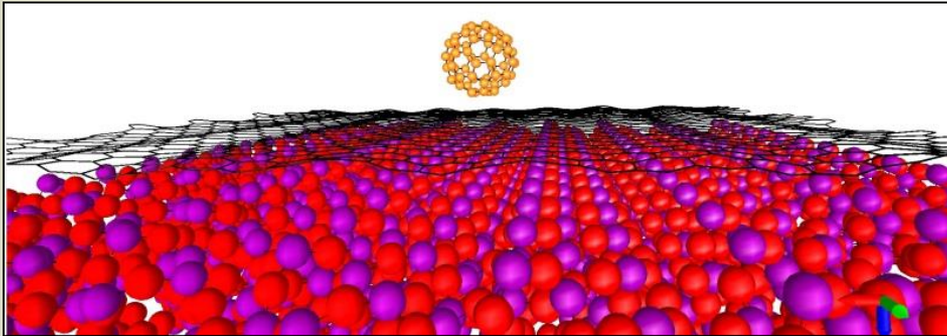
The average distance of the graphene-substrate is ~0.3 nm, the adhesion is 1.8 eV/nm² that well agrees with the experimental studies [NATURE NANOTECHNOLOGY | VOL 6 | SEPTEMBER 2011].

Monolayer graphene on a substrate

C₆₀+graphene on the ideal surface

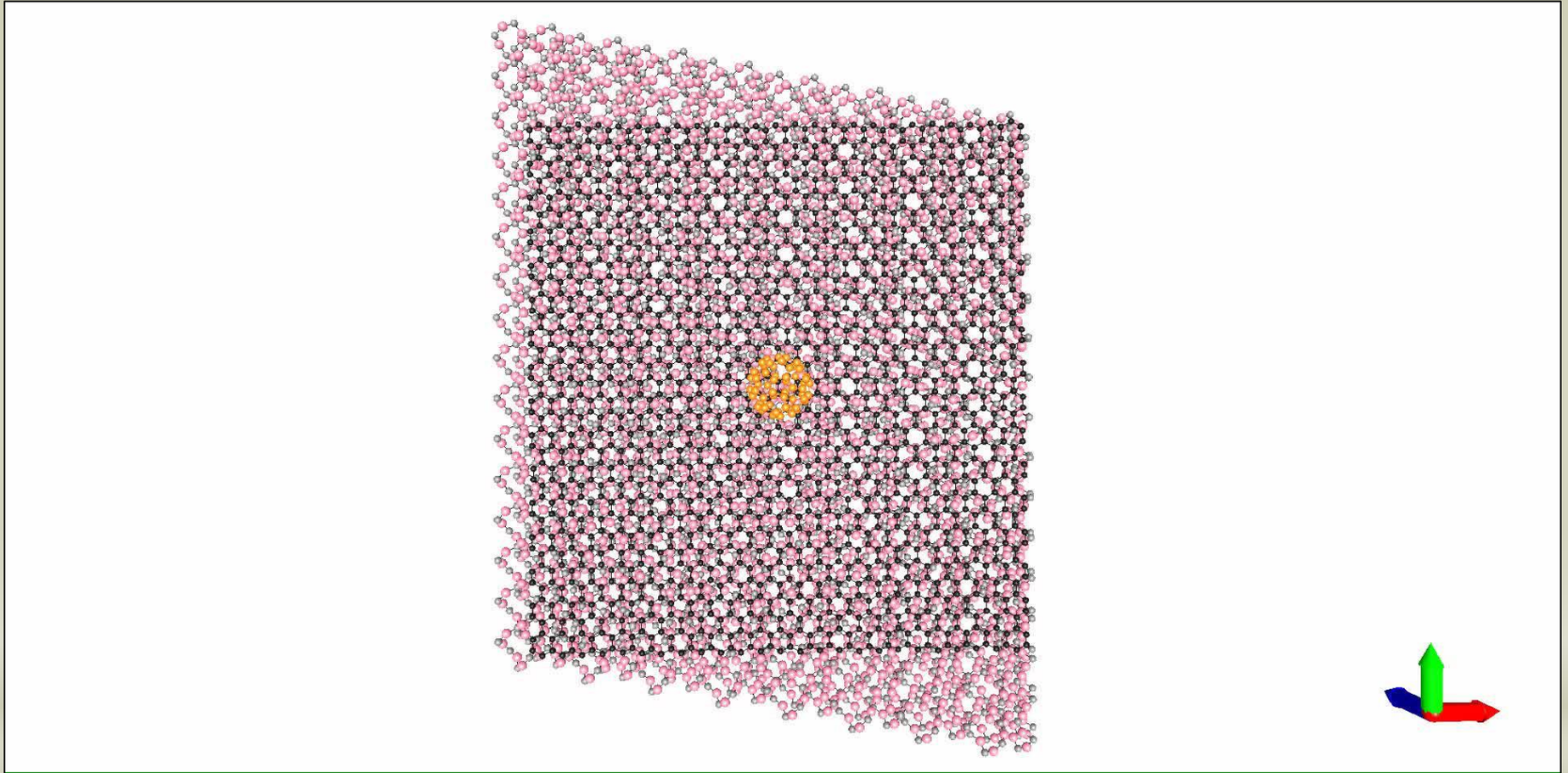


Fullerene on graphene with substrate: general view; the trajectory of the mass center at $T = 300$ K during 100 ps; the change in the energy of interaction of C₆₀ with graphene during its free motion at $T = 300$ K; the velocity of the fullerene C₆₀

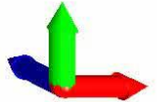
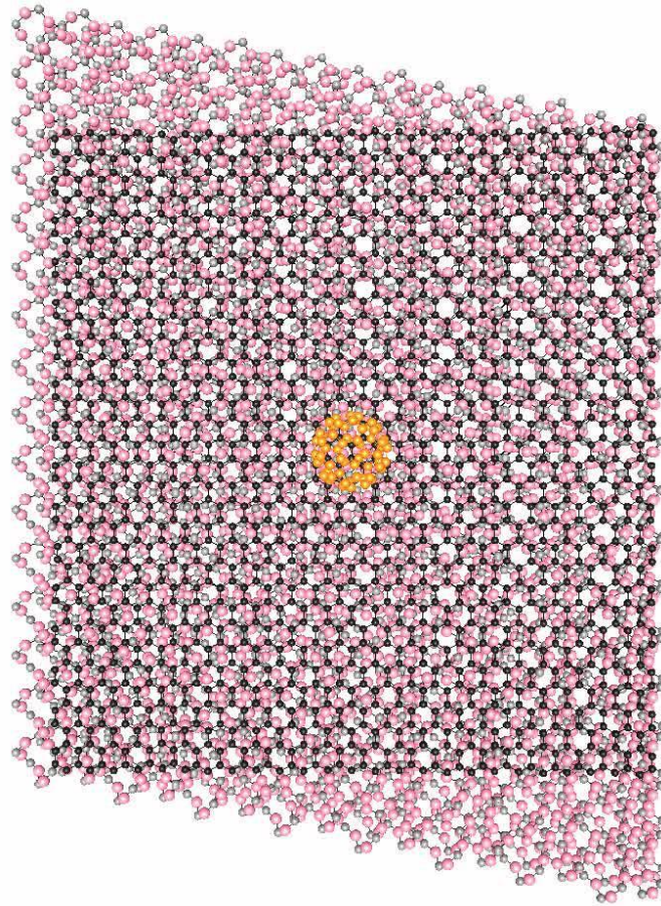


The density of electron states of complex C60+graphene near the last filled HOMO-level (vertical dotted lines indicate the position of the HOMO level).

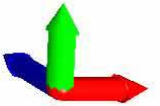
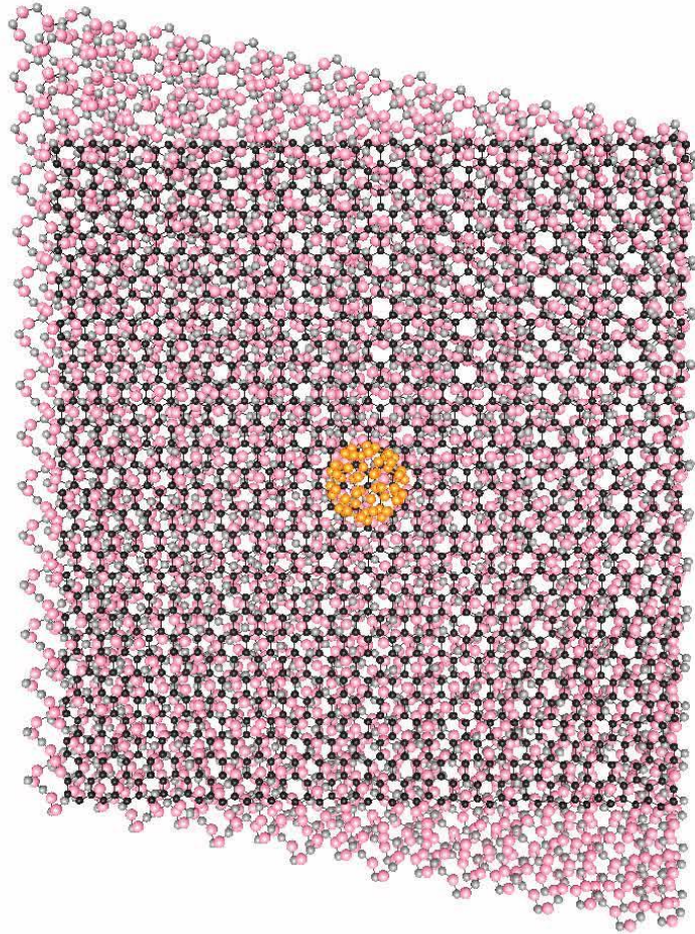
The **red curve** corresponds to the case without account of additional overlap of the electron clouds of the fullerene and graphene, **blue** - to the case with account.



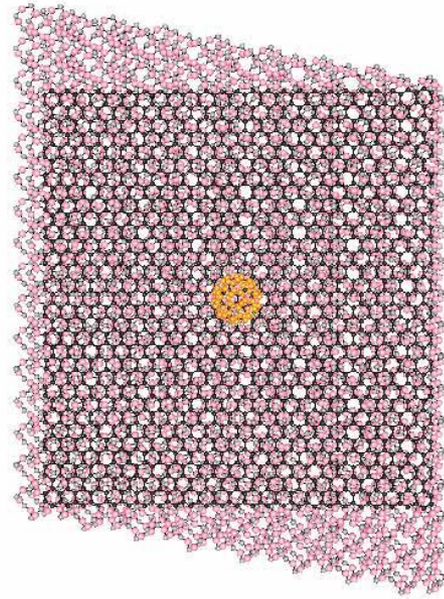
* On the ideal surface SiO_2



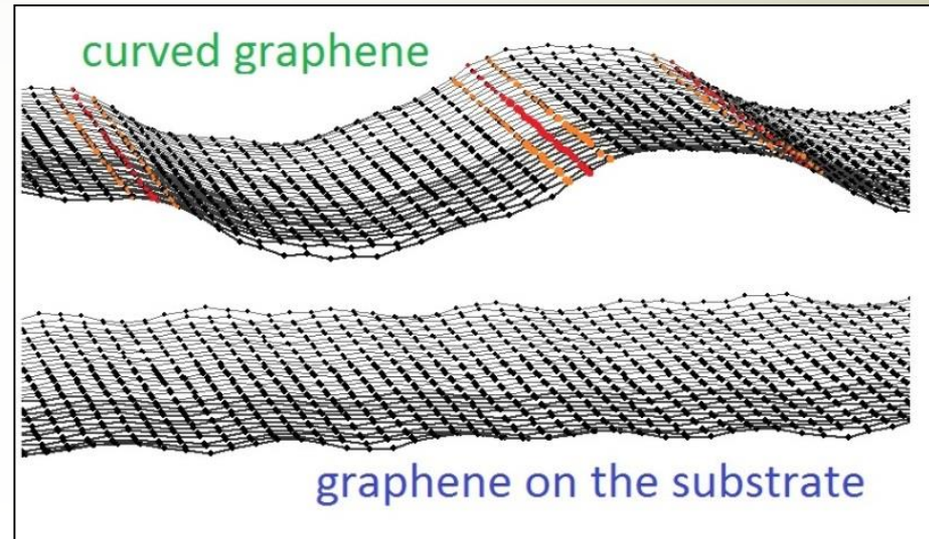
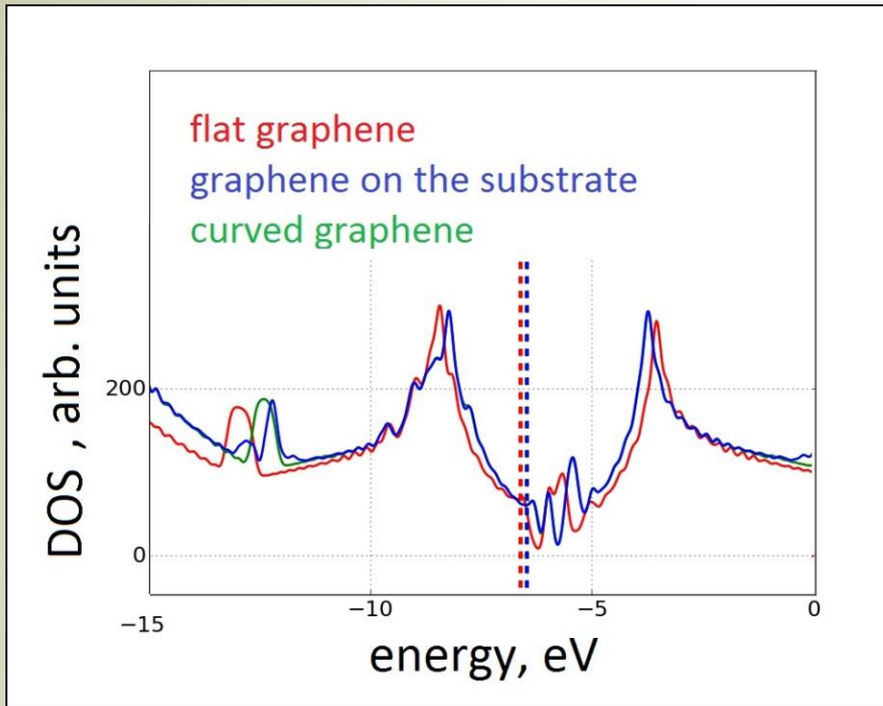
* $E_y = 10 \text{ V/nm}$



