Silicon-Doped Nanotube (Si-CNT) as Drug Delivery Nanocarrier for Osteoporosis Drug: Quantum Chemical Study

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Silicon-Doped Nanotube (Si-CNT) as Drug Delivery Nanocarrier for Osteoporosis Drug: Quantum Chemical Study

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Abstract

The aim of this study is to investigate the potential and capability of Si-CNT to detect and adsorb BMSF-BENZ ((4-bromo-7-methoxy-1-(2-methoxyethyl)-5-\{3-(methylsulfonyl)phenyl\methyl\}-2-[4-(propane-2-))yl) phenyl]-1H-1,3-benzothiazole) molecular. For this purpose, we considered different configurations for adsorbing BMSF-BENZ drug on the surface of the Si-CNT nanocluster. All considered configurations are optimized using DFT theory at the 6-31G** basis set and B3LYP level of theory, and then from optimized structures, for each nanoparticle, we selected four stable models for the adsorption of BMSF-BENZ from (Br, N8, N9, N58, O35, O41 and S) active sites on the surface the selected nanoparticle. and Quantum theory of atoms in Molecular Analysis (QTAIM), and Molecular Orbital Analysis (MO) was also established.

The calculated results indicate that the distance between nanocluster and drug from the N8 site is lower than from all other locations sites for all investigated nanoparticles, and adsorption of BMSF-BENZ from the N8 site is more favorable especially for the Si-CNT nanotube.

The adsorption energy, hardness, softness, and fermi energy results reveal that the interaction of BMSF-BENZ with Si-CNT, is an encouraging adsorbent for this drug as Eads of BMSF-BENZ/Si-CNT complexes are (-5.15, -24.21, -8.22, -17.03, -13.16, -2.22, -12.70) kcal/mol in the gas phase. As well, the appropriate and spontaneous interaction between the BMSF-BENZ drug and Si-CNT nanoparticle was confirmed by investigating the quantum chemical molecular descriptors and solvation Gibbs free energies of all atoms.

Si-CNT can be used as an amperometric sensor to detect the BMSF-BENZ drug molecule. Thus, we propose that the Si-CNT can be a promising candidate as a drug delivery vehicle for BMSF-BENZ drug molecules.
Keywords
BMSF-BENZ; Drug adsorption; Si-CNT nanotube; Drug delivery system; Density functional theory; Thermodynamic properties;

1. Introduction
Osteoporosis is related to a decrease in bone mass and damage to the microstructure of bone tissue, which leads to an increased risk of bone fractures as a result. [1]
Several recent studies [2] have shown that one in every three women and every 12 men will suffer an osteoporotic fracture at some point in their lives. Also, 50,000 forearm fractures, 40,000 symptomatic spinal fractures, and 60,000 hip fractures have been identified to occur in the UK every year[3].
Therefore, it has become necessary to develop new strategies to treat or prevent osteoporosis in men and women, such as the discovery of a drug that enhances the determination of CaSR antagonists, that is, it is rapidly absorbed (short $T_{\text{max}}$) and is eliminated fairly quickly in addition to enhancing the patient's bone mass or improving the way and delivery time of the drug to the patient, ensuring that the drug is delivered safely to the intended target[4].
Based on its appropriate and practically validated chemical and physical properties[4], one of the BMSF-BENZ derivatives used in the treatment of osteoporosis was selected in our article.
The most important properties of this compound with the chemical structure (4-bromo-7-methoxy-1-(2-methoxyethyl)-5-[[3-(methylsulfonyl)phenyl]methyl]-2-[4-(propane-2-))yl] phenyl]-1H-1,3-benzothiazole) (Figure1) is its ability to exhibit peak releasing hormone (PTH) and novel properties with respect to the CaSR receptor[4].
With the growing development of nanoscience, the facts of nanostructures like nanosheets, nano components, nanotubes, and nanocells were extensively studied as possible candidates in many different fields [5-9]. Whereas, nanotubes are best suited as drug carriers because of their high sensitivity to drug molecules, low side effects, unique surface properties, atomic structures, etc. [10-12].
Due to the wide scope of use of silicon-saturated carbon nanotubes in many fields, there have been many literature studies and scientific articles involving the use of this type of nanotube especially in relation to drug delivery system, so we will provide some literature reviews in the next paragraph.

