

Molecular Modeling, Docking and ADMET of Dimethylthiohydantoin Derivatives for Prostate Cancer Treatment

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Abstract

In silico technique was applied to screen potential of 16 compounds of 5,5-dimethylthiohydantoin derivatives as androgen antagonist. The 3D structure of the protein was obtained from PDB database. Docking analysis of the compounds was performed using hex docking. Molecular modeling analysis exhibits relatively low LUMO-HOMO energy gap of the studied molecules, indicating that it would be kinetically stable. None of the compounds violated Lipinski's parameters, making them potentially promising agents for biological activities. The title compounds exhibited the lowest docking energy of protein-ligand complex. Finally, the results indicate that these compounds are potentially as an androgen antagonist, and expected to be effective in prostate cancer treatment.

Keywords

Androgen Receptor, Prostate Cancer, Molecular Modeling, Molecular Docking, ADMET

1. Introduction

Cancer can be characterized by the arrest of cell differentiation, the inhibition of apoptosis and the accelerated proliferation of cloned cells. The understanding of the mechanisms of cell-death execution and the role that they play in different diseases opens new therapeutic strategies [1]-[4]. Prostate cancer is the most frequently diagnosed malignancy in men in Western countries [5]. The prostate is an androgen-dependent organ; androgen hormones and their executor, the androgen receptor (AR), are central drivers of prostate cancer development and progression [6]-[11]. The androgen receptor (AR), located on Xq11-12, is a 110 kDa nuclear receptor that, upon

activation by androgens, mediates transcription of target genes that modulate growth and differentiation of prostate epithelial cells AR signaling is crucial for the development and maintenance of male reproductive organs, including the prostate gland, as genetic males harboring loss of function AR mutations and mice engineered with AR defects do not develop prostates or prostate cancer [12] [13]. AR plays a major role in the development and maintenance of male sex characteristics by executing the biological functions of androgens. AR also has an important role in prostate cell proliferation and differentiation; and during androgen responsive prostate cancer [14]-[16].

However, the AR is a cytoplasmic protein and is a member of the steroid/thyroid hormone receptor superfamily [17] [18]. Upon androgen binding to its receptor, the AR migrates into the nucleus, binds to specific DNA sequences called androgen response elements and modulates the transcription of target genes [17]. For this reason, the androgen receptor is a key molecular target in the etiology and progression of prostate cancer [19]-[23]. So that, manipulations of AR function of antiandrogenic drugs are the primary mode of treatment of the malignancy [14] [15].

AR antagonists are used as a single agent (monotherapy) or in combination with castration. The latter use, referred to as “combined androgen blockade therapy”, shows significant effects by blocking adrenal androgen signals as well as suppressing the transient testosterone increase induced by GnRH analogs [24]-[29]. AR antagonists used clinically for prostate cancer include steroidal (cyproterone acetate) and nonsteroidal antagonists (flutamide, nilutamide, and bicalutamide), which all function as competitive inhibitors of AR binding to endogenous androgens testosterone and dihydrotestosterone [30].

Although, the antiandrogens exhibit good efficacy in many cases and comprise an important part of effective therapeutics [31]-[34]. But, a considerable problem with these antiandrogens is that recurrence occurs after a short period of response [35]. Hydroxyflutamide and bicalutamide have partial agonist activities at high concentrations *in vitro* [36], which may contribute to recurrence.

The thiohydantoin derivative with a carboxyl terminal side chain showed that pure antagonistic activities *in vitro* and oral AR antagonistic activity *in vivo* [37]. In this work, the molecular modeling, molecular docking and ADMET of 16 compounds of 5,5-dimethylthiohydantoin derivatives, which synthesized by [38] were conducted for the treatment of prostate cancer.

2. Experimental

2.1. Materials

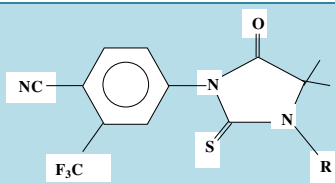
The available data sets of 16 compounds were obtained from the literature [38]. The IUPAC names and structure of 5,5-dimethylthiohydantoin derivatives and are listed in **Table 1** and **Table 2**. On the other hand, the substitute which occurs in all compounds can be seen in these tables.

2.2. Molecular Modeling

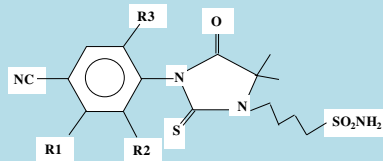
All the calculations are performed by using Gauss view 5.0 molecular visualization program and Gaussian 09 program package on the personal computer [39]. The molecular structure of the title compounds in the ground state (in a vacuum) was optimized using Semi-empirical method AM1. The frontier molecular orbital surfaces are visualized by Gauss View Molecular Visualization program [40]. All these computations were carried out for the ground states of these molecules as a single state. The dipole moment (in Debye) of the molecules and heat of formation (HOF) was directly extracted from the Gaussian09 output file. The HOMO and LUMO values were taken from the output of the Gaussian 09W calculation as molecular orbital coefficients (in a.u.).

2.3. Molecular Descriptors

Computation of molecular descriptors such as partition coefficient, topological surface area and a number of hydrogen bond acceptors & donors, was carried out using Mol inspiration online tool [41]. Using these parameters the compounds were checked for their compliance with the Lipinski rule of five [42]. Drug likeness score was computed using molsoft website [43]. The drug-likeness scores were analyzed by comparing with the earlier reports [44]. Bioactivity score predictions were also done using Mol inspiration on-line tool. Toxicity Risk Assessment, cLogP Prediction, Solubility Prediction, Molecular Weights, Drug-Likeness, Prediction and Overall Drug-Likeness Score computed using Osiris software available on-line.

Table 1. The IUPAC name of compounds from 1 to 10.


Compound No.	R	IUPAC name
1	(CH ₂) ₃ CONH ₂	4-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]butyramide
2	(CH ₂) ₃ NH ₂	4-[3-(3-Aminopropyl)-4,4-dimethyl-5-oxo-2-thioxoimidazolidin-1-yl]-2-trifluoromethylbenzonitrile
3	(CH ₂) ₃ NH CO NH ₂	3-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propylurea
4	(CH ₂) ₃ NHSO ₂ NH ₂	<i>N</i> -(3-{3-[4-cyano-3-(trifluoromethyl) phenyl]-5,5-dimethyl-4-oxo-2-sulfanylideneimidazolidin-1-yl}propyl) aminosulfonamide
5	(CH ₂) ₃ SO ₃ H	3-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl] propane-1-sulfonic acid
6	(CH ₂) ₃ SO ₂ NH ₂	3-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide
7	(CH ₂) ₄ SO ₂ NH ₂	4-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]butane-1-sulfonamide
8	(CH ₂) ₂ SO ₂ NH ₂	2-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl] ethanesulfonamide
9	(CH ₂) ₂ SO ₂ NHMe	3-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl] propane-1-sulfonic acid methylamide
10	(CH ₂) ₂ SO ₂ NHMe ₂	3-[3-(4-Cyano-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl] propane-1-sulfonic acid dimethylamide

Table 2. The IUPAC name of compounds from 11 to 16.