* $E_y = 50 \text{ V}/\mu\text{m}$



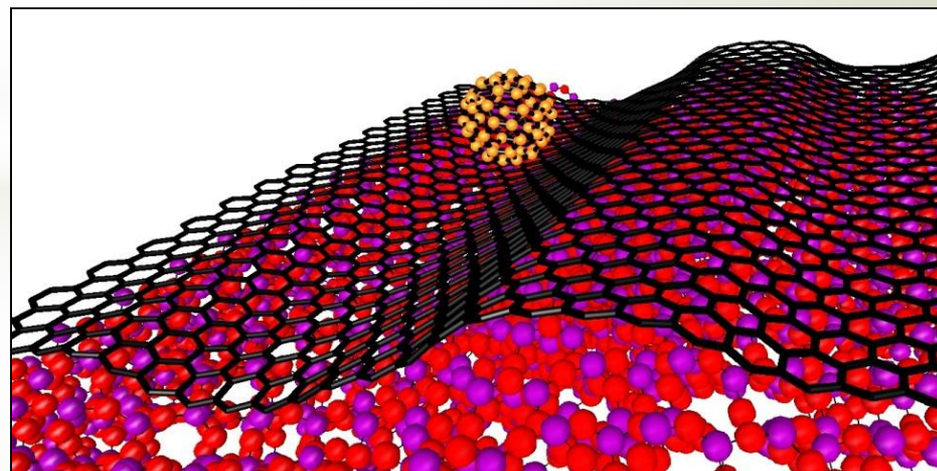
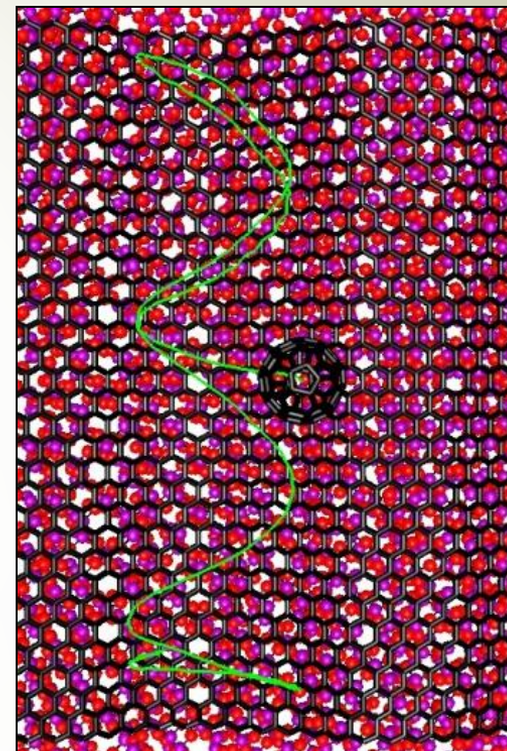
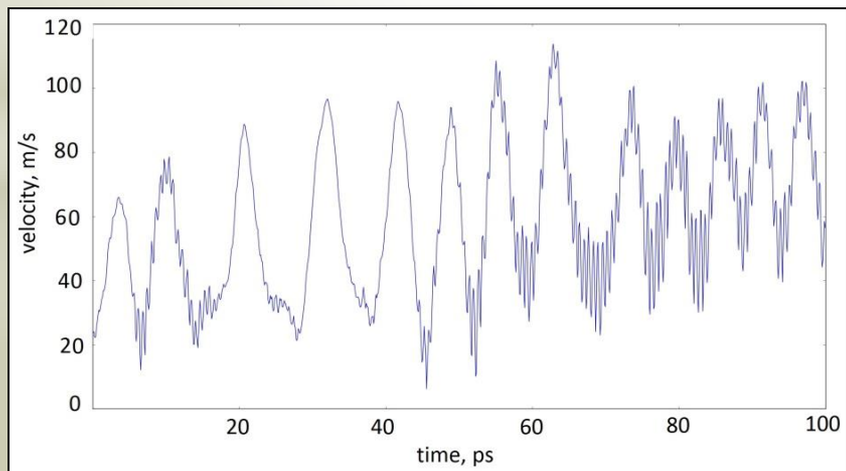
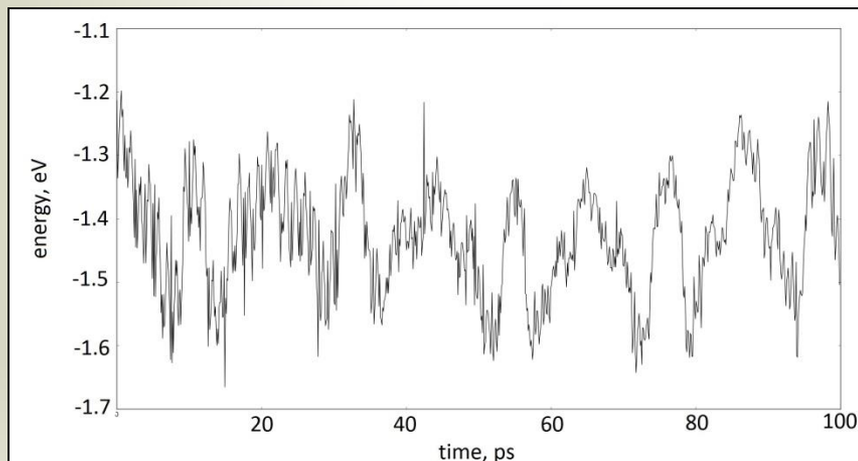
* $E_y = 100 \text{ V/mkm}$

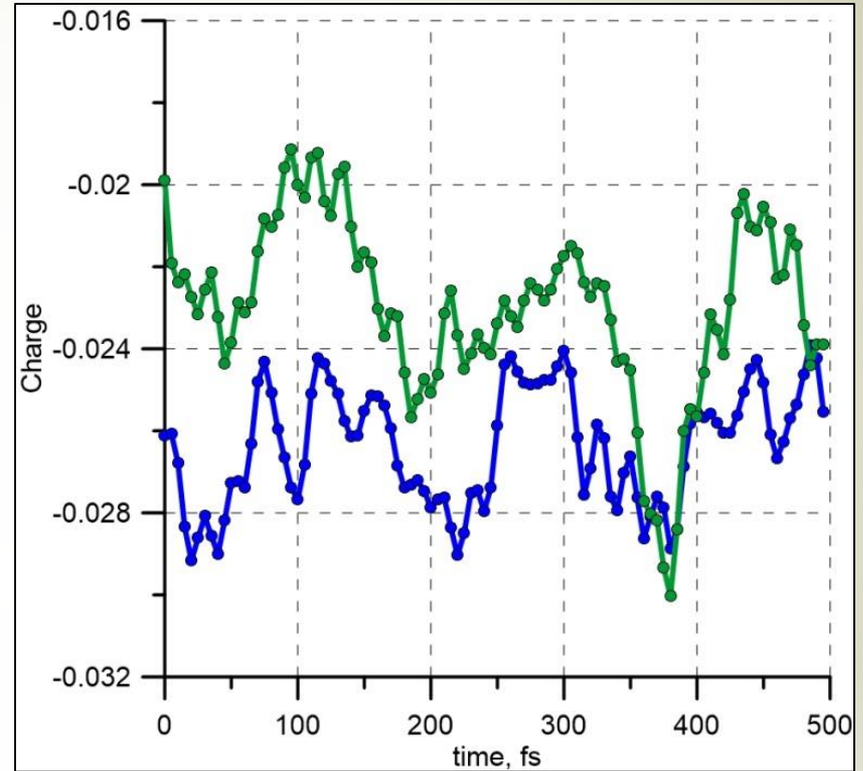
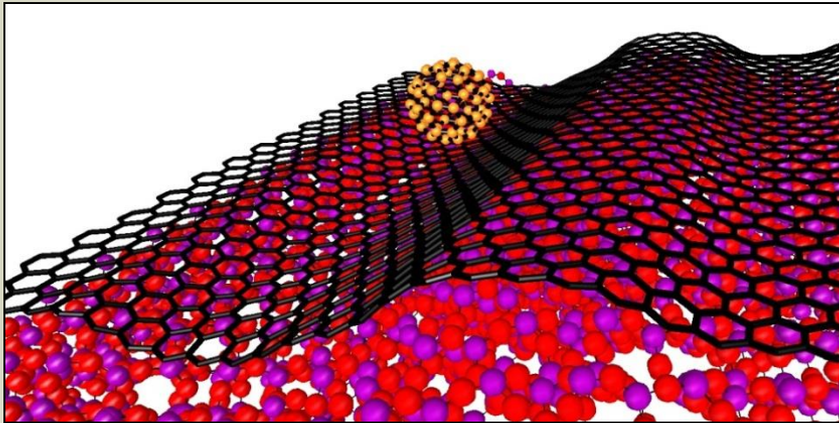
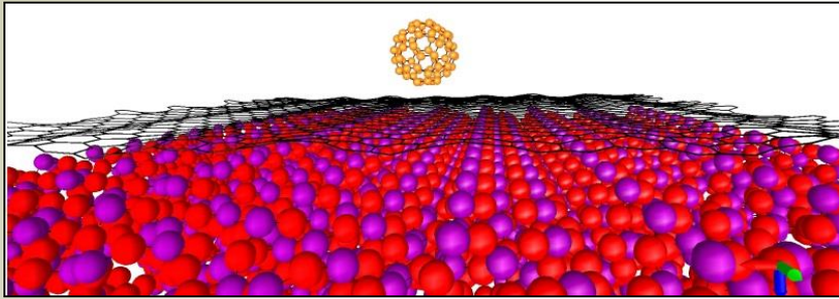


Graphene on a substrate:

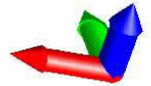
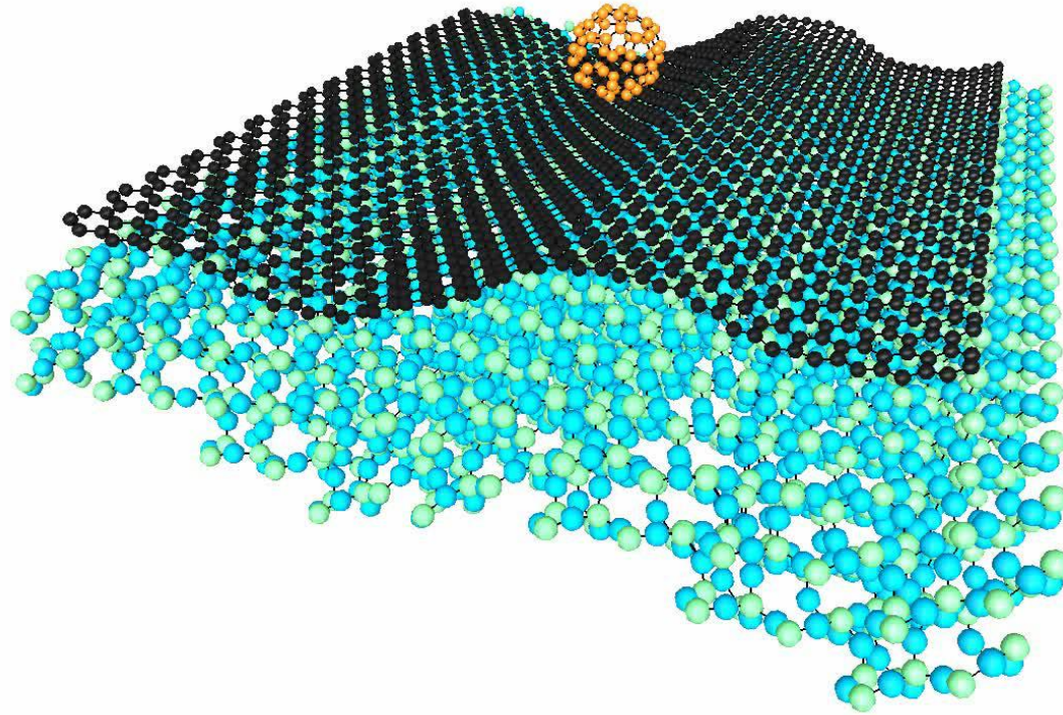
- the density of electronic states of graphene on an ideal and corrugated substrate (dashed lines indicate the position of HOMO-levels);
- fields of atomic mesh with rehybridized electron clouds (the maximum degree of rehybridization belongs to the atoms marked with red dots and orange). Some of the electrons (red highlighted) will be eventually located in $sp^{2.02}$.