### 1.1 Literature reviews

In 2012, Sazan M. Haidary et al. introduced a review on Nanoporous Silicon as Drug Delivery Systems for Cancer Therapies, and they showed that Porous silicon nanoparticles (pSi) had been established as excellent candidates for medical applications as drug delivery devices due to their excellent biocompatibility, biodegradability, and high surface area[13].

Moreover, in 2013, PA Gowri Sankar and Udhayakumar Kaithamalai conducted a comparative (DFT) study on the electronic properties of the zigzag boron silicon (10,0) single-walled carbon nanotubes, on gas molecular adsorption using nine dominant gas molecules (H₂, H₂O, O₂, CO, CO₂, NO, NO₂, NH₃, and CH₃OH) to exploit their potential applications as gas sensors. They found that all molecules are weakly adsorbed on the original carbon nanotube adsorbent with small binding energy, while they can be either charge donors or acceptors of the nanotubes. Also, they showed that the adsorbents B-CNT and Si-CNT are highly chemicals when they were exposed to NO, NO₂, O₂, and CH₃OH.
The Si-CNT adsorbent adsorbs $\text{H}_2$, $\text{H}_2\text{O}$, $\text{CO}$, $\text{CO}_2$, and $\text{NH}_3$ gas molecules moderately with binding strength compared to the B-CNT adsorbent.

F.J. Martínez-Vázquez et al. published an article in 2013 dealing with the fabrication of novel Si-doped Hydroxyapatite/Gelatine scaffolds at room temperature by fast prototyping for drug delivery system and bone regeneration, and they proved that the scaffold behaves as a hydrogel palliative the fitting to the bone defect at the time of surgery. These scaffolds show a sufficient macropore design for vascularization and a microporosity that enables fluid exchange, and the compressive conduct of such scaffolds is similar to that of trabecular bone of the same density.

Moreover, in vitro cellular investigations acquired in the well-established osteoblastic cell line $\text{MC}_3\text{T}_3$-$\text{E}_1$ exposed various benefits of gelatin inclusion in the material regarding cell differentiation and gene expression. Finally, the investigation indicated that these composite scaffolds are sufficient to be considered as matrices for bone regeneration and drug delivery[15].

In 2018, Rayeheh Bagheri et al. introduced a DFT study about Si-doped phagraphene as a drug carrier for adrucil anti-cancer drugs. Based on the results, a drucil drug very weakly interacts with phagraphene. However, after replacing one carbon atom with different atoms like Al, Si, and B, the results showed that the new complex of drug/Si-CNT is more appropriate for adrucil delivery because of moderate adsorption energy, short recovery time, and the electronically harmless adsorption property [16].

The binding energy values of the Si-CNT suggest that doping of nanotube enhances the interaction mechanism and alters the structural, chemical, and electronic properties of the complexes. When the Si-doped atom interacts with the active atoms of the BMSF-BENZ (Figure 2), the new complexes become more stable, leading the binding energies to lie in the range of chemisorption [17].

![Fig.2](image-url) location of the active atoms of BMSF-BENZ where Si-doped CNT was interacting: (1) Bromine atom, (2) Nitrogen $\text{N}_8$ atom, (3) Nitrogen $\text{N}_9$ atom, (4) Nitrogen $\text{N}_{58}$ atom, (5) Oxygen $\text{O}_{55}$ atom, (6) Oxygen $\text{O}_{61}$ atom, and (7) Sulfur $\text{S}$ atom.
To the authors' knowledge, no comparative investigation of BMSF-BENZ adsorption on Si-CNT(4,0) has been previously performed. The main objective of this article is to explore the adsorption behavior of BMSF-BENZ on the zigzag surface of Si-CNT(4,0)SWCNT. For this purpose, we have a comparative investigation of various structural and electronic properties before and after adsorption of BMSF-BENZ onto the nanotube.

2. Computational methods

All calculations were carried out by using Density Functional Theory (DFT) with B3LYP and 6-31G** basis set as employed in Gaussian 09 package [18]. In this work, the vibrational frequencies for all new complexes with the original compounds were analyzed and investigated to check the true global minima of our predicted Si-CNT Nanotubes. Adsorption energy, change in energy gap, charge transfer analysis, dipole moment, and recovery time were also examined to predict the interaction between the drug molecule with nanoparticles.