Compound No.	R1, R2 and R3	IUPAC NAME
11	R1 = R2 = R3 = H	3-[3-(4-Cyanophenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide
12	R1 = Me, R2 = R3 = H	3-[3-(4-Cyano-3-methylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide
13	R1 = OMe, R2 = R3 = H	3-[3-(4-Cyano-3-methoxyphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide
14	R1 = Cl, R2 = R3 = H	3-[3-(3-Chloro-4-cyanophenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide
15	R1 = CF ₃ , R2 = Me, R3 = H	3-[3-(4-Cyano-2-methyl-3-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide
16	R1 = CF ₃ , R2 = H, R3 = Me	3-[3-(4-Cyano-2-methyl-5-trifluoromethylphenyl)-5,5-dimethyl-4-oxo-2-thioxoimidazolidin-1-yl]propane-1-sulfonamide

2.4. Molecular Dynamics Simulation

In the present study, Bioinformatics tools are used, biological databases like PDB (Protein Data Bank) and the software's like Hex [45]. The Protein Data Bank (PDB) is the single worldwide archive of structural data of biological macromolecules, established in Brookhaven National Laboratories [46]. It contains structural information of the macromolecules determined by X-ray crystallographic and NMR methods. The structure of the androgen receptor (**Figure 1**), which is an essential target for the hydroxyflutamide drug was retrieved from PDB (2AX6). On the other hand, most favored regions of the target structure were evaluated through Ramchandran plot analysis via PROCHECK [47] from SAVES Server (<http://services.mbi.ucla.edu/SAVES/>) to investigate the quality of the target structure.

2.4.1. Activity Prediction

The activity prediction of all molecules has been achieved by PASS server and compared with hydroxyflutamide. PASS Online predicts over 3500 kinds of biological activity, including pharmacological effects, mechanisms of action, toxic and adverse effects, interaction with metabolic enzymes and transporters, influence on gene expression, etc. Prediction is based on the analysis of structure activity-relationships for more than 250,000 biologically active substances including drugs, drug-candidates, leads and toxic compounds. The structural formula only is necessary to obtain the predicted biological activity profile for any compound.

2.4.2. HEX Docking

Hex is an interactive molecular graphics program for calculating and displaying feasible docking modes of pairs of protein and DNA molecules. Hex can also calculate protein-ligand docking; assuming the ligand is rigid, and it can superpose pairs of molecules using only knowledge of their 3D shapes [48]. It uses spherical polar fourier (SPF) correlations to accelerate the calculations and its one of the few docking programs, which have been built in graphics to view the result [49].

The docking analysis of hydroxyflutamide with androgen receptor was carried by HEX docking software. Docking allows the scientist to virtually screen a database of compounds and predict the strongest binders based on various scoring functions. It explores ways in which two molecules, such as drugs and an enzyme; Human estrogen receptor fit together and docks to each other well, like pieces of a three-dimensional jigsaw puzzle. The molecules binding to a receptor, inhibit its function, and thus act as a drug.

The parameters used in the docking process were

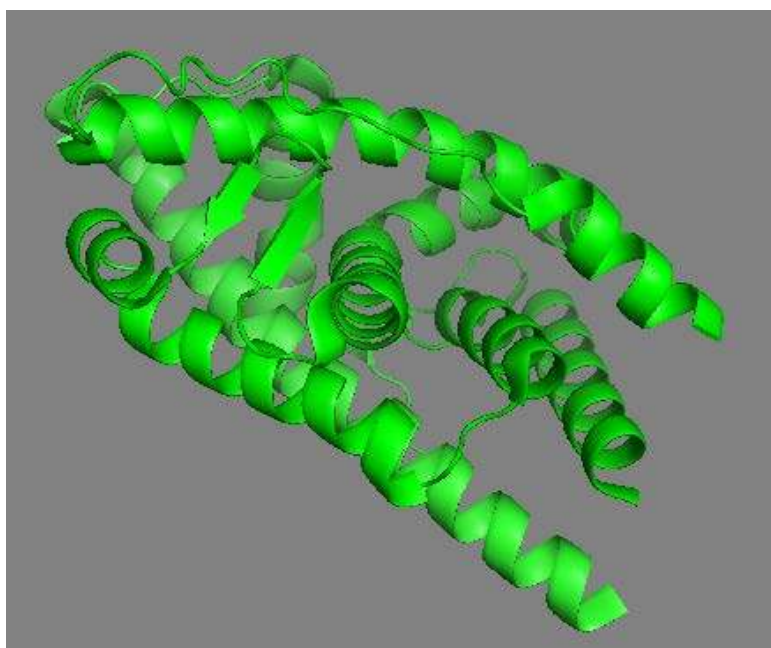


Figure 1. The structure of androgen receptor (2AX6).

- Correlation type—Shape only
- FFT Mode—3D fast lite
- Grid Dimension—0.6
- Receptor range—180
- Ligand Range—180
- Twist range—360
- Distance Range—40

All water molecules and ligands were removed from the proteins for docking studies.

The hydroxyflutamide and 5,5-dimethylthiohydantoin derivatives were docked with the receptor using the above parameters. Visualization of the docked pose had been done using a PyMol molecular graphics program [50].

2.4.3. LigPlot+ v.1.4.5

Two-dimensional representations of the best docking pose for selected 5,5-dimethylthiohydantoin derivatives inside target enzyme were generated using LigPlot+ [51]. It is a computer program which generates the schematic 2-D image of the docked protein-ligand complexes. The 3-D structure of the docked complex is input as PDB file and the software produces their interacting residues and bonds. In the present study, LigPlot+ was used to identify interacting residues as well as the interacting bonds between 2AX6 and the docked inhibitors.

2.4.4. Computed Atlas of Surface Topography of Proteins (CASTp)

Binding sites and active sites of proteins and DNAs are often associated with structural pockets and cavities. CASTp server uses the weighted Delaunay triangulation and the alpha complex for shape measurements. It provides identification and measurements of surface accessible pockets as well as interior inaccessible cavities, for proteins and other molecules. It measures analytically the area and volume of each pocket and cavity, both in the solvent accessible surface (SA, Richards' surface) and molecular surface (MS, Connolly's surface). It also measures the number of the mouth openings, area of the openings, and circumference of mouth, lips, in both SA and MS surfaces for each pocket [52].

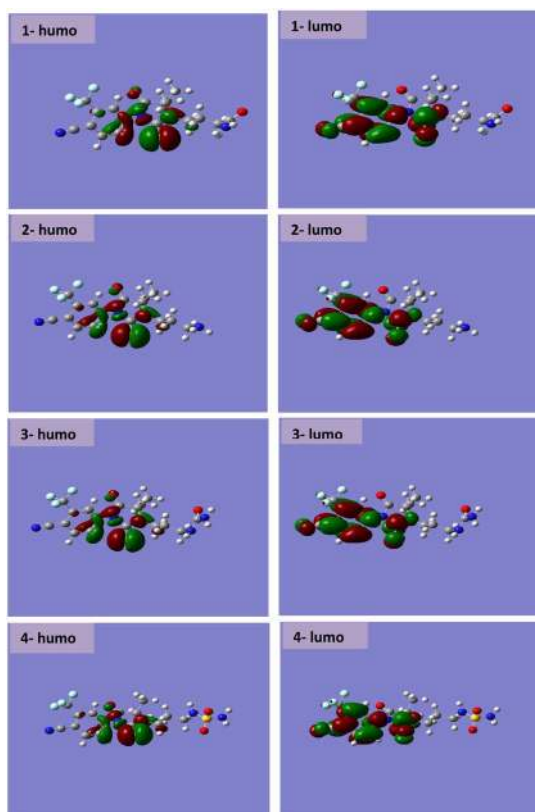
2.4.5. Prediction of ADMET Properties

The ADMET (Absorption, Distribution, Metabolism, Excretion and Toxicity) properties of the target compounds were calculated using some web-based applications. BBB (Blood-Brain Barrier) penetration, HIA (Human Intestinal Absorption), Caco-2 cell permeability and Ames test were calculated using admetSAR [53].