- * Fullerene on the corrugated substrate at $T = 300$ K:
 - general view,
 - the trajectory of the mass center,
 - changes of velocity,
 - the oscillations of the interaction energy.

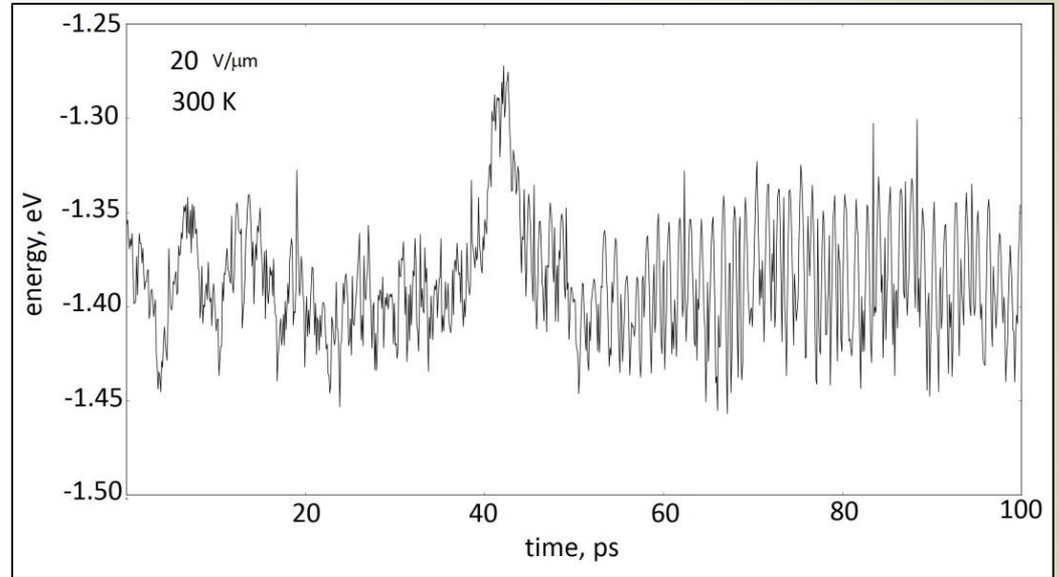
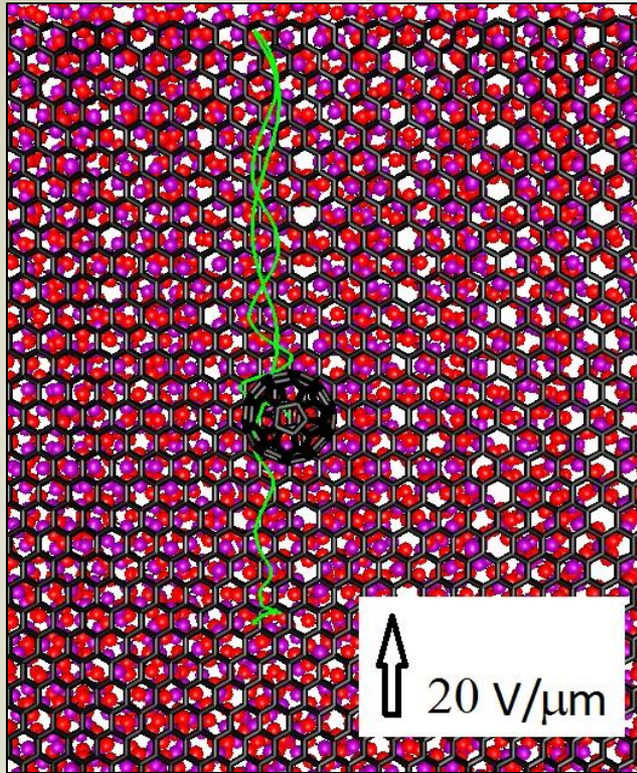




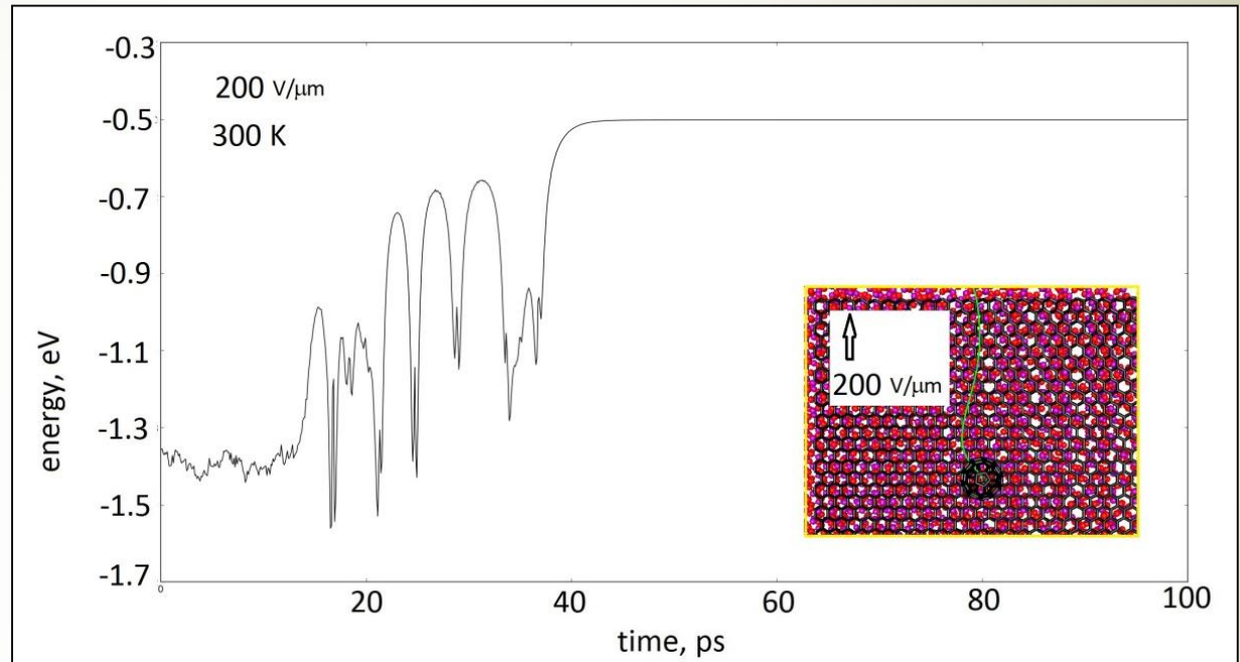
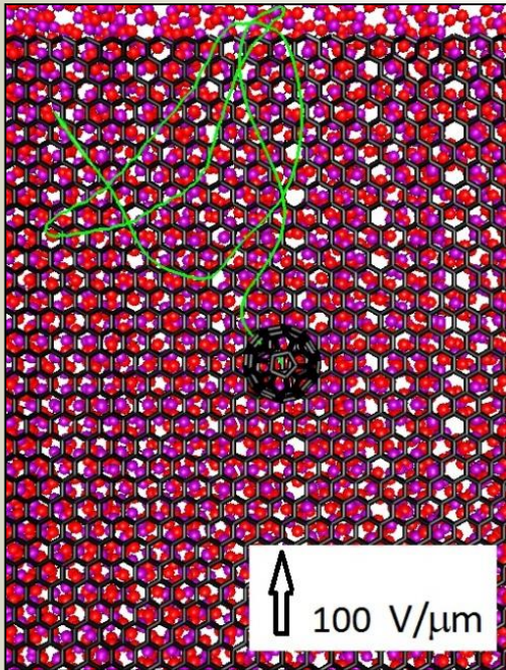
- * Change of charge on fullerene during its motion on graphene:
 - blue curve** - movement on ideal defectless substrate SiO₂. The current flowing within 10-20 fs can reach 14-17 nA;
 - green curve** - motion on graphene in corrugated substrate. Thus the current in the molecular complex reaches 16 nA.
- T = 300 K, time step - 5 fsek.



* On the corrugated SiO₂



Fullerene on the corrugated substrate in external electric field at $T = 300$ K:
- the trajectory of the mass center and the change in the interaction energy for 100 psec at a field strength of 20 V/ μm



- * Fullerene on the corrugated substrate in external electric field at $T = 300$ K:
 - at a field strength of 100 V/mkm;
 - at a field strength of 200 V/mkm

* Giga- and terahertz range nanoemitter based on a peapod structure

M.M. Slepchenkov¹, A.S. Kolesnikova¹, G.V. Savostiyanov¹, Igor S. Nefedov²,
Ilya V. Anoshkin³, Alexandr V. Talyzin⁴, Albert G. Nasibulin^{3,5}, Olga E. Glukhova¹

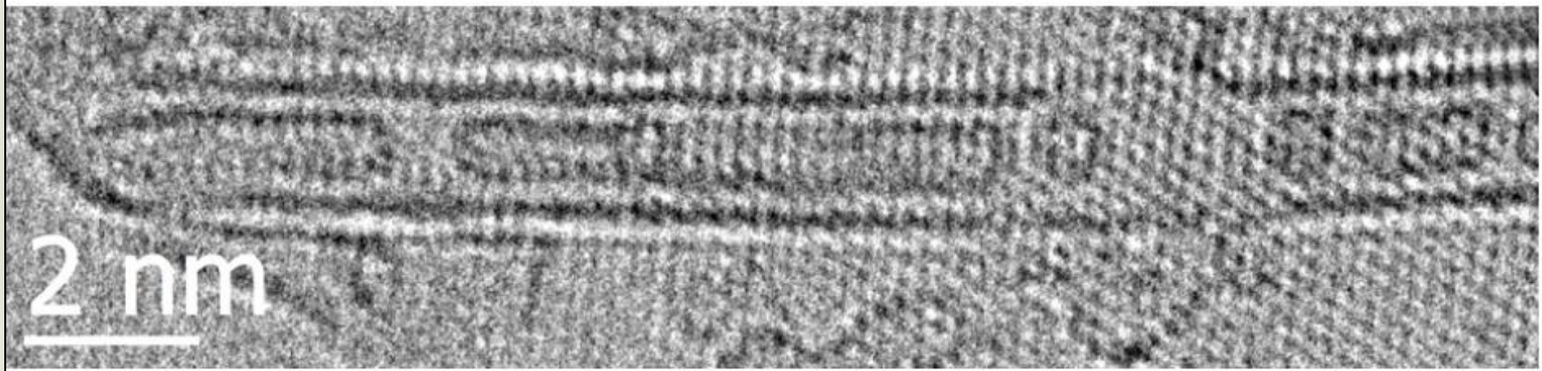
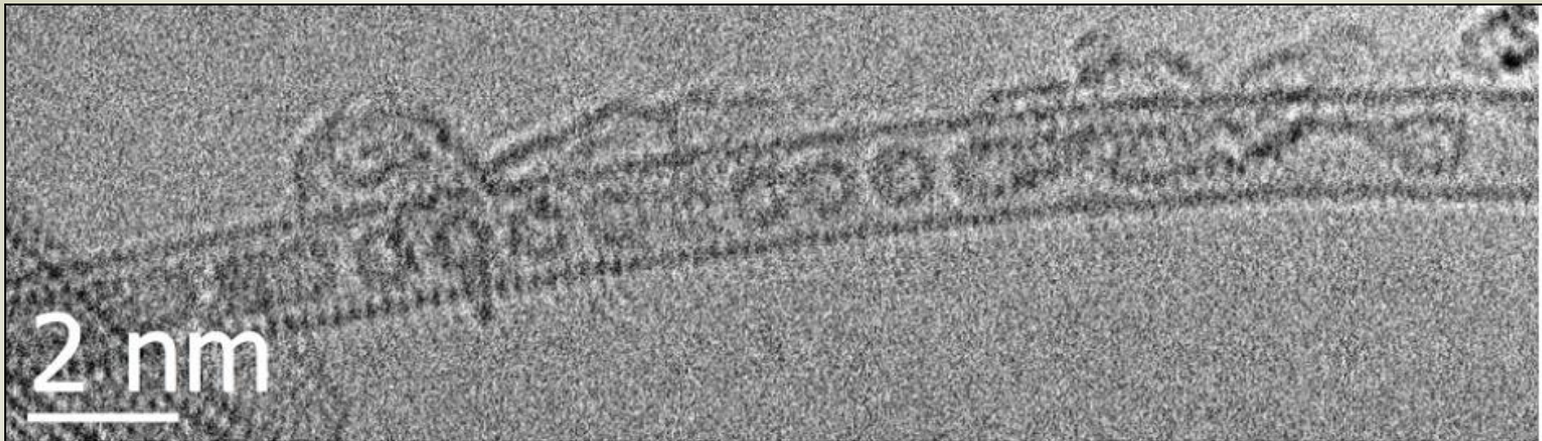
¹ Saratov State University, Department of Physics, Russian Federation