In order to obtain electrical properties of complexes, the energies of $E_{\text{HOMO}}$ and $E_{\text{LUMO}}$ correspond to the highest occupied molecular orbital (HOMO) and the lowest occupied molecular orbital (LUMO) level, respectively were computed. Also some physico chemical parameters based on $E_{\text{HOMO}}$ and $E_{\text{LUMO}}$ energies, such as, electronegativity ($\chi$), Fermi energy level ($E_f$), energy gap ($E_g$) can be calculated by the following equations : [19-21]

$$\chi = -\frac{1}{2} (E_{\text{HOMO}} + E_{\text{LUMO}})$$ .................................................................(1)

$$E_f = E_{\text{HOMO}} + \left(\frac{E_{\text{LUMO}} - E_{\text{HOMO}}}{2}\right) = -\chi$$ .........................................................(2)

$$E_g = E_{\text{LUMO}} - E_{\text{HOMO}}$$ .................................................................(3)

Several calculations were executed to calculate the total energies of the molecules depending on the position of the nanotube attached to the drug molecule. The adsorption energies of drug on the surfaces of the nanotubes were obtained by: [22]

$$E_{\text{ads}} = E_{\text{complex}} - (E_{\text{nanotube}} + E_{\text{drug}})$$ .........................................................(4)

where $E_{\text{complex}}$, $E_{\text{nanotube}}$ and $E_{\text{drug}}$ donate the energy of the complex composed with nanotubes–drug and isolated energies of nanotube and drug, respectively. The $E_{\text{ads}}$ energy was determined from the summation of the interaction energy ($E_{\text{int}}$) and deformation energies ($E_{\text{def}}$) of the drug ($E_{\text{def-drug}}$) and nanotubes ($E_{\text{def-nanotube}}$) during the adsorption process. [23]

$$E_{\text{int}} = E_{\text{complex}} - (E_{\text{nanotube in complex}} + E_{\text{drug in complex}})$$ .........................................................(5)
\[ E_{\text{def}} = (E_{\text{drug in complex}} - E_{\text{drug}}) + (E_{\text{nanotube in complex}} - E_{\text{nanotube}}) \] ..............................(6)

where  \(E_{\text{nanotube in complex}}, E_{\text{drug in complex}}\) are the energies of nanotube and drug with their geometries in the complex, respectively.

The thermodynamical parameters were also investigated such as changing in Gibbs free energy (\(\Delta G\)), entropy (\(\Delta S\)), and enthalpy (\(\Delta H\)), to examine the structural stability using the following equations:[24]

\[ \Delta G = G_{\text{complex}} - G_{\text{nanotube}} - G_{\text{drug}} \] ..............................(7)

\[ \Delta H = H_{\text{complex}} - H_{\text{nanotube}} - H_{\text{drug}} \] ..............................(8)

\[ \Delta S = \frac{\Delta H - \Delta G}{T} \] ..............................(9)

where  \(G_{\text{complex}}\) and  \(H_{\text{complex}}\) are the Gibbs free energy and the enthalpy of drug adsorbed upon nanotubes,  \(G_{\text{nano}}\) and  \(H_{\text{nano}}\) are the Gibbs free energy and enthalpy of the nanotubes and  \(G_{\text{drug}}\) and  \(H_{\text{drug}}\) are the Gibbs free energy and enthalpy of the drug respectively. (T) is the room temperature equal to (298.15K).

The nature of interactions between BMSF-BENZ and Si-CNT has been determined Quantum Theory of atoms in molecules (QTAIM) analysis of Bader’s [25] which is implemented in MULTIWFN program [26]. The molecular orbital (MO) analysis has been evaluated by MULTIWFN 3.7 program using density of states (DOS). Recovery time has predicted to understand the desorption process of the drug molecule from Si-CNT surface when the adsorbents in the gas phase.