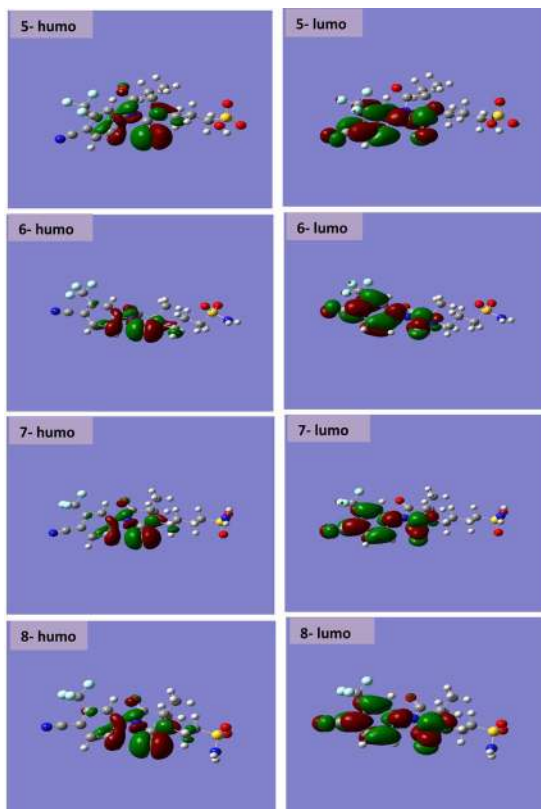
3. Results and Discussions

3.1. Molecular Modeling

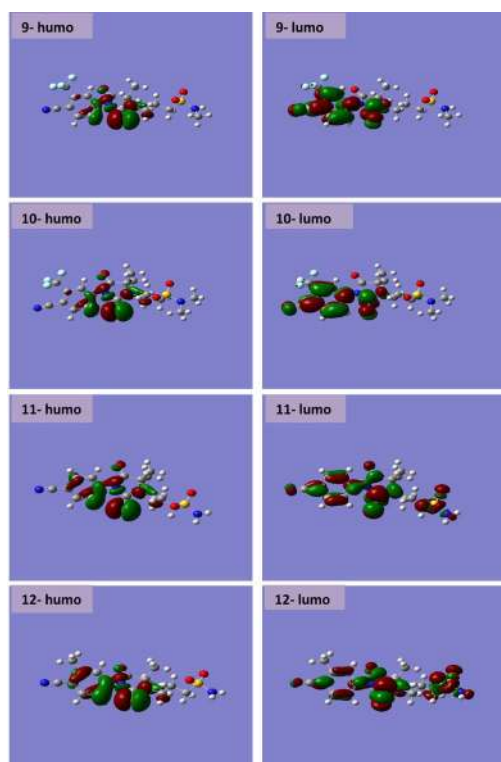
The frontier molecular orbitals for all compounds have been presented in **Figures 2(a)-(d)**. The molecular orbital shows that, the location of possible sites responsible for electron transfer between molecules and its biological target. Thus, we can discover that how molecules react and where is the active sites in reaction. For the molecules **1-10**, the HOMO are delocalized on thiohydantoin system and part of the benzene ring, while in the LUMO, the electron density predominantly are located on the benzene ring, C≡N group, thiohydantoin system except N atom which linked to benzene ring and C=O group. On the other hand, in the case of the molecules **11** and **12**, the HOMO are delocalized on thiohydantoin system and part of the benzene ring, while in the LUMO, the electron density predominantly are located on the benzene ring, C≡N group, thiohydantoin system and CH₂-SO₂-NH₂ system. However, in the case of the molecule **13**, the HOMO are delocalized on thiohydantoin system, the benzene ring and O atom linked to phenyl ring, while in the LUMO, the electron density predominantly are located on the benzene ring, C≡N group, thiohydantoin system and CH₂-SO₂-NH₂ system. In the case of molecule **14**, the HOMO are delocalized on thiohydantoin system, Cl atom and part of the benzene ring, while in the LUMO, the electron density predominantly are located on the benzene ring, C≡N group, thiohydantoin system, Cl atom and SO₂ system. But, the HOMO of molecule **15** are delocalized on thiohydantoin system, while in the LUMO, the electron density predominantly located are on the benzene ring, C≡N group and thiohydantoin system. However, the HOMO of molecule **16** are delocalized on thiohydantoin system, while in the LUMO, the electron density predominantly are located on the benzene ring, C≡N group and part of



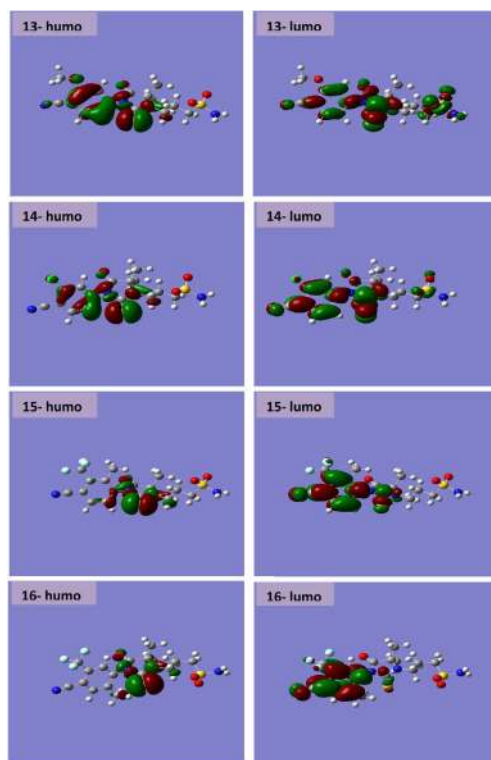
(a)



(b)



(c)



(d)

Figure 2. (a) The 3D graph of HOMO and LUMO of 1- 4 molecules; (b) The 3D graph of HOMO and LUMO of 5 - 8 molecules; (c) The 3D graph of HOMO and LUMO of 9 - 12 molecules; (d) The 3D graph of HOMO and LUMO of 13 - 16 molecules

thiohydantoin system.

The energies of the frontier orbital (EHOMO and ELUMO) obtained with the AM1 semiempirical method are presented in **Table 3**. From these results, we can consider the compounds with larger values of EHOMO as being more electron donor and the compounds with smaller values of ELUMO as being more electron acceptor. From **Table 3**, it can be seen that, the EHOMO and ELUMO calculated values present significant differences: the values of EHOMO vary from -0.32712 au to -0.34521 au and the ELUMO values vary from -0.04819 au to -0.02938 au. HOMO-LUMO refers to the energetic difference of HOMO and LUMO. Such an energetic gap can be used as an indicator of chemical reactivity, where the low energy gap correlates to high chemical reactivity and vice versa. Considering the chemical hardness, large HOMO-LUMO gap means a hard molecule and small HOMO-LUMO gap means a soft molecule. One can also relate the stability of the molecule to hardness, which means that the molecule with least HOMO-LUMO gap means it, is more reactive. Relatively low LUMO-HOMO energy gap of the studied molecules indicates that, it would be kinetically stable. According to AM1 calculation, the heat of formation (HF) energy of the studied molecules has a negative sign (**Table 3**), and they are exothermic. However, the exothermic nature of HF makes the studied molecules thermodynamically stable [54].