² Aalto University School of Electrical Engineering, Department of Radio Science and
Engineering, Finland

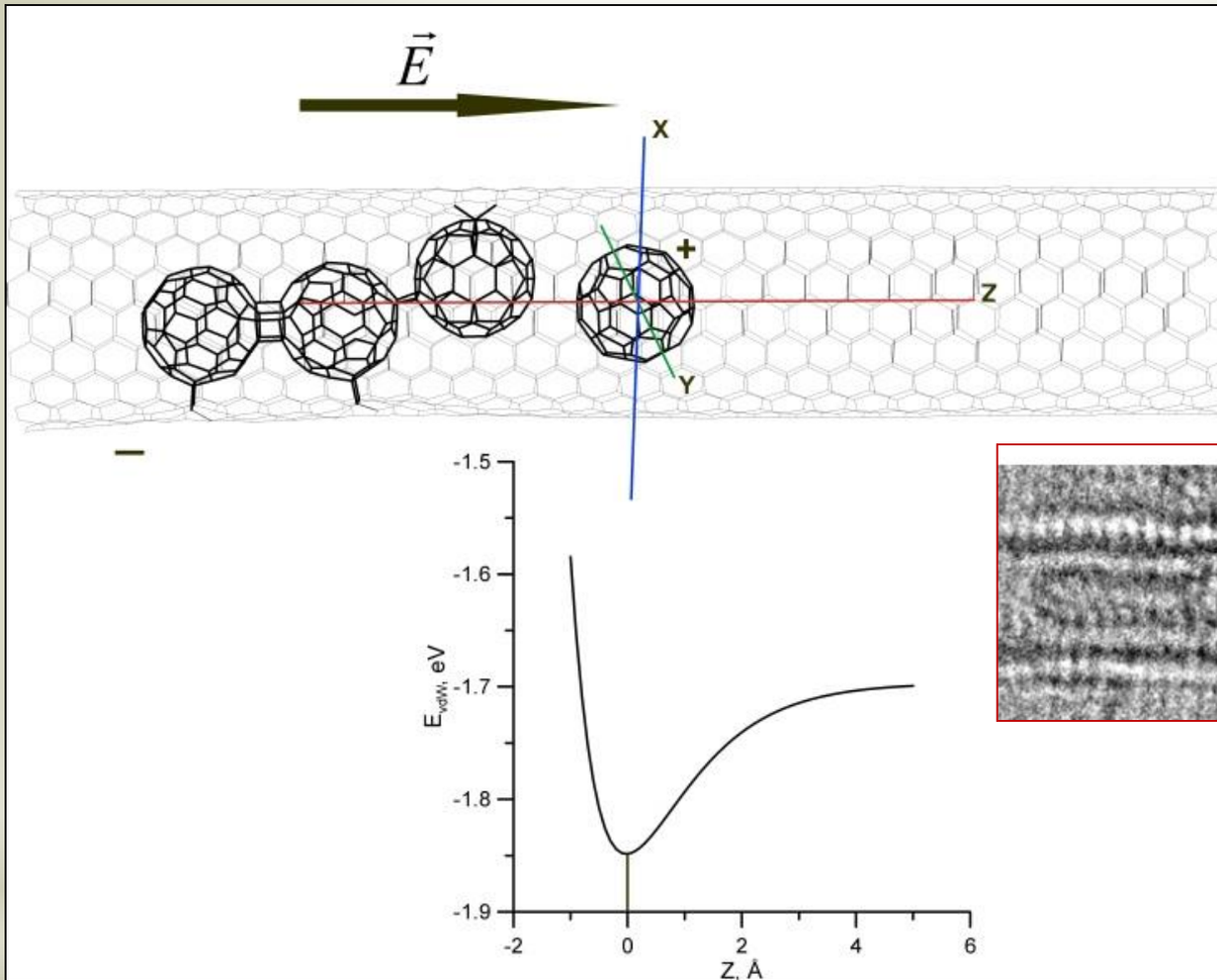
³ Aalto University School of Science, Department of Applied Physics, Espoo, Finland

⁴ Umeå University, Department of Physics, S-90187 Umeå, Sweden

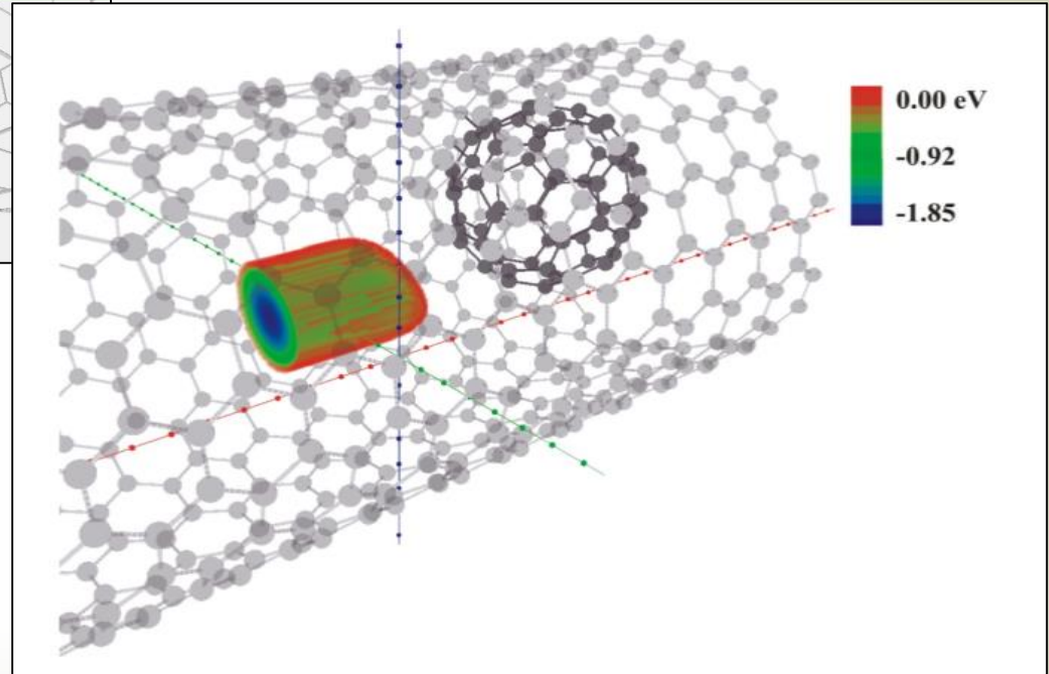
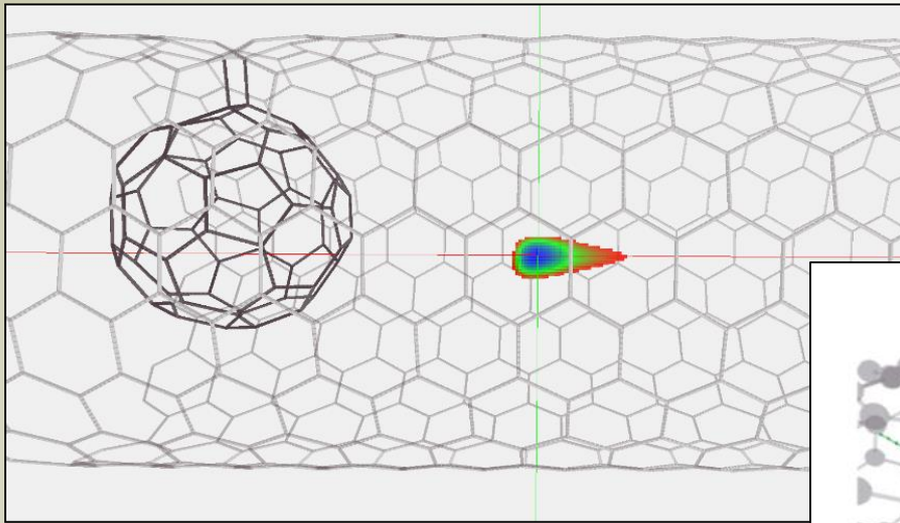
⁵ Skolkovo Institute of Science and Technology, 100 Novaya st., Skolkovo, 143025,
Russia



* HR-TEM images illustrating the partial polymerization of fullerene molecules inside CNTs



* Model of a nanoemitter: configuration of fullerenes inside (10,10) carbon nanotube



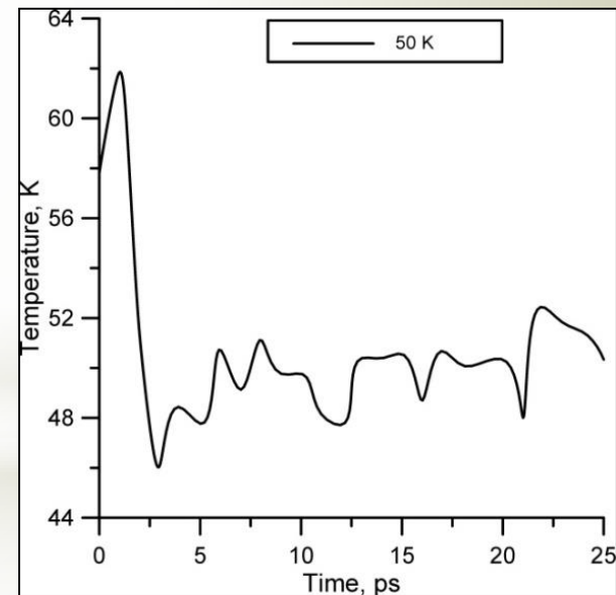
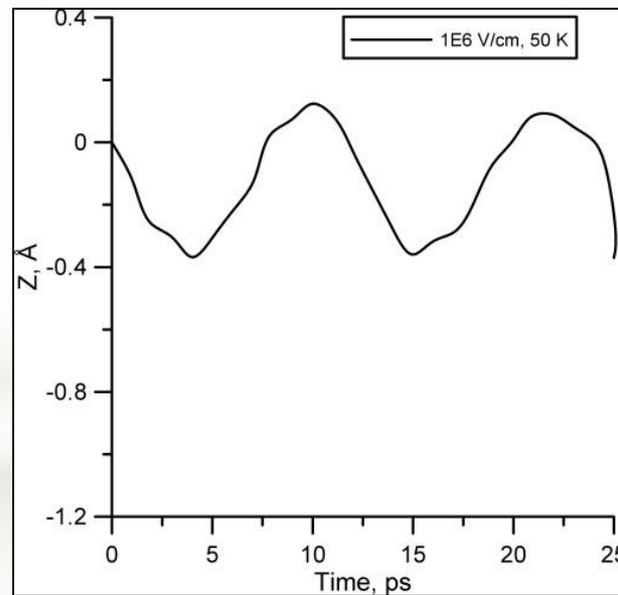
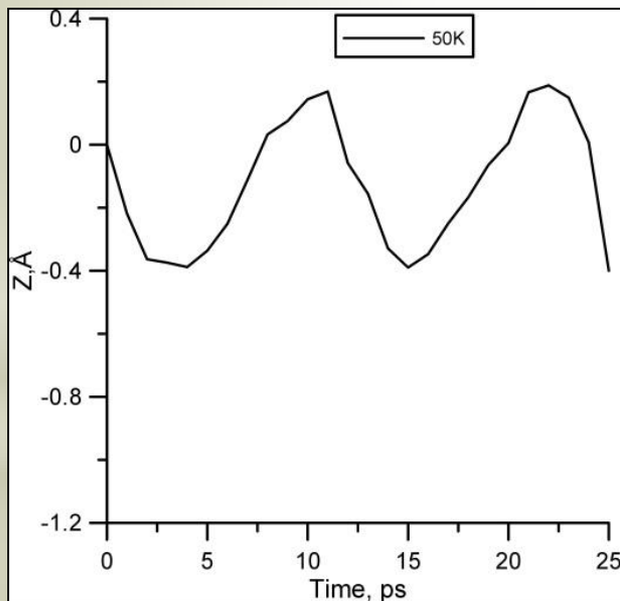
- * Color image of the potential well for free charged C60; field image of the center of gravity for charged C60. Only one attached to the CNT wall fullerene closest to the free fullerene is shown here.