3. Results and discussions

3.1. The optimized geometry of the adsorbents

An adsorbent nanotube of zigzag Si-doped SWCNT (n,0) was considered for the BMSF-BENZ drug as an osteoporosis disease drug delivery vehicle. Initially, the nanotube and BMSF-BENZ molecule were optimized using DFT calculation in the gas media. The optimized geometries of adsorbent Si-CNT and drug molecules with their corresponding HOMO, LUMO, are shown in Figure 3.
The vibrational modes for the new complexes and the original compounds were calculated to check the structural stability in the presence of IR. Where all compounds give vibrational modes with a positive frequency range of (8.39, 10.12, 6.92, 6.71, 10.43, 12.02, 4.24) cm\(^{-1}\) as a minimum value, and (3241.55, 3249.13, 3229.78, 3244.58, 3241.58, 3244.41, 3236.36) cm\(^{-1}\) as a maximum value for the active atom binding (Br, N\(_8\), N\(_9\), N\(_{58}\), O\(_{35}\), O\(_{41}\), and S) with Si-CNT nanotube (Figure 2). Furthermore, the HOMO and LUMO energies were also calculated to find the energy gap of nanotubes.

HOMO levels are located on the center of the C–C bonds throughout the nanotube, whereas LUMO levels were located all over the nanotube structure. The predicted energy gap of Si-CNT (1.66, 1.54, 1.63, 1.37, 1.68, 1.64, 1.66) eV for all locations, respectively.

### 3.2. BMSF-BENZ adsorption on Si-CNT nanotube

To detect suitable adsorbents for BMSF-BENZ, we attempted to verify various properties like geometric, electronic, adsorption properties, etc., of BMSF-BENZ/Si-CNT complexes. Initially, Si-CNT was exposed to the drug compound at different adsorption sites and the preferred adsorption sites for Si atoms were found as it approached (Br, N\(_8\), N\(_9\), N\(_{58}\), O\(_{35}\), O\(_{41}\), S) atoms of BMSF-Benz (Fig. 2).

In the case of the BMSF-BENZ adsorbed compound, the Si-CNT geometry remains unchanged, and the lengths of the C-C bonds varied by about 0.03 Å at the proximal adsorption site. It claims that there is an interaction between BMSF-BENZ and Si-CNT nanotubes, and this was confirmed by the adsorption energy and changes in the electronic properties analysis except when the nanotube approached the N\(_9\) nitrogen atom, we observed that the bond length was the same before and after adsorption which It means that there is no interaction between the nanotube and the drug compound at this point.
The BMSF-BENZ drug molecule was adsorbed on the Si-CNT nanotube with an adsorption energy of about (-5.15, -24.21, -8.22, -17.03, -13.16, -2.22, and -12.70) kcal/mol at a distance of at least about (3.27, 1.91, 8.32, 1.94, 1.90, 3.91, and 2.37) Å in the B3LYP method as shown in Table 1. Due to the proximity of the bromine atom to the interaction region between the nanoparticle and the nitrogen atom (N₉), the bromine atom forced the silicon atom to move away from the nitrogen atom and prevented it from forming a bond between them, but instead, hydrogen bonds were formed between two hydrogen atoms in the nanoparticle with an atom Nitrogen in the drug compound and with spaces (1.91, 1.85) Å respectively.

In order to analyze the adsorption and desorption process together, the charge Q(e) values for Si-CNT by adsorption process and recovery time were obtained Under vacuum UV light conditions with a frequency of (3 × 10¹⁶) sec⁻¹ at room temperature for the different complexes in the gas phase shown in Table 1.

Note that the length of the bonds between the silicon atom and the adjacent carbon atoms were (1.80, 1.89, 1.89) Å, respectively, and the distance between the Si-CNT and every active atoms was (2.4) Å.

Table 1. Adsorption energy (E_ads) in (kcal/mol), the distance between BMSF-BENZ molecule, and adsorbents (d) in (Å), Bond length between active atoms of the nanotubes after adsorption of BMSF-BENZ drug for the different complex in gas phase, the DM denotes the dipole moment values of Si-CNT in (Debye) and recovery time (τ) in sec.

