

# Sensing 6-Mercaptopurine with a Thiophene-carbazole Copolymer Matrix: A First Principles Approach

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## ABSTRACT

This study discusses the impact of cytostatic drugs, particularly 6-mercaptopurine(6-MP), as emerging pollutants in aquatic ecosystems. Computational analyses using Density Functional Theory analyzed the electronic properties and adsorption behavior of the 3-methoxy thiophene (3-MeOTH) monomer matrix and the 3-methoxy thiophene(3-MeOTH)–n-vinyl carbazole (NVK) copolymer matrix in the presence of 6-MP. Results indicate that 6-MP is adsorbed on the matrices of 3-MeOTH polymer and 3-MeOTH-NVK copolymer with weak Van der Waal forces as indicated by negative adsorption energy and a significant charge transfer. Additionally, 6-MP shows notable sensitivities with both matrices, with 3-MeOTH-NVK demonstrating a 37.8% higher sensitivity compared to 3-MeOTH polymer. Enhanced conductance and peak amplitude enhancements in the presence of 6-MP suggest the potential of these materials for drug sensing applications.

Keywords: Density functional theory; 6-mercaptopurine; 3-methoxy thiophene; N-vinyl carbazole; Emerging pollutants.

### **1. INTRODUCTION**

Recently identified synthetic or natural substances that enter the biological systems of humans and animals with limited regulatory oversight, called emerging pollutants (EPs) pose a challenge in environmental management (Saboorizadeh et al. 2021). Despite their presence from trace levels to high concentrations in aquatic systems, they show significant adverse impact on the ecosystems because EPs lack comprehensive monitoring protocols or legislative frameworks (Mousavi et al. 2020). Among these emerging contaminants, cytostatic drugs used in chemotherapy have become particularly concerning environmental pollutants due to their genotoxic properties and resistance to degradation (Santana et al. 2016). Notably, 6-mercaptopurine (6-MP), a widely utilized cytostatic agent to treat acute lymphocytic leukemia, poses a specific hazard due to its low biodegradability requiring advanced oxidation processes for its effective removal (Ensafi et al. 2012; González et al. 2021). The limited research available underscores the critical need for further investigation into the occurrence, environmental fate and ecological effects of cytostatic drugs. Developing comprehensive studies to assess their presence, concentration and impact on the environment after disposal is crucial in understanding and mitigating the risks they pose to aquatic ecosystems and human health.

The precise identification of anticancer drugs in aquatic environments holds significant importance, driving extensive research efforts towards developing efficient detection materials. Carbon quantum dots are a promising material for detecting 6-MP drug residues. Numerous studies have explored the efficacy of carbon quantum dots across varied sample matrices. Aminefunctionalized and fluorescent carbon quantum dots, multiwall carbon nanotubes-TiO2 hybrids(González et al. 2021), ZnO-graphene capped quantum dots embedded in polymer matrices have been employed as fluorometric sensors for aqueous and biological samples (Agrawal et al. 2021; 2022; Ensafi et al. 2012; Ipte et al. 2023; Mousavi et al. 2020) Additionally, investigations have extended to alternative materials like conjugated structures, namely, Fe<sub>3</sub>O<sub>4</sub>@polypyrrole-Pt core-shell nanoparticles for the detection of 6-MP (Hatamluyi et al. 2018). These diverse approaches reflect ongoing efforts to synthesize robust and adaptable detection materials for 6-MP in various environmental and biological contexts, addressing crucial requirements in environmental surveillance and medical applications. Nevertheless, there exists a research gap concerning the utilization of conjugated materials like aniline, carbazole, acetylene, and thiophene for 6-MP detection. Thiophene and n-vinyl carbazole, notable among these materials, due to their synthetic flexibility and favorable optoelectronic properties, offer distinct advantages. Thiophene and its derivatives facilitate polymer backbone planarization, carbazole its nitrogen-containing and aromatic



heterocyclic derivatives find broad applications across industries, including pharmaceuticals and light-emitting photosensors (Sharma *et al.* 2023; Tajik *et al.* 2021). Consequently, this research embarks on a comprehensive comparative analysis of the sensing ability of 3-MeOTH and its copolymer with n-vinyl carbazole (3-MeOTH-NVK), through studying their electronic properties and adsorption behavior using Density Functional Theory (DFT).

## 2. COMPUTATIONAL DETAILS

The Density Functional Theory was performed to analyze the structural and electronic properties of synthesized polymer matrices. The analysis exploited the Perdew-Burke-Ernzerhof (PBE) functional within the generalized gradient approximation (GGA), applying double-zeta-polarized (DZP) basis sets for optimization. Matrices were optimized until forces on each atom reached below 0.05 eV/Å. A Rydberg cut-off energy of 75 and convergence threshold energy of  $10^{-5}$  eV were chosen, alongside a K-point sampling of  $1\times1\times1$  for accurate calculations of Molecular Energy Spectrum (MES) and Density of States (DOS).