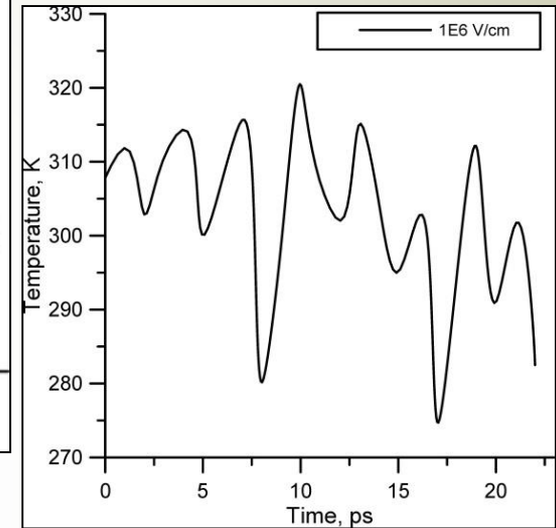
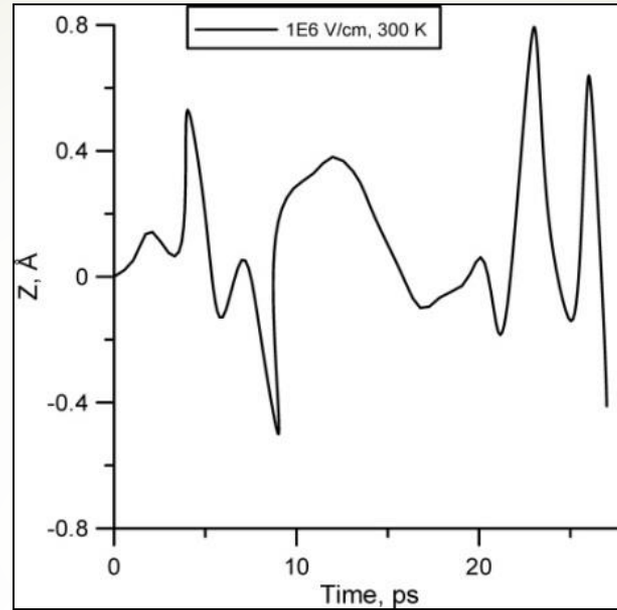
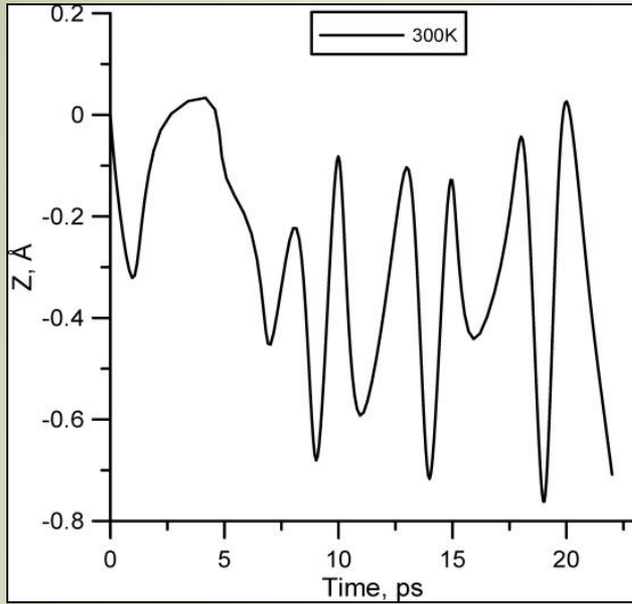
* C_{60}^+ oscillations in the potential well at $T = 50$ K:

a) the position of the gravity center without electric field;

b) the position of the gravity center in the external electric field with the strength of $1 \text{ V}/\mu\text{m}$;

c) the change of the system temperature

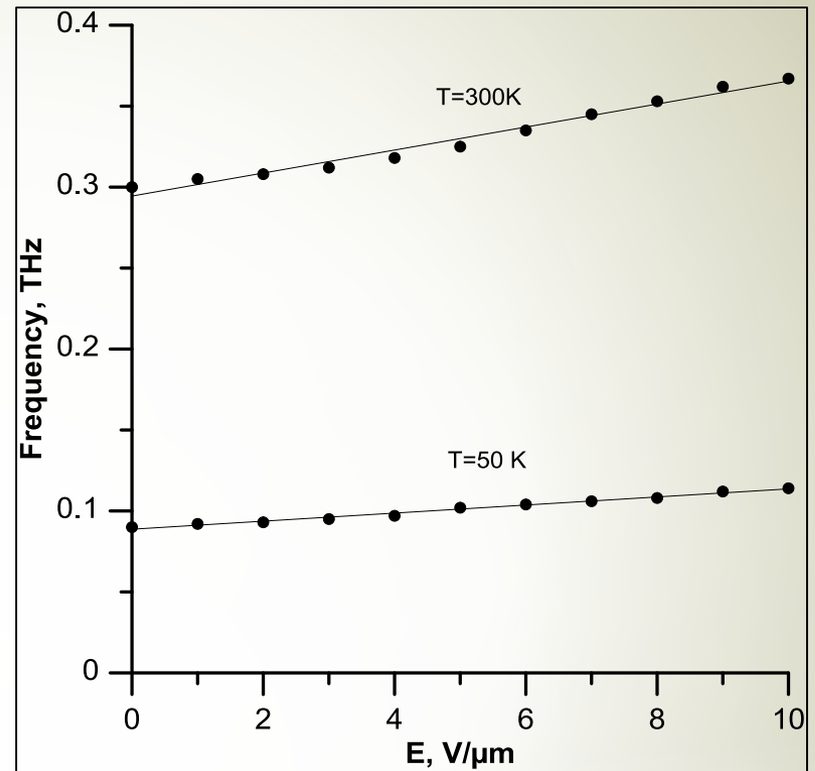
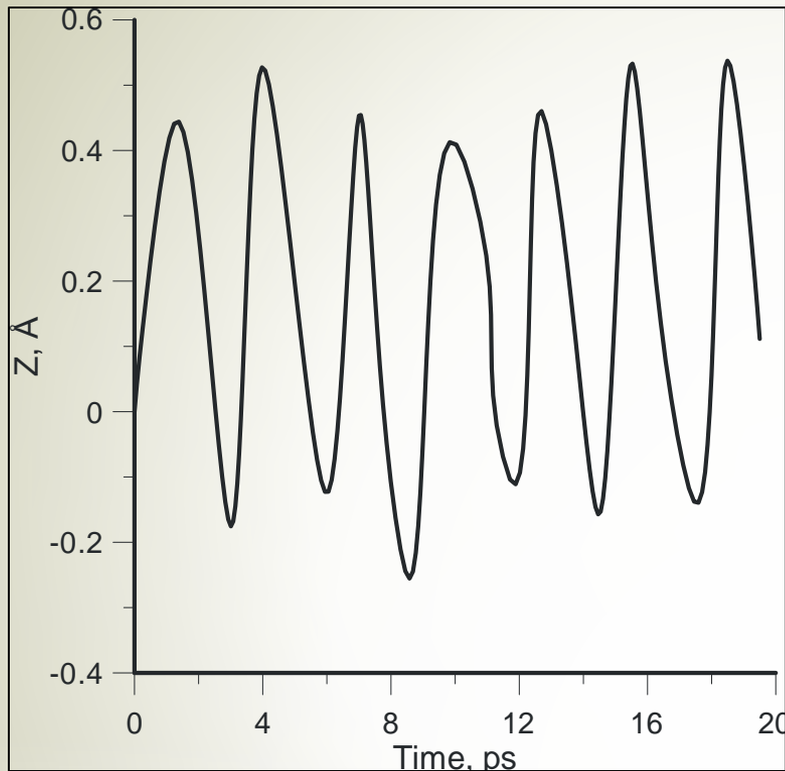




* **C₆₀⁺ oscillations in the potential well at $T = 300$ K:**

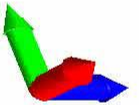
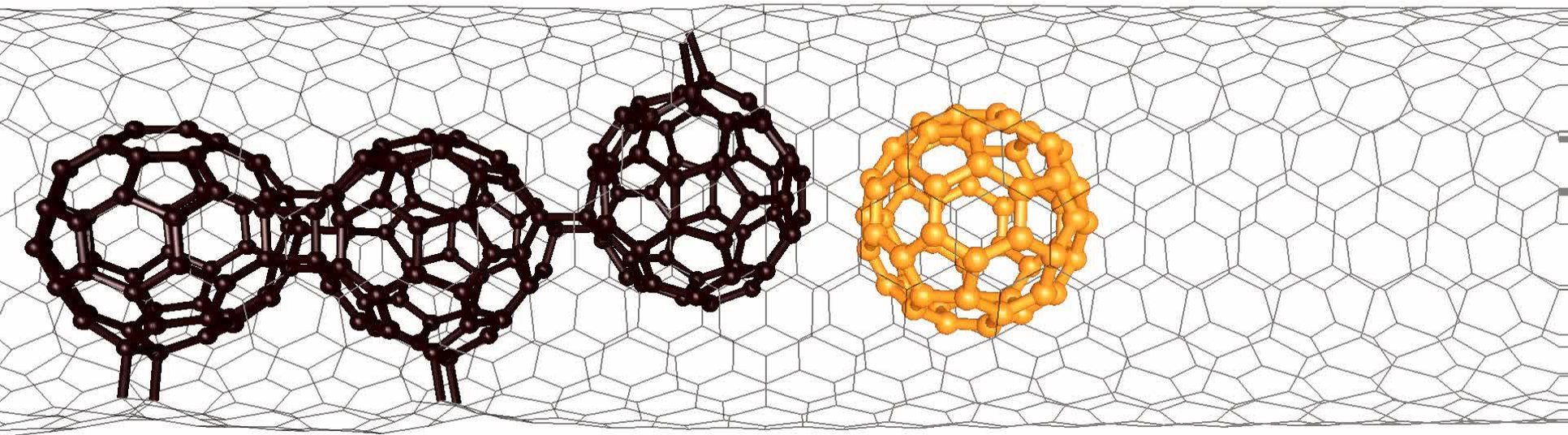
- a) the trajectory of the center of gravity without field;
- b) the trajectory of the center of gravity in external electric field with the strength of $1 \text{ V}/\mu\text{m}$;
- c) the change of the system temperature

The C₆₀⁺ oscillations in the GHz range are found to be stable at 50 K, while after the temperature increase to 300 K the C₆₀⁺ oscillation frequency falls in the THz range.



- * 1) Position of the gravity center of C60 + oscillating in the potential well under the external field with the strength of $10 \text{ V}/\mu\text{m}$
- 2) Oscillation frequency versus intensity of the electric field strength.

The oscillations are generated only at the external electric field of $10 \text{ V}/\mu\text{m}$. We also demonstrated the experimental possibility to synthesize such kind of structures by hydrogen annealing of the carbon nanopeapods.