| Structure     | d   | E_ads | Bond location | Bond length after adsorption | DM     | τ (sec)   |
|---------------|-----|-------|---------------|-------------------------------|--------|-----------|
| BMSF-BENZ     | -   | -     | -             | -                            | 1.82   | -         |
| Si-CNT        | -   | -     | -             | -                            | 0.64   | -         |
| Br/Si-CNT     | 3.27| -5.15 | Si-C1         | 1.79                          |        |           |
| N₅/Si-CNT     | 1.91| -24.21| Si-C1         | 1.82                          | 2.20   | 1.97×10⁻¹³|
| N₆/Si-CNT     | 8.32| -8.22 | Si-C1         | 1.80                          | 1.70   | 3.49×10⁻⁷ |
| N₅/O₃/Si-CNT  | 1.94| -17.03| Si-C1         | 1.78                          | 3.02   | 9.88×10⁻⁵ |
| O₃/Si-CNT     | 1.90| -13.16| Si-C1         | 1.78                          | 1.63   | 1.45×10⁻⁷ |
| O₂/Si-CNT     | 3.91| -2.22 | Si-C1         | 1.79                          | 0.71   | 1.41×10⁻¹⁵|
Dipole moment refers to the polarity of the compound as it appears due to the asymmetric charge distribution as well as the difference in electronegativity between the resultant molecules or atoms, where the large dipole moment refers to the more polar structure which results in a more stable interaction of the structure [27].

The results of the dipole moment showed that the new compounds obtained from the adsorption of the drug compound on the surface of the nanotube had different values greater than the value that the drug had before adsorption. A more stable reaction occurred when the drug compound was adsorbed on the surface of the nanotube at N\textsubscript{8} atom with the value of (5.92) Debye which was greater than the rest of the other active atoms of the drug compound.

**Quantum Theory of Atoms in Molecules (QTAIM) Analysis**

In order to understand the nature of interactions of BMSF-BENZ drug on Si-CNTs, Topology Analysis based on Bader’s quantum theory of atoms in molecules (QTAIM) method has been employed with MULTIWFN program.

In QTAIM analysis, the nature of intermolecular interactions between drug and CNT is determined in terms of electron densities at bond critical points (BCPs). The electron density of $\rho(r)$ and laplacian $\nabla^2 \rho(r)$ characteristics are widely used to understand the bonding interactions nature. However, the other two parameters of total energy density of $H(r)$ and the ratio of $|V(r)|/G(r)$ are more remarkable parameters used on bonding characteristics. The covalent bonding characteristics correspond to $\nabla^2 \rho(r) < 0, H(r) < 0, |V(r)|/G(r) > 2$. The strong hydrogen bonds as the intermediate type of interaction related to $\nabla^2 \rho(r) > 0, H(r) < 0, 1 < |V(r)|/G(r) < 2$. The weak and medium-strength hydrogen bonding and Van der Waals interactions due to the $\nabla^2 \rho(r) > 0, H(r) > 0, |V(r)|/G(r) < 1$.

In this work, the QTAIM theory has been applied for the N\textsubscript{s}/Si-CNT and N\textsubscript{58}/Si-CNT complexes which are appeared having the first two higher adsorption energy values (in magnitude) given in Table 1. The calculated QTAIM parameters are displayed in Table 2 and the interactions between the drug and Si-CNT for the selected complexes are shown in Figure 4.
Table 2 The QTAIM parameters of selected complexes at the BCPs. Electron density($\rho(r)$), Laplacian of electron density ($\nabla^2 \rho(r)$), the kinetic electron density ($G(r)$) , potential electron energy density ($V(r)$), total electron energy density ($H(r)$), the ratio $|\frac{V_{BCP}}{G_{BCP}}|$, the ratio $|\frac{V_{BCP}}{G_{BCP}}|$. 

| Structure      | BCPs Drug/Si-CNT | $\rho_{BCP}$ | $\nabla^2 \rho_{BCP}$ | $G_{BCP}$ | $V_{BCP}$ | $H_{BCP}$ | $|\frac{V_{BCP}}{G_{BCP}}|$ |
|----------------|-----------------|--------------|----------------------|----------|-----------|-----------|-----------------|
| N8/Si-CNT      | $N_{41}$-Si$_{19}$ | 0.0826       | 0.2943               | 0.1021   | -0.1306   | -0.0285   | 1.2793          |
|                | $B_{50}$-C$_{26}$ | 0.0127       | 0.0371               | 0.0077   | -0.0061   | 0.0015    | 0.7941          |
|                | $B_{50}$-I1      | 0.0074       | 0.0210               | 0.0042   | -0.0032   | 0.983 x 10$^{-3}$ | 0.7694          |
| N58/Si-CNT     | $N_{41}$-Si$_{19}$ | 0.0796       | 0.2479               | 0.0914   | -0.1208   | -0.0294   | 1.3218          |
|                | $H_{37}$-C$_{11}$ | 0.0072       | 0.0244               | 0.0048   | -0.0035   | 0.0013    | 0.7297          |
|                | $H_{37}$-C$_{26}$ | 0.0093       | 0.0312               | 0.0062   | -0.0047   | 0.0015    | 0.7544          |
|                | $H_{24}$-H$_{33}$ | 0.0039       | 0.0137               | 0.0025   | -0.0016   | 0.899 x 10$^{-3}$ | 0.6441          |