In the framework of density functional theory (DFT) global descriptors of chemical reactivity corresponds to global responses of systems to global perturbations (for instance, changes in the number of electrons N), whereas the external potential remains constant. Among such types of indexes, chemical potential (μ), chemical hardness (η) and softness (s) can be used as complementary tools in the description of thermodynamic aspects of chemical reactivity.

$$\mu = \frac{1}{2} \left(\frac{\partial E}{\partial N} \right)_{v(r)} \quad (1)$$

$$\eta = \frac{1}{2} \left(\frac{\partial^2 E}{\partial N^2} \right)_{v(r)} \quad (2)$$

From equation (1), the first order partial derivatives of total energy (E) with respect to the number of electrons

Table 3. The frontier orbital's energies and heat of formation.

Compound No.	Lumo	Homo	Lumo-Homo	Hardness	Hf kcal/mol
1	-0.04604	-0.34227	0.29623	0.148115	-134.037
2	-0.04247	-0.33717	0.29470	0.147350	-90.6825
3	-0.04198	-0.33679	0.29481	0.147405	-125.751
4	-0.04377	-0.33874	0.29497	0.147485	-163.745
5	-0.04559	-0.34209	0.29650	0.148250	-208.489
6	-0.04221	-0.33773	0.29552	0.147760	-159.117
7	-0.04333	-0.33785	0.29452	0.147260	-166.572
8	-0.04819	-0.34521	0.29702	0.148510	-153.122
9	-0.04217	-0.33766	0.29549	0.147745	-155.617
10	-0.04205	-0.33745	0.29540	0.147700	-150.352
11	-0.03027	-0.32958	0.29931	0.149655	-8.83272
12	-0.02941	-0.32823	0.29882	0.149410	-15.6560
13	-0.02938	-0.32712	0.29774	0.148870	-42.8924
14	-0.03430	-0.33500	0.30070	0.150350	-13.4327
15	-0.04058	-0.33698	0.29640	0.148200	-160.789
16	-0.04315	-0.34082	0.29767	0.148835	-167.172

(N) at constant external potential, $V(\mathbf{r})$, define chemical potential (μ). On the other hand according to equation (2) the second partial derivatives of total energy (E) with respect to the number of electrons (N) at constant external potential, $V(\mathbf{r})$, define the global hardness (η) of the system [55].

Operational schemes for the calculation of chemical hardness are based on a finite difference method and thus,

$$\mu \approx -\frac{1}{2}(I \cdot P + E \cdot A) \quad (3)$$

$$\eta \approx \frac{1}{2}(I \cdot P + E \cdot A) \quad (4)$$

where, $I \cdot P$ = Ionization Potential and $E \cdot A$ = Electron Affinity. Using the Koopmans' theorem in terms of the energies of highest occupied molecular orbital (EHOMO) and lowest unoccupied molecular orbital (ELUMO), according equation (5) & (6) as follow

$$I \cdot P \approx -EHOMO \quad (5)$$

$$E \cdot A \approx -ELUMO \quad (6)$$

So that, the equation 3 & 4 can be expressed as follows.

$$\mu \approx \frac{1}{2}(EHOMO + ELUMO) \quad (7)$$

$$\eta \approx \frac{1}{2}(ELUMO - EHOMO) \quad (8)$$

Electron affinity refers to the capability of the ligand to accept precisely one electron from a donor. However, in many kinds of bonding viz. covalent hydrogen bonding, partial charge transfer takes places. Softness (S) is a property of the compound that measures the extent of chemical reactivity. It is the reciprocal of hardness.

$$S = \frac{1}{2\eta} \quad (9)$$

Recently Parr *et al.* [56] have defined a new descriptor quantity of the global electrophilic power as an electrophilicity index (ω) of the compound, which defines a quantitative classification of the global electrophilic nature of a compound. Parr *et al.* have proposed electrophilicity index (ω) as a measure of energy lowering due to maximal electron flow between donor and acceptor. They defined electrophilicity index (ω) as follows.

$$\omega = \frac{\mu^2}{2\eta} \quad (10)$$

This index measures the stabilization in energy when the system acquired an additional electronic charge from the environment. Electrophilicity encompasses both the abilities of an electrophile to acquire an additional electronic charge and the resistance of the system to exchange electronic charge with the environment. It contains information about both electron transfer (chemical potential) and stability (hardness) and is a better descriptor of global chemical reactivity. On the other hand, electron affinity (EA) ionization potential (IP), molecular softness, electrophilic index and electronegativity (X) were derived from these results of HOMO and LUMO as per **Table 4**. It is seen that the chemical potential of the title compound is negative, and it means that the compound is stable. They do not decompose spontaneously into the elements they are made up of. The hardness signifies the resistance towards the deformation of the electron cloud of chemical systems under small perturbation encountered during the chemical process.

3.2. Molinspiration Calculations

A computational study for prediction of ADME properties of all molecules is presented in **Table 5**. The number of rotatable bonds and Lipinski's rule of five were also calculated [57]. The rule states that most molecules with

Table 4. Electron affinity, ionization potential, molecular softness, electrophilic index and electronegativity of the title compounds.

Compound No.	Electron affinity (EA)	Ionization potential(IP)	Molecular softness	Electrophilic index	Electronegativity X
1	0.04604	0.34227	6.751510651	0.127253026	-0.1941550
2	0.04247	0.33717	6.786562606	0.122265465	-0.1898200
3	0.04198	0.33679	6.784030392	0.121660318	-0.1893850
4	0.04377	0.33874	6.780350544	0.124007442	-0.1912550
5	0.04559	0.34209	6.745362563	0.126724943	-0.1938400
6	0.04221	0.33773	6.767731456	0.122118980	-0.1899700
7	0.04333	0.33785	6.790710308	0.123334742	-0.1905900
8	0.04819	0.34521	6.733553296	0.130263585	-0.1967000
9	0.04217	0.33766	6.768418559	0.122060669	-0.1899150
10	0.04205	0.33745	6.770480704	0.121885790	-0.1897500
11	0.03027	0.32958	6.682035348	0.108158784	-0.1799250
12	0.02941	0.32823	6.692992437	0.107009546	-0.1788200
13	0.02938	0.32712	6.717270101	0.106714121	-0.1782500
14	0.03430	0.33500	6.651147323	0.113387504	-0.1846500
15	0.04058	0.33698	6.747638327	0.120235791	-0.1887800
16	0.043735	0.33855	6.783915337	0.123926718	-0.1911425

Table 5. Prediction of molecular properties descriptors of the title compounds.