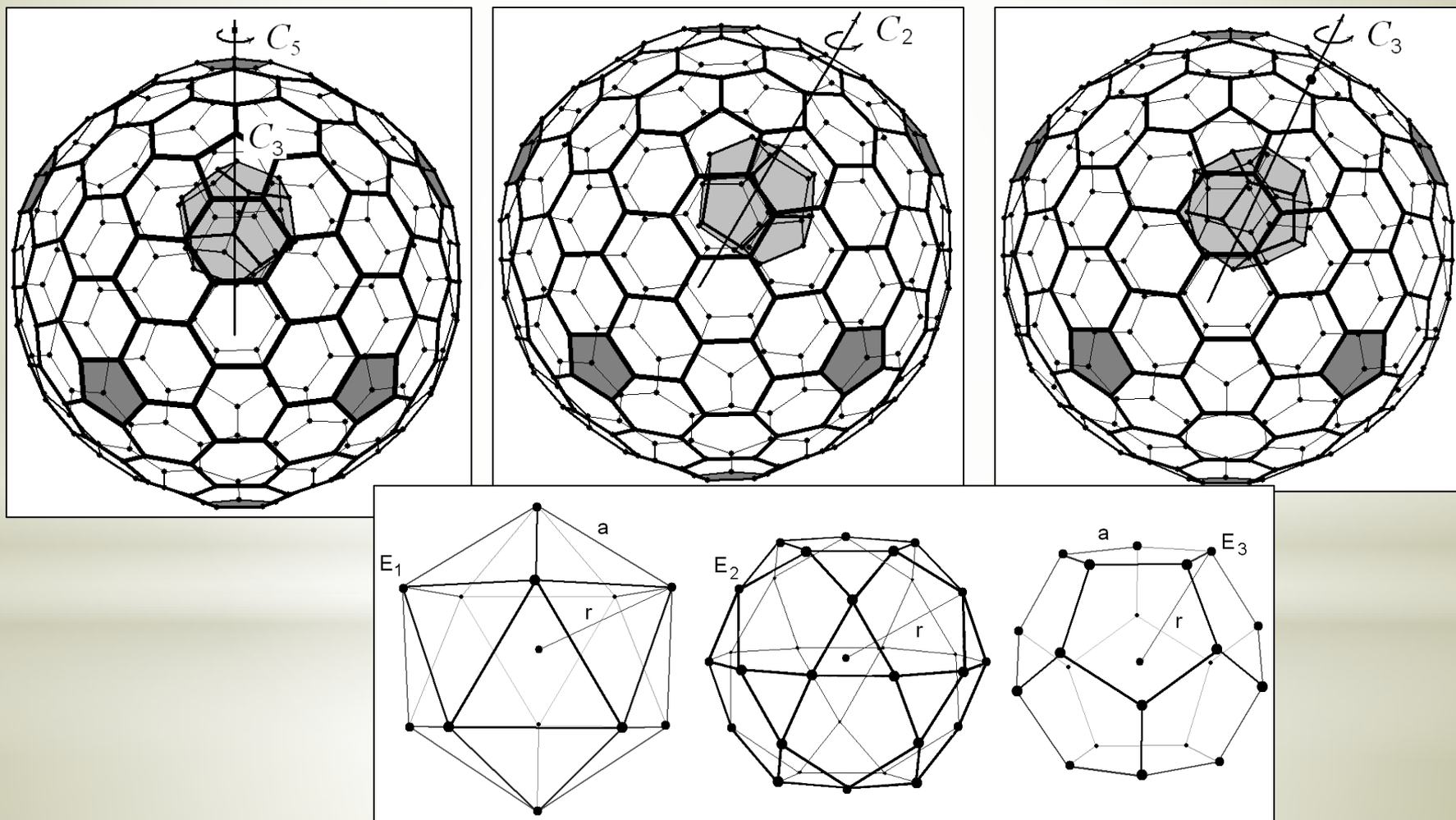


*The radiation

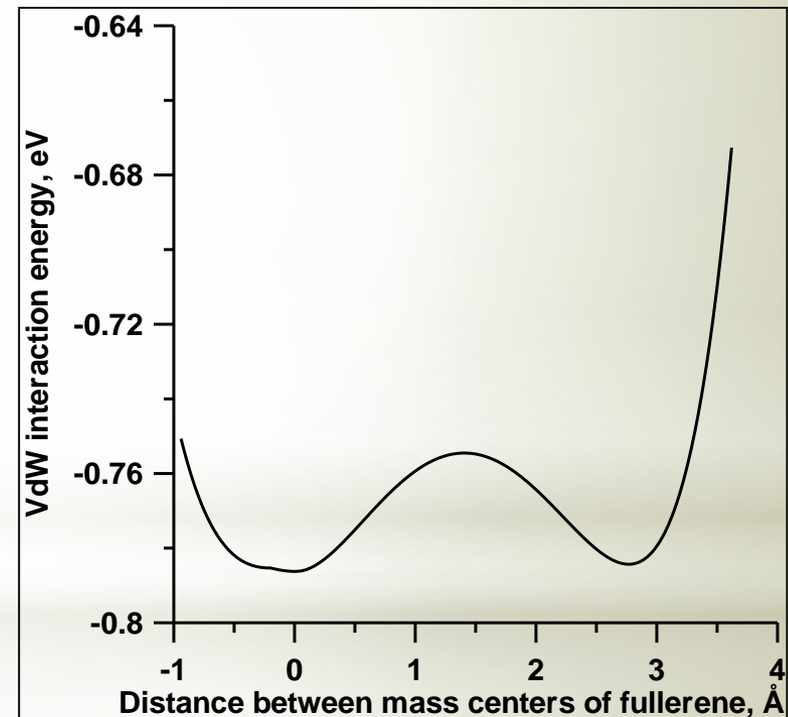
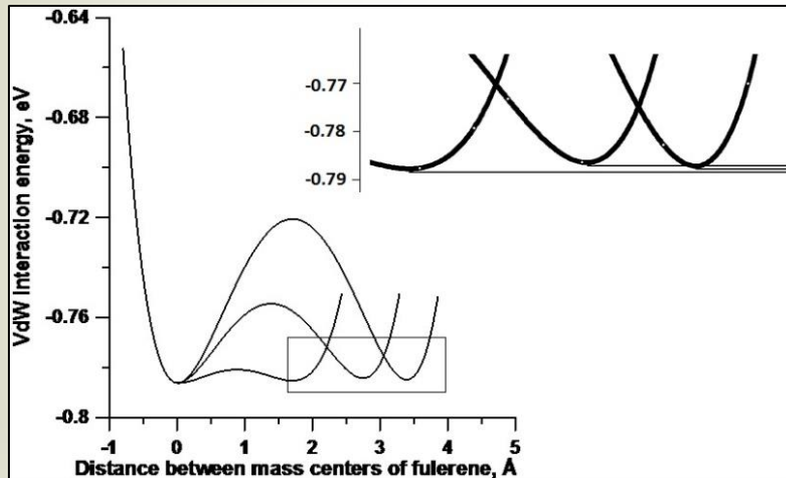
* The theoretical investigation
of bilayer fullerenes C60@C540
and C20@C240

O.E. Glukhova, A.S. Kolesnikova, M.M. Slepchenkov,
V.V. Shunaev Moving of Fullerene Between Potential
Wells in the External Icosahedral Shell // J. Comput.
Chem. 2014. 35(17):1270-7.

* Positioning of the C₂₀ in the field of fullerene C₂₄₀ retaining potential a) for interaction energy E_1 b) for energy E_2 , c) for energy E_3 .



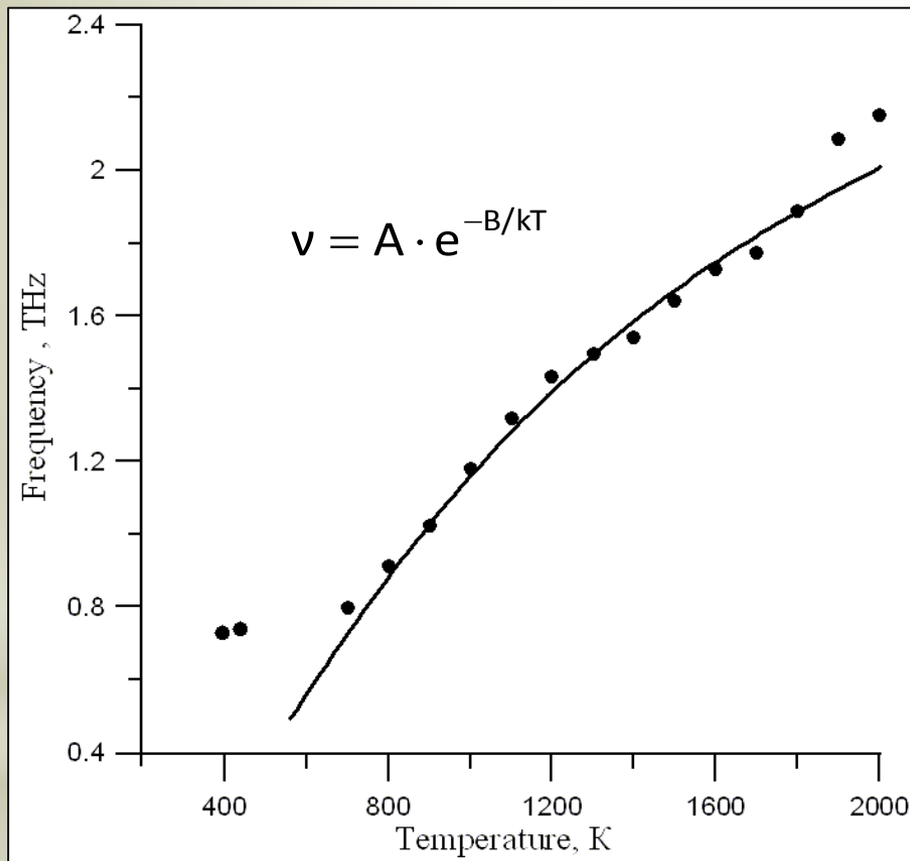
* Surface relief of the van der Waals interaction energy between C20@C240 nanoparticle layers at different variants of C20 moving. a) from the well with energy E1 to the same well, from the well with energy E1 to the well with the energy E2 from the well with energy E1 to the well with the energy E3; b) from the well with the energy E2 to the well with the energy E3.



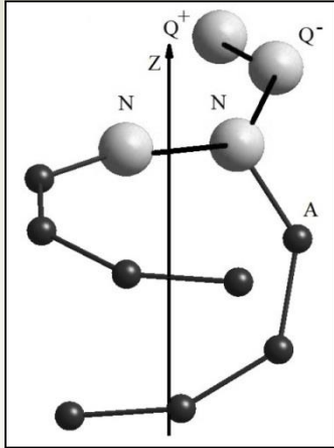
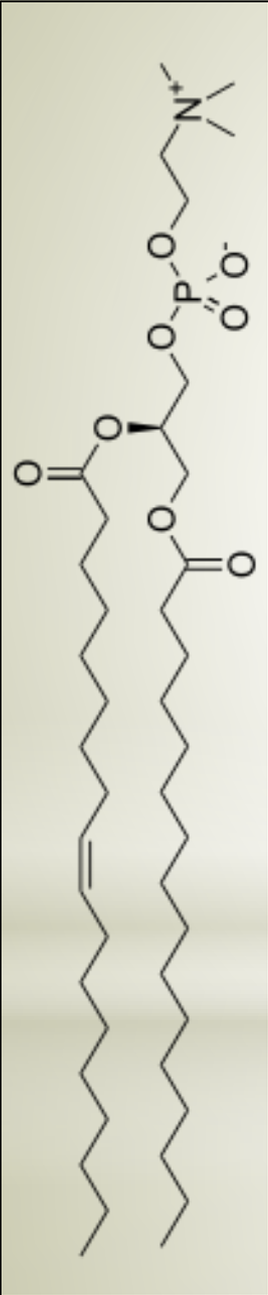
Geometry and Energy Parameters for Fullerenes of Nanoparticle C₂₀@C₂₄₀

C _n	E _g , eV	P _{ioniz} , eV	E _b , eV	ΔH, kcal mol · atm	R, Å	Min/max Bond length, Å
C ₂₀ (D _{3d})	2.88	6.71	6.34	25.10	2.06	1.43/1.51
C ₂₄₀ (I _h)	1.29	7.06	7.17	5.99	7.20	1.39/1.44

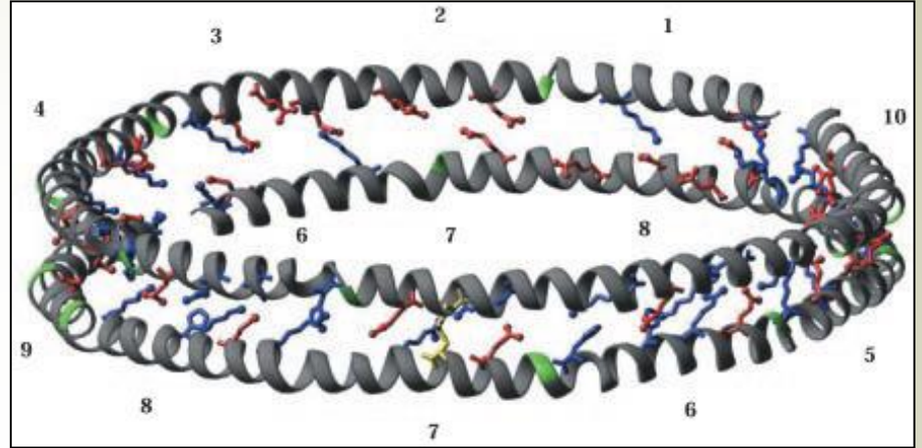
* Dependence of jumping frequency on temperature in the case when fullerene C₂₀ jumps from one potential well to another in the field of C₂₄₀ keeping potential.