It is clear in Table 2, the nature of interaction between $N_{41}$ and Si$_{19}$ atoms is semi-covalent character with the ratio of $|\frac{V_{BCP}}{G_{BCP}}| > 1$ which are for both studied complexes as 1.2793 and 1.3219. Those interactions of $N_{41}$ can be classified as the intermediate type of interaction with ($\nabla^2 \rho_{BCP} > 0$) with ($H_{BCP} < 0$) rule. All remaining interactions in Table 2 exhibit mainly van der Waals type weak and with medium-strength hydrogen bonding due to the ratio of $|\frac{V_{BCP}}{G_{BCP}}| < 1$.

The quantum descriptors denoted by Equations (1-4) are calculated for BMSF-BENZ /Si-CNT complex structures and given in Table 3. In the BMSF-BENZ/Si-CNT structure, the electronic parameters such as $E_{\text{HOMO}}$, $E_{\text{LUMO}}$, $E_g$, and Fermi energies have a noticeable change. The values...
of $E_{\text{HOMO}}$ and $E_{\text{LUMO}}$ of Si-CNT nanotubes before the adsorption of molecules are (-4.91 and -3.30) eV, and after adsorption of BMSF-BENZ drug are mentioned in Table 3. The sensors are related to the variation of their electrical conductance after drug adsorption due to the electron exchange between the drug and sensor. On this line, the CNT sensitivity to the drug is based on the changing of HOMO and LUMO energies with $E_g$.

In order to understand the sensing mechanism of the Si-CNT to drug, the percentage variation of $\% \Delta E_g$ energy gap during the adsorption process is taken into account by the following equation:

$$\%\Delta E_g = 100 \times \left( \frac{E_{g2} - E_{g1}}{E_{g1}} \right)$$ .........(9)

where $E_{g1}$ and $E_{g2}$ are the $E_g$ values of CNT and complex structure, respectively. The percentage change values, of $\%\Delta E_g$ for each structure are given in Table 3. The percent change in the work function as $\%\Delta \Phi$ can be derived by logic in band gap energies given in Eq.(9).

The corresponding frontier molecular orbital map is shown in Figure 5, and the Frontier molecular orbital (HOMO and LUMO) of all investigated complexes is shown in Figure 6. Thus, the BMSF-BENZ drug molecule was significantly affecting the HOMO energy level and LUMO energy level as shown in Table 3.

Table 3. Calculated $E_{\text{HOMO}}$, $E_{\text{LUMO}}$ energies, HOMO-LUMO gap ($E_g$), Fermi level energy ($E_f$), and work function ($\Phi$), the percentage variation of $E_g$ and $\Phi$ respect to bare Si-CNT for the studied complexes.