Compound no.	Logp	TPSA A ²	(% ABS)	MW	HBA	HBD	N vio	Nrotb	Volume A ³
	≤5	-	-	<500	<10	<5	≤1	-	-
1	2.319	90.4330	77.80062	398.410	6	2	0	6	324.075
2	2.121	73.3620	83.69011	370.400	5	2	0	5	305.091
3	2.378	102.460	73.65130	413.425	7	3	0	6	336.477
4	1.862	119.531	67.76181	449.480	8	3	0	7	348.925
5	0.400	101.709	73.91040	435.449	7	1	0	6	333.252
6	2.102	107.504	71.91112	434.465	7	2	0	6	336.523
7	0.671	101.709	73.91040	449.476	7	1	0	7	350.054
8	1.832	107.504	71.91112	420.438	7	2	0	5	319.721
9	2.477	93.5080	76.73974	448.492	7	1	0	7	354.197
10	2.722	84.7190	79.77195	462.519	7	0	0	7	371.140
11	1.279	107.504	71.91112	366.468	7	2	0	5	305.225
12	1.656	107.504	71.91112	380.495	7	2	0	5	321.786
13	1.264	116.738	68.72539	396.494	8	2	0	6	330.771
14	1.885	107.504	71.91112	400.913	7	2	0	5	318.761
15	2.479	107.504	71.91112	448.492	7	2	0	6	353.083
16	2.479	107.504	71.91112	448.492	7	2	0	6	353.083
HOF	2.076	95.1500	76.17325	292.213	6	2	0	4	227.700

LogP, logarithm of compound partition coefficient between n-octanol and water; TPSA, topological polar surface area; % ABS, percentage of absorption; MW, molecular weight HBA, number of hydrogen bond acceptors; HBD, number of hydrogen bond donors; Nrotb, number of rotatable bonds; Nvio, Number of violations.

good membrane permeability have $\log P \leq 5$, molecular weight ≤ 500 , a number of hydrogen bond acceptors ≤ 10 , and a number of hydrogen bond donors ≤ 5 . This rule is widely used as a filter for drug-like properties. Furthermore, none of the compounds violated Lipinski's parameters, making them potentially promising agents for biological activities. On the other hand, the number of rotatable bonds is important for conformational changes of molecules under study and ultimately for the binding with receptors or channels. It is revealed that for passing oral bioavailability criteria, a number of rotatable bonds should be ≤ 10 [58]. The compounds in this series, in general, possess a high number of rotatable bonds (5 - 7) and therefore, exhibit conformational flexibility. Topological Polar Surface Area (TPSA) is calculated based on the methodology published by Ertl *et al.*, as a sum of fragment-based contributions [59] [60] in which O- and N- centered polar fragments are to be considered and calculated by surface areas that are occupied by oxygen and nitrogen atoms, and by hydrogen atoms attached to them. TPSA has been shown to be a very good descriptor characterizing drug absorption, including intestinal absorption, bioavailability, Caco-2 permeability and blood-brain barrier penetration. Thus, the TPSA is closely related to the hydrogen bonding potential of a compound [61]. It was found that passively absorbed molecules with a TPSA more than 140 Å are thought to have low oral availability [62]. In respect of TPSA, all the compounds were found within the limit, *i.e.* 140 Å, which implies that molecules are fulfilling the optimal requirement for drug absorption. TPSA was used to calculate the percentage of absorption (% ABS) following the equation:

$\% \text{ ABS} = 109 - (0.345 \times \text{TPSA})$ as reported [63]. From all these parameters, it can be observed that all the title compounds exhibited a moderate % ABS ranging from 67.76% to 83.69%.

3.3. Calculation of Bioactivity Scores

The activity of all test compounds and the standard drug (Flutamide) were rigorously analyzed under four criteria of known successful drug activity in areas of GPCR ligand, ion channel modulator, kinase inhibitor and nuclear receptor ligand. For average organic molecules, the probability is that, if the bioactivity score is more than 0 then it is active if -0.5 to 0 then moderately active [64]. As per **Table 6**, it is readily seen that all the compounds

Table 6. Prediction of bioactivity by Molinspiration of title compounds.

Comp.	GPCR	ICM	KI	NRL	PI	EI
1	0.00	-0.12	-0.11	0.51	0.02	-0.04
2	0.09	0.01	-0.04	0.50	0.06	0.02
3	0.00	-0.16	-0.12	0.29	-0.01	-0.05
4	0.09	-0.26	-0.11	0.51	0.17	0.21
5	-0.01	-0.11	-0.26	0.49	0.07	0.07
6	0.03	-0.12	-0.25	0.57	0.13	0.02
7	-0.01	-0.13	-0.27	0.50	0.08	0.11
8	-0.05	-0.23	-0.20	0.49	0.09	0.02
9	0.03	-0.17	-0.24	0.49	0.00	-0.12
10	0.03	-0.15	-0.29	0.40	0.06	-0.08
11	0.02	-0.22	-0.30	0.46	0.13	0.06
12	-0.05	-0.31	-0.38	0.46	0.07	-0.03
13	-0.01	-0.36	-0.31	0.39	0.01	0.02
14	-0.05	-0.31	-0.34	0.48	0.04	0.04
15	0.05	-0.23	-0.35	0.46	0.04	-0.06
16	0.01	-0.16	-0.27	0.54	0.08	-0.01
HOF	-0.36	-0.04	-0.25	0.10	-0.28	-0.30

GPCR = GPCR ligand, ICM = Ion channel modulator, KI = Kinase inhibitor, NRL = Nuclear receptor ligand, PI = Protease inhibitor and EI = Enzyme inhibitor. HOF = Hydroxyflutamide.

are expected to have near similar activity to standard drugs used based upon these four rigorous criteria (GPCR ligand, ion channel modulator, kinase inhibitor, and nuclear receptor ligand).

3.4. Osiris Calculations

The toxicity risks (mutagenicity, tumorigenicity, irritation, reproduction) were calculated by the methodology developed by Osiris. The toxicity risks predictor locates fragments within a molecule, which indicates a potential toxicity risk. Toxicity risk alerts are an indication that the drawn structure may be harmful concerning the risk category specified. From the data evaluated in [Table 7](#), it is obvious that all molecules are supposed to be non-mutagenic, non-irritating with no-reproductive effects when run through the mutagenicity assessment system in comparison with the standard drug.

3.5. The Aqueous Solubility

The aqueous solubility (S) of a compound significantly affects its absorption and distribution characteristics. Typically, a low solubility goes along with a bad absorption, and therefore the general aim is to avoid poorly soluble compounds. Our estimated logS value is a unit stripped logarithm (base 10) of a compound's solubility measured in mol/liter. There are more than 80% of the drugs on the market have an (estimated) logS value greater than -4. In the case of title compounds, values of logS are around -4. Further, [Table 8](#) shows the drug-likeness (DL) of title compounds which are in the comparable zone with that of standard drug used for comparison. We have calculated overall drug score (DS) for the compounds 1 - 16 and compared with that of standard Hydroxyflutamide. The drug score combines drug-likeness, m_i LogP, logS, molecular weight and toxicity risks in one handy value then may be used to judge the compound's overall potential to qualify for a drug. This value is calculated by multiplying contributions of the individual properties according to the following equation.

$$DS = \Pi(1/2 + 1/2 S_i) \Pi t_i$$

Table 7. Prediction of toxicity risks by Osiris of title compounds.