A device simulator with gold (Au) electrodes was used to calculate the transmission spectrum of synthesized polymer matrices. The PBE function with local density approximation (LDA), 75 Hartee cut-off energy and  $3\times3$  K-point are applied for optimization of the device.

## 3. RESULTS AND DISCUSSION

In this study, we analyzed the most stable optimized geometries among various orientations to conduct a comprehensive analysis of the electronic properties of 6-mercaptopurine adsorbed onto both 3-MeOTH and its copolymer 3-MeOTH-NVK.

#### **3.1 Stability Analysis**

To understand the interaction between the 3-MeOTH-NVK and 3-MeOTH with the 6-MP drug, the adsorption energy of both 3-MeOTH+6-MP and 3-MeOTH-NVK+6-MP systems was computed and analyzed using equation 1 (Table 1) (Agrawal et al. 2022). The calculated adsorption energy values for 3-MeOTH and 3-MeOTH-NVK with 6-MP drug are -1.918 eV and -3.173 eV, respectively. The negative value of the adsorption energy indicates an exothermic reaction and weak Van der Waals interaction between both the systems. This interaction is illustrated in Fig.1, where after full relaxation, the distance of 6-MP from 3-MeOTH and 3-MeOTH-NVK systems was found to be 3.33 Å and 2.85 Å, respectively. This distance, defined as the molecule-drug distance, represents the minimum atom-to-atom distance between the molecule and the drug.

The comparison between both the systems reveals that 3-MeOTH-NVK exhibits a significantly higher release of interaction energy, precisely 65.43% more than that of 3-MeOTH. This suggests that the 3-MeOTH-NVK copolymer is more stable than the 3-MeOTH monomer. Furthermore, due to the weak Van der Waals interaction between the drug and the molecule, no significant changes are observed in their geometry. Consequently, the bond length of the molecules remains nearly unchanged after the adsorption of 6-MP.

$$E_{ads} = TE_{(matrix+6-MP)} - (TE_{(matrix)} + TE_{(6-MP)}) \qquad \dots \dots (1)$$

Here,  $E_{ads}$ ,  $TE_{(matrix+6-MP)}$ ,  $TE_{(matrix)}$  and  $TE_{(6-MP)}$  are adsorption energy, energy of optimized molecule-6-MP system and energy of 6-MP drug, respectively.





#### **3.2 Electron Density Analysis**

The charge transfer between the drug and the molecule was computed through Mulliken population analysis using equation 2 (Table 1) (Agrawal *et al.* 2022). Notably, the 3-MeOTH-NVK+6-MP system shows almost double the charge transfer to 6-MP compared to the 3-MeOTH+6-MP system. Specifically, 0.053e is transferred from 3-MeOTH to 6-MP, while 0.106e is transferred from 3-MeOTH-NVK to 6-MP. Fig. 2 presents the electron density and positioning of 6-MP on 3-MeOTH and 3-MeOTH-NVK, where the jade-green color indicates the significant accumulation of electron



densities on nitrogen and oxygen atoms due to their electronegative nature.

Fig. 2: Schematic view of electron densities of 3-MeOTH+6-MP (a) and 3-MeOTH-NVK+6-MP (b) systems at isoscales value of 1.9

$$\Delta \rho = \rho_{(\text{matrix}+6-\text{MP})} - \rho_{(6-\text{MP})} \qquad (2)$$

Here  $\Delta \rho$ ,  $\rho_{(matrix+6-MP)}$ , and  $\rho_{(6-MP)}$  are charge transfer, charge on 6-MP after adsorption on the matrix, and charge on 6-MP before adsorption on the matrix, respectively.

Table 1. Calculated adsorption energies and Mulliken charges of the 3-MeOTH+6MP and 3-MeOTH-NVK+6MP systems

System	$\mathbf{E}_{\mathrm{ads}}$	Q(e) <sup>1</sup>
6-MP on 3-MeOTH-NVK	-3.173	0.106e
6-MP on 3-MeOTH	-1.918	0.053e

## **3.3 Electronic Properties Analysis**

The molecular orbital analysis highlights the significant influence of 6-MP adsorption on the HOMO-

LUMO (E<sub>H</sub>-E<sub>L</sub>) gap for both 3-MeOTH and 3-MeOTH-NVK molecules. Upon 6-MP adsorption, the E<sub>H</sub>-E<sub>L</sub> gap of 3-MeOTH decreases from 4.08 eV to 0.2 eV, while 3-MeOTH-NVK experiences a reduction from 2.8 eV to 0.1 eV. A comparison reveals that the 3-MeOTH-NVK+6-MP system has half the E<sub>H</sub>-E<sub>L</sub> gap of the 3-MeOTH+6-MP system, confirming the respective reduction of the E<sub>H</sub>-E<sub>L</sub> gap. This reduction is evident in the shifting of HOMO and LUMO energy levels towards the Fermi level (Fig. 3). A smaller  $E_H$ - $E_L$  gap indicates a good conductive behaviour of systems. Additionally, the impact of 6-MP on the 3-MeOTH and 3-MeOTH-NVK matrices was analyzed by computing its DOS profile. The right panel of Fig. 3 provides insight into the DOS profiles for 3-MeOTH and 3-MeOTH-NVK before and after the adsorption of 6-MP. The DOS profile for both molecules support the observed changes in the E<sub>H</sub>-E<sub>L</sub> gap, with successive shifts of DOS curve near the Fermi level. Remarkably, the DOS plot of both molecules after the interaction with 6-MP shows remarkable changes in the position, count, and strength (height) of peaks due to alteration in the electronic property, further emphasizing the robustness of the studied systems for drug sensing.