| Structure   | $E_{\text{HOMO}}$ (eV) | $E_{\text{LUMO}}$ (eV) | $E_g$ | $\%\Delta E_g$ | $E_f$ (eV) | $\Phi$ | $\%\Delta \Phi$ |
|-------------|------------------------|------------------------|-------|-----------------|-----------|-------|-----------------|
| BMSF-BENZ   | -5.40                  | -0.88                  | 4.52  | -3.14           | 3.14      | -     |
| Si-CNT      | -4.91                  | -3.30                  | 1.61  | -4.10           | 4.10      | -     |
| Br/Si-CNT   | -4.62                  | -2.96                  | 1.66  | 3.10            | 3.79      | 3.79  | -7.56           |
| N$_8$/Si-CNT| -3.43                  | -1.88                  | 1.54  | -4.35           | 2.66      | 2.66  | -35.12          |
| N$_9$/Si-CNT| -4.57                  | -2.93                  | 1.63  | 1.24            | 3.75      | 3.75  | -8.53           |
| N$_{58}$/Si-CNT| -3.93                | -2.56                  | 1.37  | -14.90          | 3.25      | 3.25  | -20.73          |
| O$_{35}$/Si-CNT| -4.18               | -2.50                  | 1.68  | 4.35            | 3.34      | 3.34  | -18.54          |
| O$_{41}$/Si-CNT| -4.95               | -3.30                  | 1.64  | 1.86            | 4.12      | 4.12  | 0.49            |
| S/Si-CNT    | -4.08                  | -2.42                  | 1.66  | 3.10            | 3.25      | 3.25  | -20.73          |
As reported energies of $E_{\text{HOMO}}$, $E_{\text{LUMO}}$ and $E_g$ in Table 3, some electronic properties of Si-CNT are affected weakly by the adsorption of the BMSF-BENZ drug. The decreasing order of percentage change values are obtained for the interactions of N$_8$, and N$_{58}$ atoms of the BMSF-BENZ drug with Si-CNT as -4.35%, -14.90%, but for other atoms of (Br, N$_9$, O$_{35}$, O$_{41}$, and S) are about (3.10%, 1.24%, 4.35 %, 1.86%, 3.10%) by increasing.

It is known that the lower $E_g$ values indicate higher electrical conductivity, reactivity, and sensitivity. Therefore, decreasing the $E_g$ through adsorption of the BMSF-BENZ drug molecule indicates that the Si-CNT can detect the drug. Thus, among all studied geometries of complex structures, the highest variation value of $\%E_g$ has been obtained by the N$_{58}$ atom of the BMSF-BENZ interacts with Si-CNT nanotube.

According to the values of percentage variation in the work function of $\%\Phi$, the structure corresponding to O$_{41}$/Si-CNT has the highest value of $\%0.49$ by increasing the conduction electrons.
Figure 6. The Frontier molecular orbital (HOMO and LUMO) of all investigated new complexes; the considered iso value is 0.02 electron/bohr^3.

The recovery time (adsorption time) is one of the important parameters used for both gas sensors and drug delivery systems, which predicts the amount of time required for drug adsorption from the adsorbent, as it is highly related to the adsorption energy, and it is known that a high adsorption reaction needs a high adsorption time and vice versa.

The recovery time is calculated by the equation: [28]

$$
\tau = \frac{1}{v} \exp\left(\frac{-E_{ads}}{kT}\right) \tag{10}
$$

Where T, k, and v are the temperature, Boltzmann’s constant and the attempted frequency, respectively. (k~2 × 10^{-3} kcal/mol K)
As the BMSF-BENZ molecule seems to be adsorbed on the Si-CNT nanotube with maximum adsorption energy, a different recovery time (Table 1) was obtained in vacuum UV-light conditions with frequencies of $3 \times 10^{16}$ sec$^{-1}$ at room temperature. When the adsorption energy is more than 1 eV (23.061 kcal/mol) (in magnitude), a strong chemical interaction between adsorbate and adsorbent is considered to occur, and the adsorption process is then leveled as chemisorption. Therefore, the more negative value of the adsorption energy, the stronger the connection of the system and represents a higher static system, so the adsorption process between drug molecules and the adsorbent is supposed to give negative values of adsorption energy.

It is also known that in drug delivery systems, the adsorbent must have a strong interaction with the drug, and for drug sensors, there must be less energy expenditure [29].

By combining the corresponding recovery time values of those complexes in Table 1 with the percentage variation of work function given in Table 3, it has been evaluated that those complexes cannot be used as amperometric drug sensor applications of BMSF-BENZ drug molecule on Si-CNT nanotube.

Those values indicate that the adsorption process of N$_8$/Si-CNT and N$_{58}$/Si-CNT complexes are in physisorption nature with the desorption time values of 17.9 sec, and $9.88 \times 10^{-5}$ sec, respectively. Among them, the complex of N$_8$/Si-CNT can be extended as a drug delivery system with a reasonable recovery time.