Compound no.	MUT	TUM	IRRIT	RE
1	■	■	■	■
2	■	■	■	■
3	■	■	■	■
4	■	■	■	■
5	■	■	■	■
6	■	■	■	■
7	■	■	■	■
8	■	■	■	■
9	■	■	■	■
10	■	■	■	■
11	■	■	■	■
12	■	■	■	■
13	■	■	■	■
14	■	■	■	■
15	■	■	■	■
16	■	■	■	■
HOF	■	■	■	■

MUT: mutagenic; TUM: tumorigenic; IRRIT: irritant; RE: reproductive effective.

Table 8. Prediction of Bioavailability and Drug Score by Osiris of title compounds.

Comp.	S	DL	DS
1	-4.39	-8.360	0.35
2	-4.14	-9.530	0.38
3	-4.60	-6.810	0.34
4	-4.60	-6.540	0.32
5	-3.80	-16.98	0.37
6	-5.09	-9.990	0.30
7	-4.07	-16.39	0.35
8	-4.82	-10.00	0.33
9	-4.74	-10.02	0.31
10	-4.33	-9.790	0.32
11	-4.31	-4.530	0.38
12	-4.65	-4.950	0.21
13	-4.33	-5.150	0.37
14	-5.05	-2.970	0.27
15	-5.43	-9.330	0.28
16	-5.43	-9.330	0.28
HOF	-3.18	-13.19	0.35

S: Solubility, DL: Drug likeness, DS: Drug Score.

where; $S = 1/1 + e^{ap+b}$

DS is the drug score. S_i is the contributions calculated directly from m_i LogP; logS, molecular weight and drug likeness (π) via the second equation, which describes a spline curve. Parameters a and b are (1, -5), (1, 5), (0.012, -6) and (1, 0) for m_i LogP, logS, molecular weight and drug-likeness, respectively. The t_i is the contributions taken from the four toxicity risk types and the values are 1.0, 0.8 and 0.6 for no risk, medium risk and high risk, respectively. From this work, all compounds showed moderate to good drug score as compared with the standard drug used.

3.6. Ramachandran Plot Analysis

The Ramachandran plot analysis for androgen receptor PDB (2AX6) revealed that, the 2AX6 is an excellent choice which is in good quality to use as a target for docking studies with the title compounds. The percentage of residues in the most favored-regions was 94.1% in 2AX6 (above the cut off 75%), and the percentage of residues in additional allowed regions was 5.9% in 2AX6. This result suggests that the target enzyme's structure was of good quality.

3.7. Prediction of Activity

PASS (Prediction of Activity Spectra) [64] is an online tool which predicts almost 900 types of activities based on the structure of a compound. The activity prediction of all title compounds has been achieved by PASS server and compared with hydroxyflutamide. As it can be seen in **Table 9**, the designed ligands as well hydroxyflutamide when examine in the server as a smile format shows the significant properties like "Androgen antagonist and prostate cancer treatment" which predicts that the designed ligands will probably be effective to treat prostate cancer as that of hydroxyflutamide. Based on this we decided to evaluate the in silico anti-cancer activity of the title compounds against the human androgen receptor enzyme (2AX6).

Table 9. Activity prediction of 5,5-dimethylthiohydantoin derivatives and hydroxyflutamide with PASS server.

COMP.	Androgen antagonist		Prostate cancer treatment	
	pa	pi	Pa	pi
1	0.730	0.003	0.738	0.004
2	0.654	0.004	0.765	0.004
3	0.558	0.004	0.740	0.004
4	0.802	0.003	0.711	0.005
5	0.500	0.005	0.623	0.005
6	0.995	0.002	0.706	0.004
7	0.991	0.002	0.697	0.004
8	0.995	0.002	0.714	0.004
9	0.957	0.002	0.638	0.005
10	0.987	0.002	0.655	0.005
11	0.787	0.003	0.693	0.007
12	0.892	0.002	0.652	0.005
13	0.731	0.003	0.584	0.005
14	0.865	0.003	0.627	0.005
15	0.981	0.002	0.669	0.004
16	0.978	0.002	0.684	0.004
HOF	0.403	0.006	0.368	0.027

Pa = Probability of Active, Pi = Probability of Inactive, Pa > Pi confirms significant activity

3.8. Hex Docking

Docking results between androgen receptor 2AX6 and 5,5-dimethylthiohydantoin derivatives are reported in **Table 10**. The molecular docking study of the title compounds with human androgen receptor shows that, all the title compounds are showing better docking score than that of hydroxyflutamide which predicts that the title compounds have the better binding affinity to the androgen receptor than hydroxyflutamide. Among all the designed compounds, the compound 8 shows the better docking score (**Figure 3**). This prediction leads us to believe that, the title compounds will possibly be suitable for treatment of prostate cancer.

3.9. Active Sites Identification

In order to find the active sites, the Castp Server is used. The PDB file is used as an input and the results are obtained from this tool which explains the total number of active sites present in the Query PDF file along with information on their amino acid sequence. In addition to that, it also gives their area and volume of the pockets. Pockets are empty concavities on a protein surface into which solvent (probe sphere 1.4 Å) can gain access, *i.e.*, these concavities have mouth openings connecting their interior with the outside bulk solution. Currently, shallow depressions are excluded from the calculation. In 2AX6 there are 33 pockets presented in **Figure 4**. On the other hand, the details of the pocket area, volume, positions and amino acids are given in **Table 11**. The results obtained from the Castp server revealed that, the 32nd pocket has a maximum area of 379.7 mm² and Volumes of 473.2 mm³. Further the amino acids like ILE, MET, PHE, LEU, ALA, PHE, LEU, ALA, PHE, LEU, MET, MET, PHE, ARG, MET, VAL, MET, MET, TRP, GLN, GLY, LEU, ASN, LEU, LEU are presented in this pocket.

3.10. Lig Plot⁺

The interactions of each compound with the functional residues of 2AX6 demonstrated that all the ligands interact