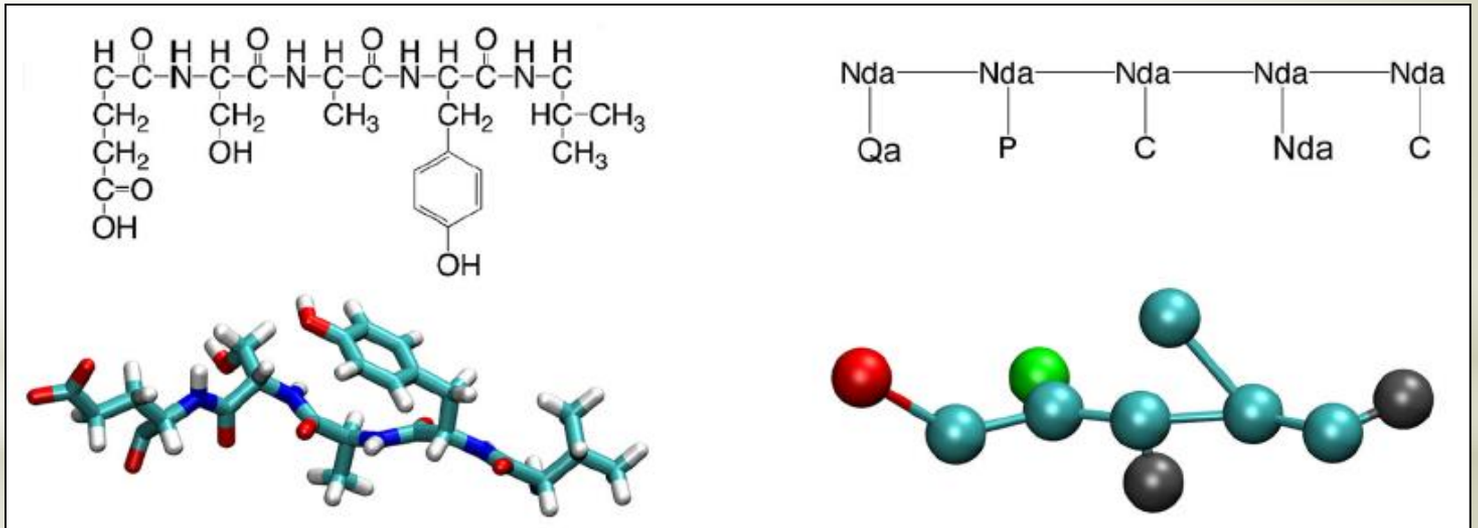
Nannclus- ters with non-central effect	C ₂₀ @ C ₁₈₀	C ₂₀ @ C ₂₄₀	C ₂₀ @ C ₅₄₀	C ₆₀ @ C ₅₄₀	C ₈₀ @ C ₅₄₀
$\Delta H_{\text{formation}}$ (C _n @C _m) kcal/mol	-40.29	-25.72	-12.33	-2.00	-71.78
E ₁ , eV	-1.734	-1.126	-0.869	-1.972	-2.569
E ₂ , eV	-1.700	-1.114	-0.665	-1.691	-2.288
E ₂ - E ₁ , K	394.6	139.2	2367.6	3261.3	3261.3
E ₃ , eV	-1.699	-1.113	-0.658	-1.643	-2.270
E ₃ - E ₂ , K	11.6	64	81.2	557.1	208.9



Lipid molecular (POPC)

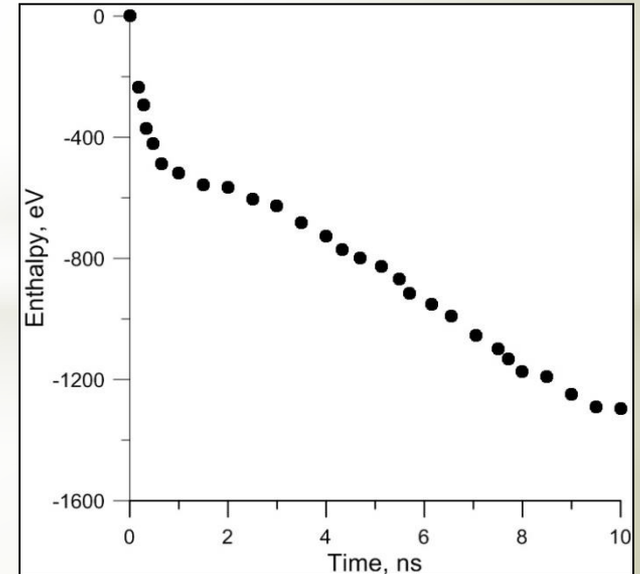
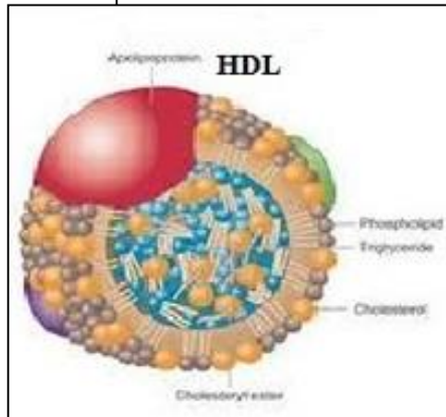
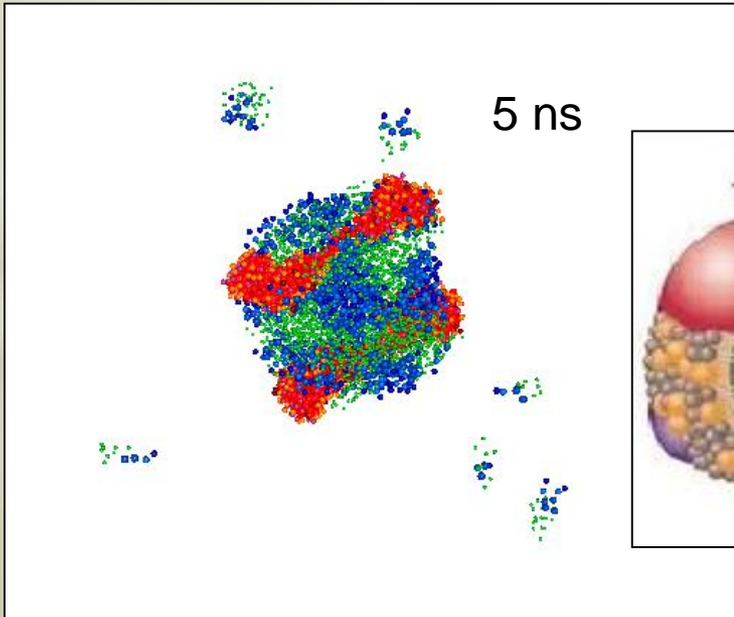
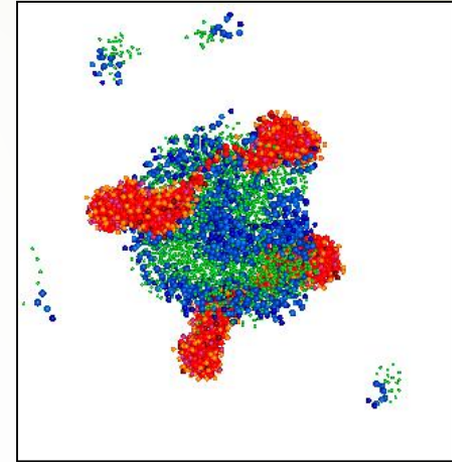
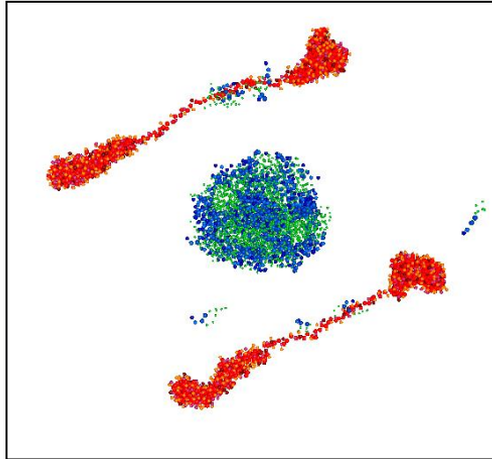
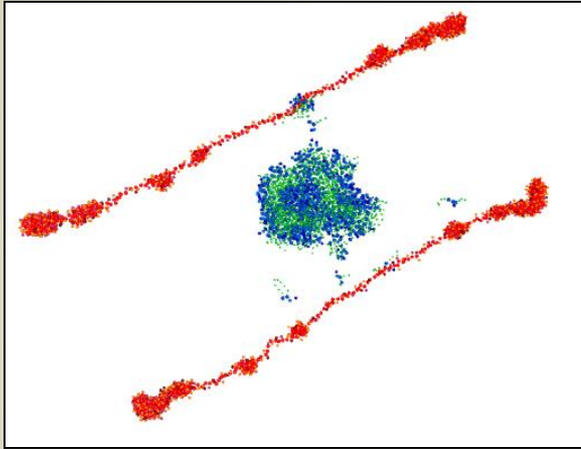