In order to prove the importance of (DOS) calculations in the field of quantum chemistry, researchers interested in this topic have found gradual relationships between the absorption energies between different adsorbents on the surfaces of nanoparticles and the scaling relationships between adsorption energies and interaction transfer barriers that lead to the so-called volcanic activity cutoff. These schemes indicate that the optimal catalyst must have a strong bonding with the molecules and atoms participating in the interaction, which enables these molecules to bond well on the surface of the nanoparticles, as it does not reduce or impede the absorption of products.[30]

The lower Eg values indicate higher electrical conductivity, reactivity, and sensitivity. Therefore, decreased the Eg through adsorption of BMSF-BENZ drug indicates the CNT can detect the drug.

The density of states (DOS) diagram of the complex structure of the N$_{58}$ atom of the BMSF-BENZ interacting with Si-CNT was selected to plot in Figure 7 because of the significant %ΔEg values are given in Table 3, each was also calculated in order to better understand the stability of the system. The change in Eg by DOS analysis can be confirmed as shown in Figure7 by the MULTIWFN program.
It is clear in Figure 7 that the bandgap of the complex is getting narrow concerning the Eg of Si-CNT. The recovery time is related to the adsorption energy, as a high adsorption reaction needs a high adsorption time and vice versa.

It is known that positive values of ΔH and G indicate that the reaction process is endothermic rather than spontaneous. In contrast, negative values of ΔH and G indicate that the reaction is an exothermic and spontaneous process [31] (see Table 4).

![Figure 7. TDOS plot for complex structure of N58-BMSF-BENZ /Si-CNT. The dashed line shows the HOMO energy level.](image)

| Structure | ΔG  | ΔH  | ΔS  | v_{min} | v_{max} |
|-----------|-----|-----|-----|---------|---------|
| 1         | Br/Si-CNT | 7.04 | -4.87 | -0.03 | 8.39 | 3241.55 |
| 2         | N8/Si-CNT | -23.07 | -36.45 | -0.04 | 10.12 | 3249.13 |
| 3         | N9/Si-CNT | 3.25 | -8.32 | -0.03 | 6.92 | 3229.78 |
| 4         | N58/Si-CNT | -18.72 | -29.68 | -0.03 | 6.71 | 3244.58 |
| 5         | O35/Si-CNT | -11.03 | -24.55 | -0.04 | 10.43 | 3241.58 |
| 6         | O41/Si-CNT | -1.88 | -8.53 | -0.02 | 12.02 | 3244.41 |
| 7         | S/Si-CNT | -1.14 | -15.82 | -0.04 | 4.24 | 3236.36 |

The obtained data for ΔH and ΔG were negative values in all the examined complexes except upon adsorption of the drug at the bromine atom, that is, these complexes are dynamically stable, and the most negative value of ΔS means the most ordered compound. Therefore, thermodynamic parameters predict that the investigated complexes are more suitable and thermodynamically stable for BMSF-BENZ drug delivery.

4. Conclusions
In order to obtain a suitable drug delivery method for BMSF-BENZ, we examined the adsorption behavior of BMSF-BENZ on Si-CNT surface using DFT calculations. It was observed that the BMSF-BENZ molecule was adsorbed onto the nanostructures with different adsorption energy depending on the nanotube location of the active drug atoms. The HOMO energy ($E_{\text{HOMO}}$), LUMO energy ($E_{\text{LUMO}}$), HOMO-LUMO energy gap, and Fermi level energies were changed after the adsorption of the BMSF-BENZ drug on the surface of Si-CNT. The BMSF-BENZ drug was adsorbed on the Si-CNT nanotube with an adsorption energy of $(-5.15, -24.21, -8.22, -17.03, -13.16, -2.22, \text{and} -12.70)$ kcal/mol in the gas phase at the B3LYP method, which is more favorable for the drug delivery system and the work function type sensor applications.

Moreover, Gibbs free energy shows that the BMSF-BENZ/Si-CNT formation exhibits spontaneous and favorable adsorption energy. Therefore, we can suggest that the Si-CNT nanotube will be a promising drug delivery medium for BMSF-BENZ drug molecules. Based on the calculated results, it can be considered that the best site for adsorption of BMSF-BENZ on the Si-CNT surface is when the nanotube approaches the nitrogen atom ($N_8$) where the best energy and appropriate recovery time were obtained.

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| Conflicts of interest/Competing interests: | (Not applicable' for that section) |
| Availability of data and material: | (All the required data and calculations included in this research are original and not extracted from previous research, except for some paragraphs in the Introduction section taken as citations.) |
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