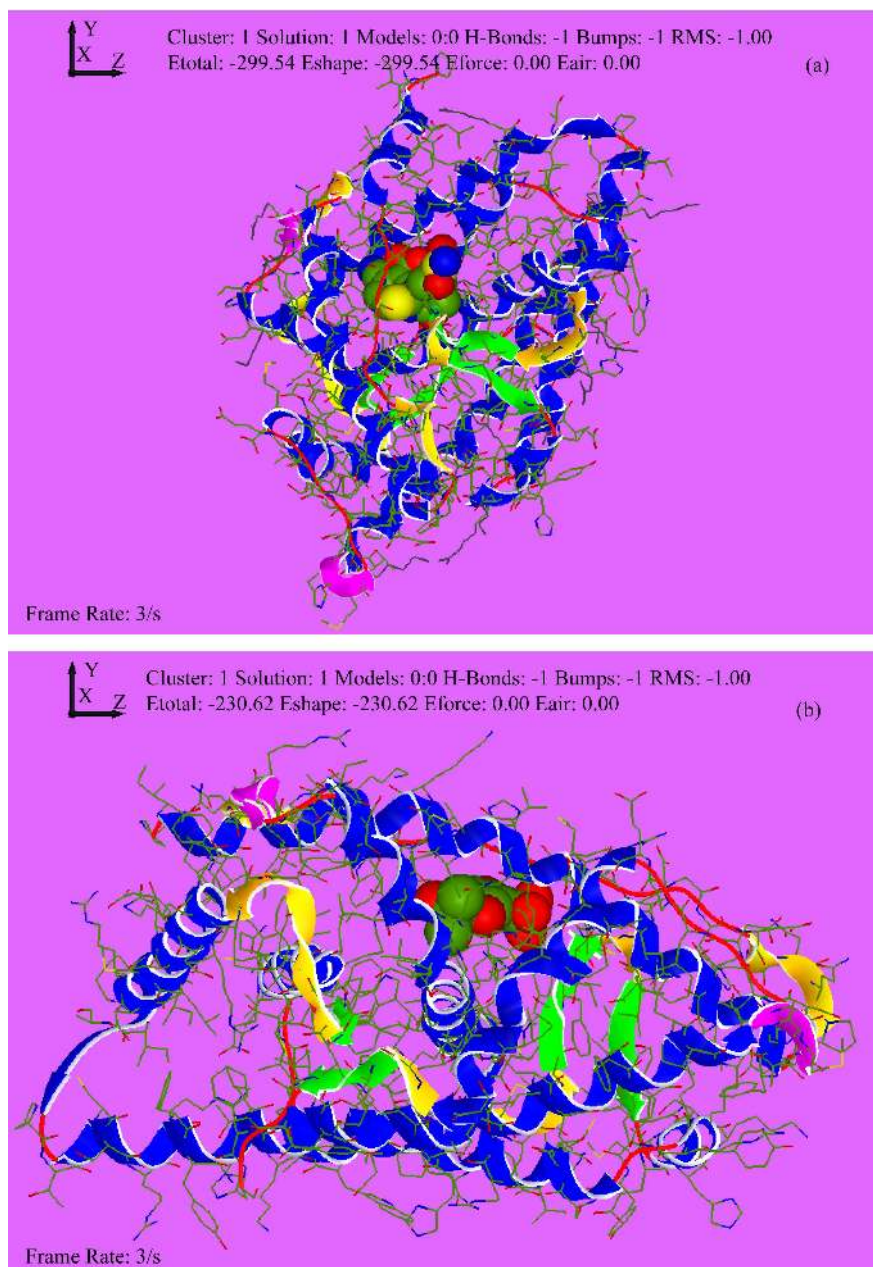
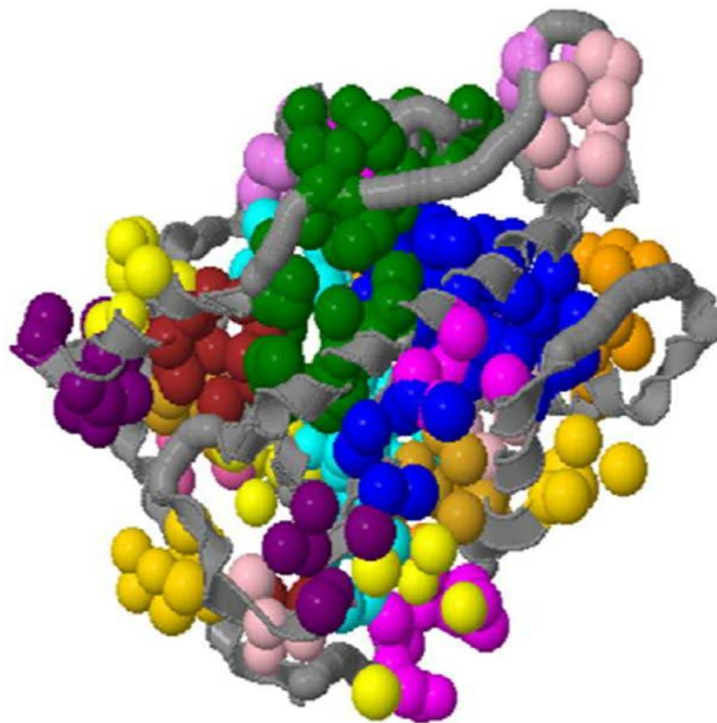


Figure 3. Interaction and binding energy of compound 8 (a) and hydroxyflutamide (b) with androgen receptor (2AX6) respectively.

with most of the residues in the binding pocket as shown in **Figure 5(a)-(d)**. Compound **1** had no hydrogen bond interaction with the protein, but got two external bonds with His789. Also, compound **1** got hydrophobic interaction with Leu790, Val785, Ile869, Arg786, Ser865, Lys861, Leu862 and Glu793. Also, compound **2** had no hydrogen bond interaction with the protein, but got the hydrophobic interaction with Val713, Leu712, Met894, Glu893, Glu897, Gln738, Met734 and Val716. Compound **3** was found to show two hydrogen bonds interaction with active site amino acid residues Gln902 and Met734 at a distance of 2.43 and 2.9 respectively, and the hydrophobic interaction with Lys720, Gln733, Val730, Lys822, Asp732, Ala735, Asp731 and Gln738. Compound **4** was found to show three hydrogen bonds interaction with Ala735, Gln902 and Lys822 a distance of 2.65, 3.02 and 2.84 respectively, and the hydrophobic interaction with Val911, Tyr739, Lys910, Pro817, Lys905, Lue821, Gly820, Asp732, Asp731 and Gln738. On the other hand, compound **5** had no hydrogen bond interaction with

Table 10. Docking Results of 2AX6 enzyme with 5,5-dimethylthiohydantoin Derivatives.

Compound no.	E-value
1	-283.40
2	-265.67
3	-271.19
4	-275.11
5	-264.86
6	-267.75
7	-263.37
8	-299.54
9	-277.55
10	-276.51
11	-262.84
12	-258.33
13	-264.64
14	-265.53
15	-263.89
16	-281.81
HOF	-230.62