Fig. 3: Schematic view of HOMO-LUMO gap (left panel) and DOS profiles (Right panel) of 3-MeOTH and 3-MeOTH-NVK matrices before and after adsorption of 6-MP drug

The study analyzed conductance and detection range through transmission spectra of 6-MP adsorbed 3-MeOTH and 6-MP adsorbed 3-MeOTH-NVK systems to validate the suitability of the reduced  $E_{H}$ - $E_{L}$  gap for drug sensing. A two-probe simulation system featuring gold electrodes was used in the device (Fig.4) (Xue *et al.* 2008). A detailed analysis of conductance through the transmission spectrum of the studied systems is shown in Fig. 5. The adsorption of 6-MP results in nearly a threefold enhancement in amplitude for the peaks of 3-MeOTH (Fig. 5a). Moreover, a peak in the range 0.2eV to -0.3eV indicates a change in conductivity. The greater amplitude suggests an increase in conductance(Agrawal *et al.* 2021). Fig. 5b compares 6-MP adsorbed 3MeOTH-NVK with 6-MP adsorbed 3-MeOTH and shows peak amplitude enhancement between 0.1eV to -0.2eV. Moreover, two additional peaks appear at -0.6 eV and -0.7 eV with notable amplitudes, affirming the enhanced conductance of the 3-MeOTH-NVK copolymer for the 6-MP drug. This enhanced conductance is substantiated by a reduced  $E_{H}$ - $E_{L}$  gap.



Fig. 4: (a) Two-probe semi-infinite linear and 3×3×2 gold (Au) chain electrodes with (b) 3-MeOTH+6-MP and (c) 3-MeOTH-NVK+6-MP

Further analysis is directed towards examining variations in conductance within the studied systems concerning the spatial distance between the 6-MP drug and the 3-MeOTH and 3-MeOTH-NVK matrices, particularly within the optimized adsorption range. An inverse relationship between conductance and the distance of the 6-MP drug to the matrices within the optimized adsorption range is observed (Fig. 6), highlighting the sensitivity of both matrices in the presence of the 6-MP drug. Utilizing Equation 3 (Agrawal et al. 2021) (Table 2), the sensitivity of these matrices to the 6-MP drug was calculated. The resulting sensitivities for 3-MeOTH+6-MP and 3-MeOTH-NVK+6-MP are 34.24% and 72.04%, respectively, confirming the significant sensitivities exhibited by both matrices. Notably, 3-MeOTH-NVK+6-MP demonstrates a 37.8% higher sensitivity compared to 3-MeOTH+6-MP system.

$$S = G - G_0 / G \qquad \dots \dots \dots (3)$$

Here, G represents the conductance of the 6-MP adsorbed system, and  $G_0$  is the conductance of the matrix.



Fig. 5: Comparative transmission spectrum of (a) 3-MeOTH and 3-MeOTH+6-MP, and (b) 3-MeOTH+6-MP and 3-MeOTH-NVK+6-MP



Fig. 6: Conductance vs distance graph of (a) 3-MeOTH+6-MP and (b) 3-MeOTH-NVK +6-MP systems

System	Conductance (mS)	Sensitivity (%)
3-MeOTH	3.14	-
3-MeOTH+6-MP	4.78	34.24
3-MeOTH-NVK	1.67	-
3-MeOTH-NVK+6-MP	5.97	72.04

Table 2. Conductance and sensitivity of 3-MeOTH, 3-MeOTH-NVK, 3-MeOTH+6-MP, and 3-MeOTH-NVK+6-MP systems

## 4. CONCLUSION

Computational analyses reveal that 3-MeOTH-NVK is a more reliable matrix for detecting the 6-MP drug than 3-MeOTH. This superiority arises from the 6-MP adsorption on 3-MeOTH-NVK being 1.6 times more stable and involving two-fold charge transfer than with 3-MeOTH. This transformation is supported by the MES and DOS profile, along with approximately 55% conductivity enhancement in the 3-MeOTH-NVK+6-MP system compared to the 3-MeOTH+6-MP system, as calculated using transmission spectra. The sensitivity to the 6-MP drug is significantly enhanced to 37.8% through the copolymerization of 3-MeOTH with NVK. The combination of improved conductance and significant sensitivity establishes 3-MeOTH-NVK as a reliable sensor for detecting the 6-MP drug.

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#### DECLARATION OF COMPETING INTEREST

The author declares no conflict of interest in the presentation of this research.

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