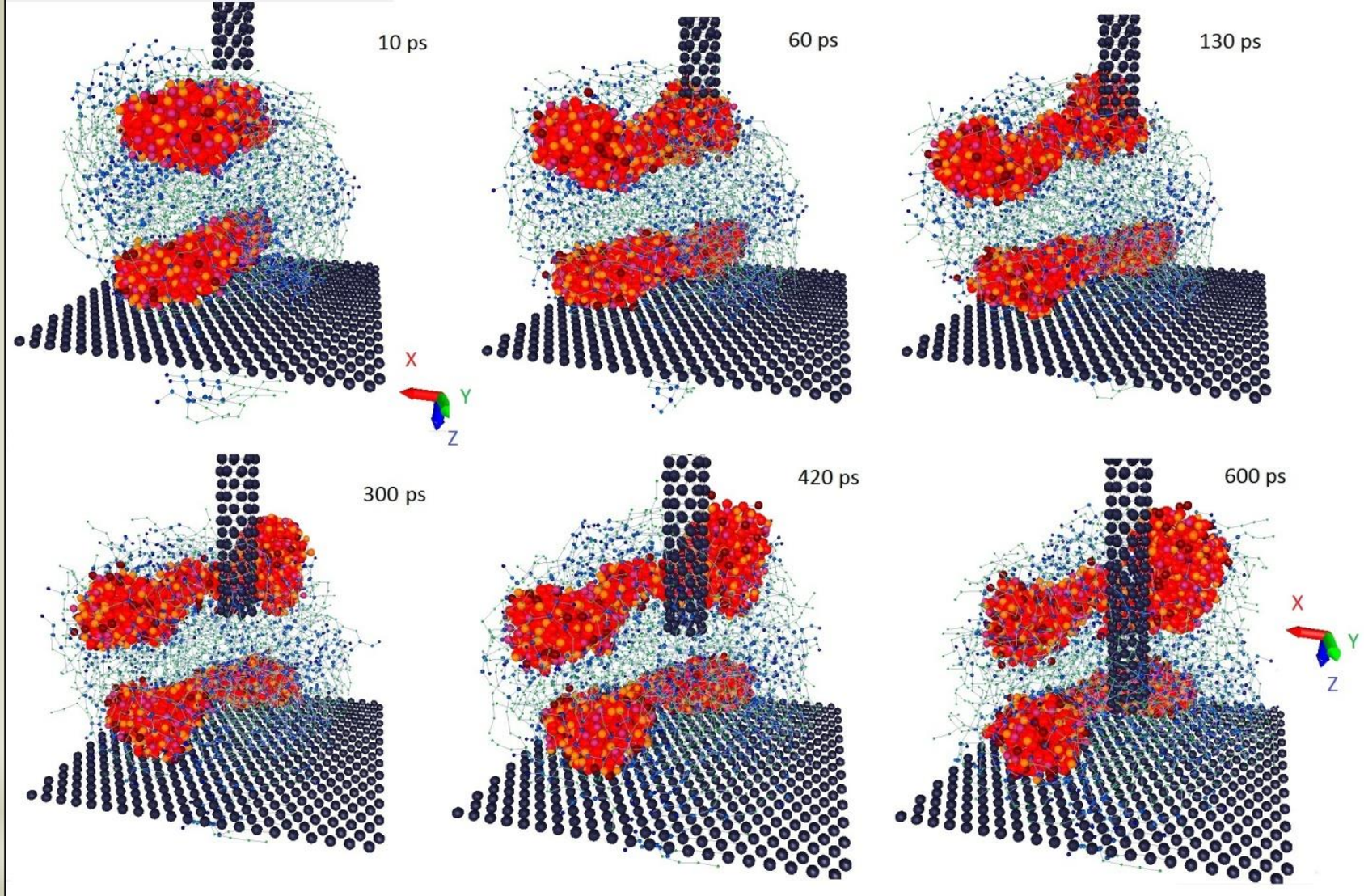
Two belts of Apo A-I



* CG-models

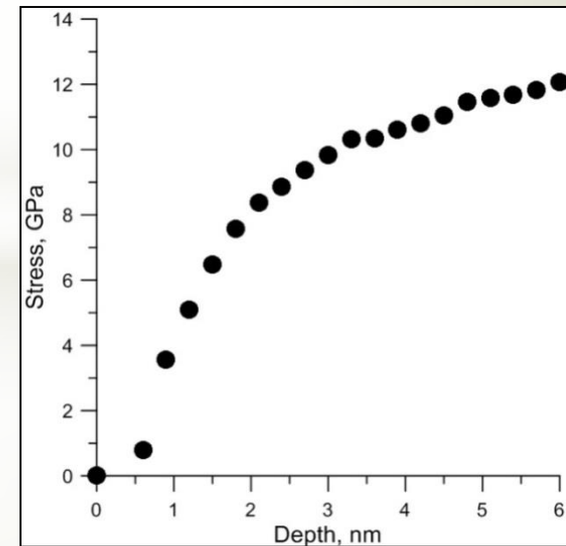
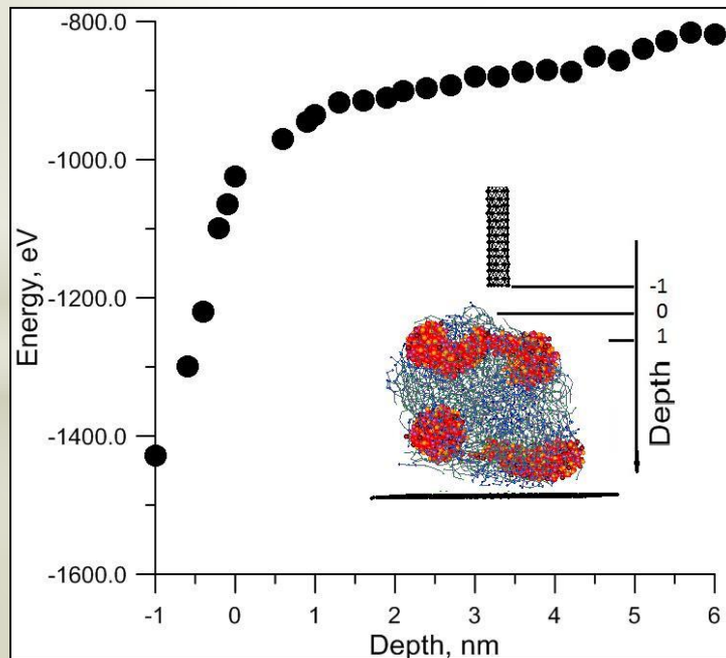
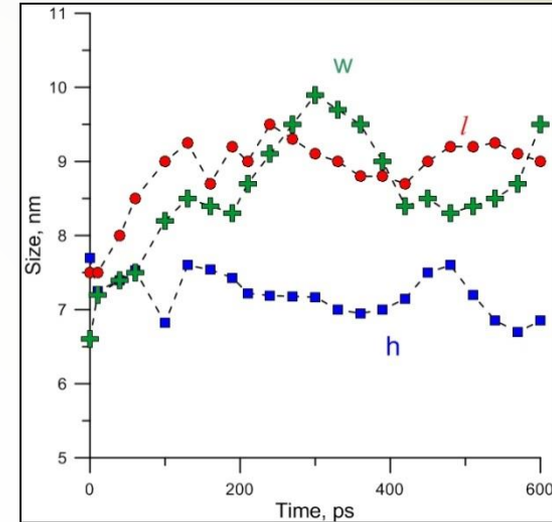
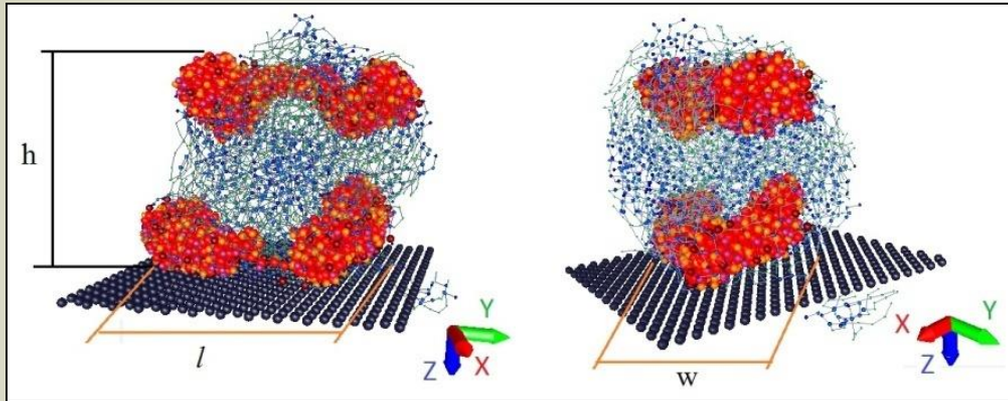
* self-assembled lipoprotein HDL

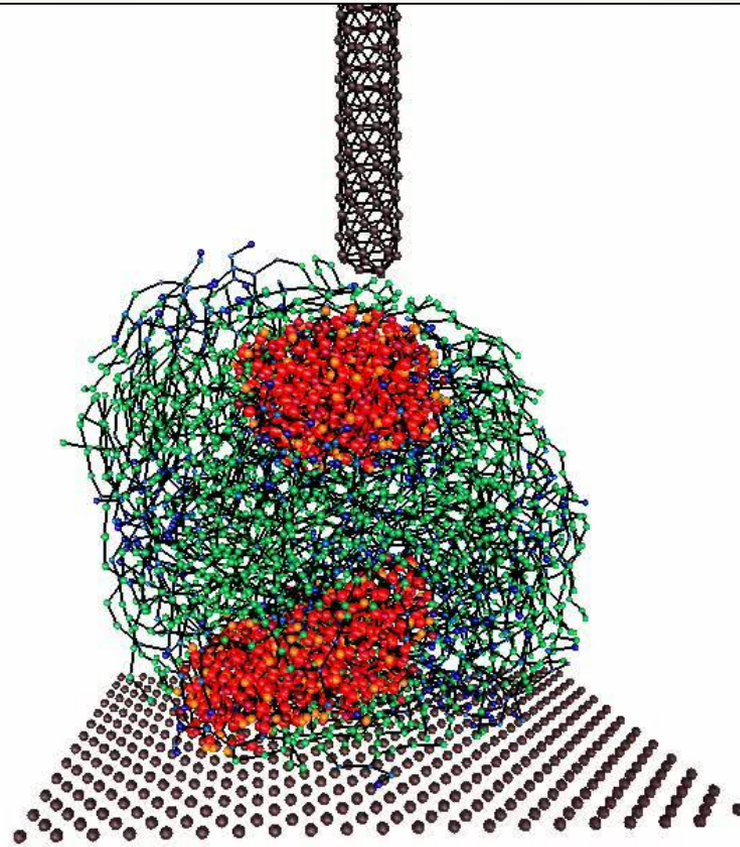




* Indentation of HDL by open CNT

* Interaction of HDL with open CNT



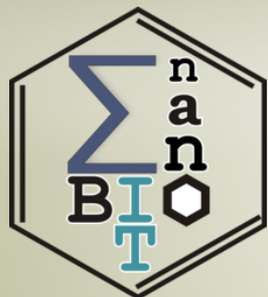
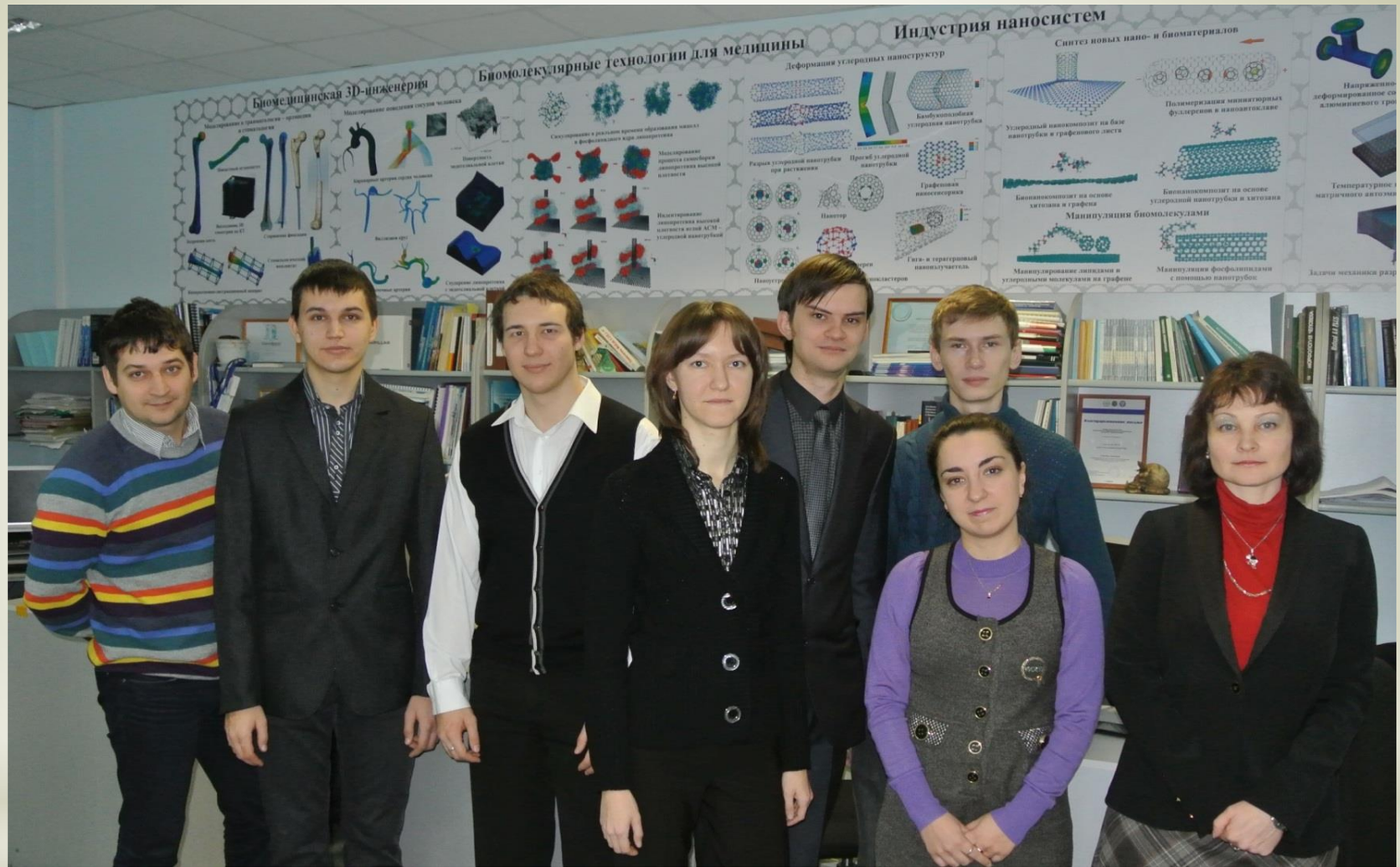


* Interaction of HDL with closed CNT

* In Russia

- * The Bogoliubov Laboratory of Theoretical Physics (BLTP), Joint Institute for Nuclear Research (Dubna, Moscow Region)
- * National Research University of Electronic Technology (Moscow)
- * The Institute of Radioengineering and Electronics (IRE) of Russian Academy Science (Moscow)
- * Outside Russia
- * NCTS, National Cheng Kung University, Taiwan
- * Institute of High Performance Computing, A*STAR, Singapore
- * Physics Department, Aalto University, Finland
- * Chapman University, Orange, CA (USA)

* **COLLABORATION**



Thank you for attention!