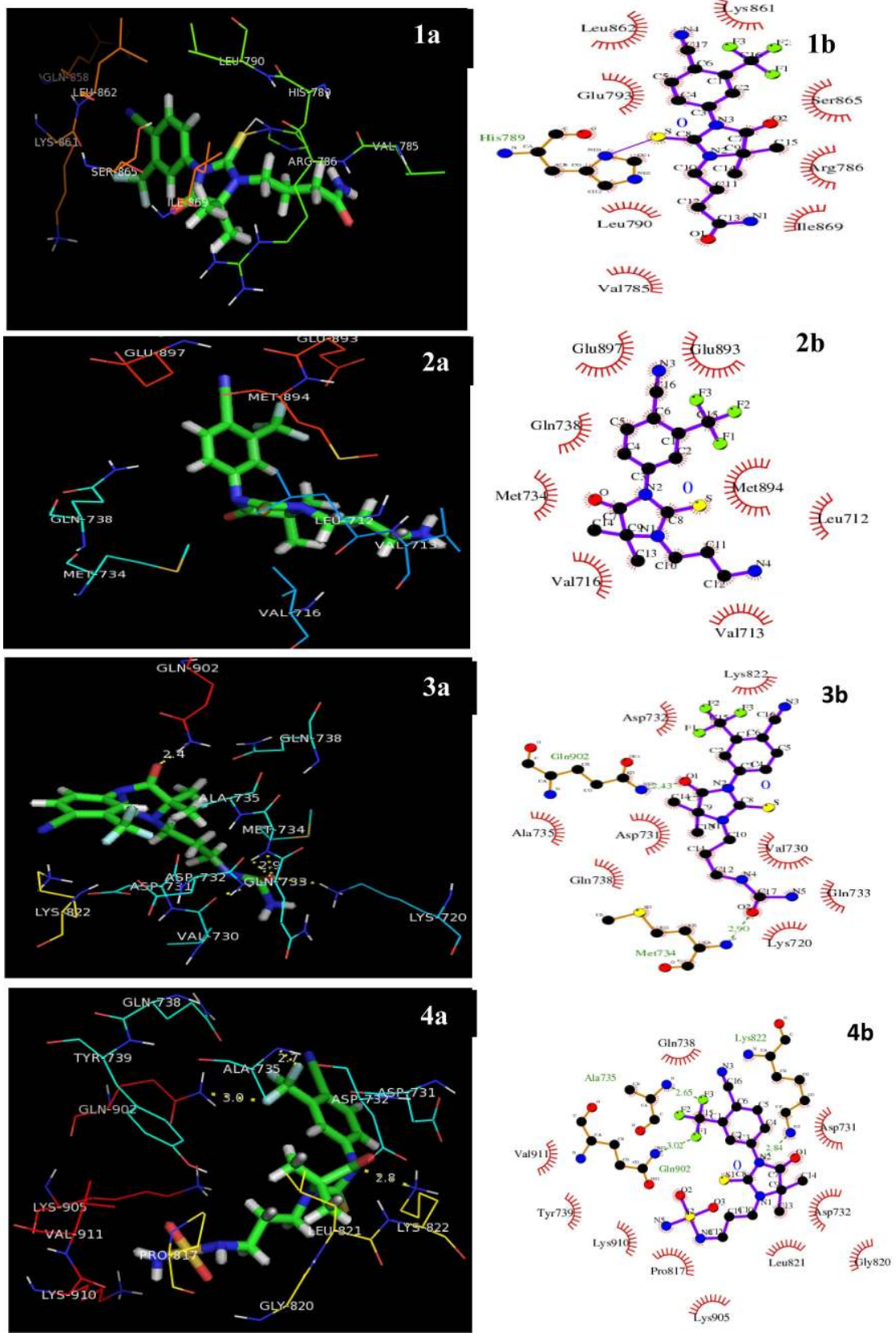


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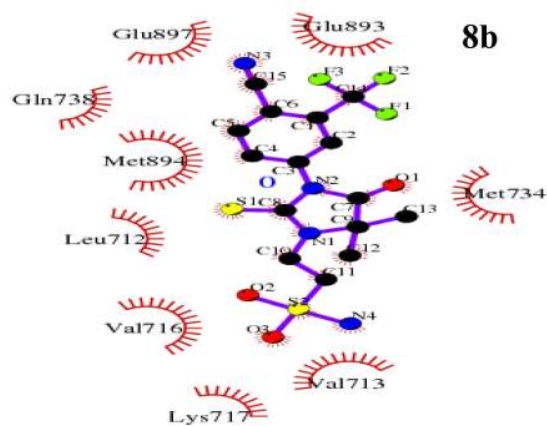
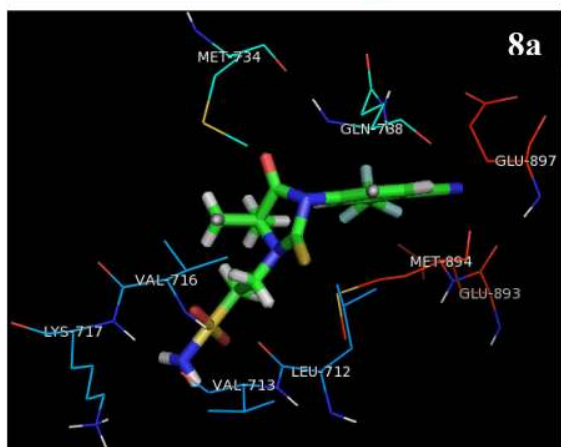
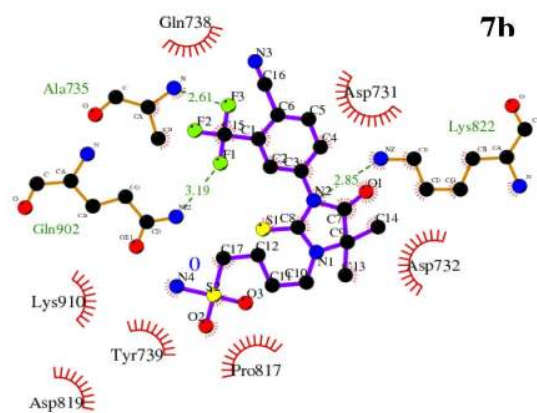
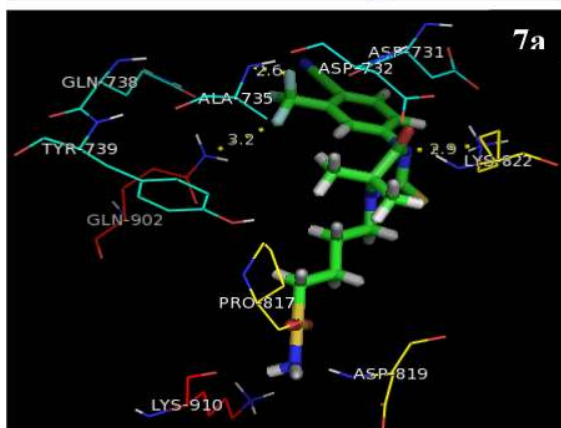
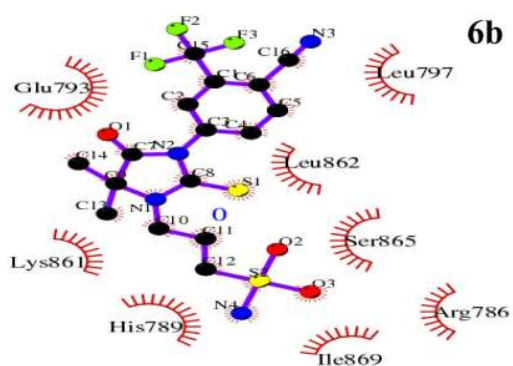
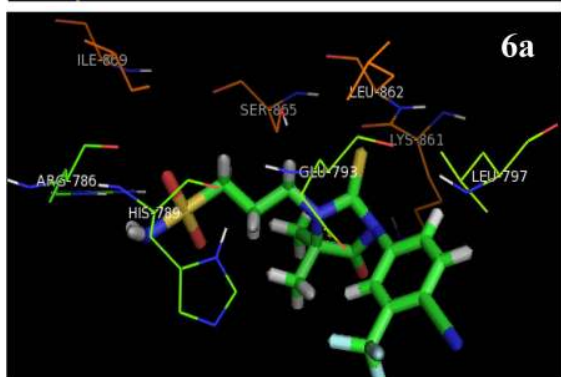
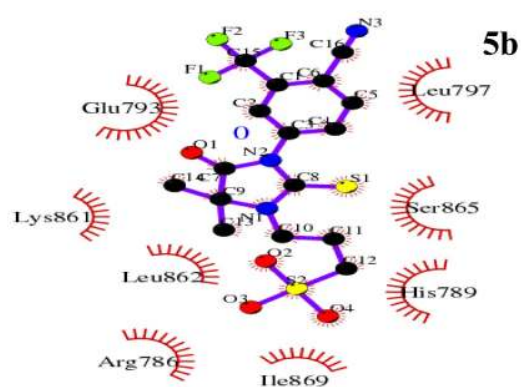
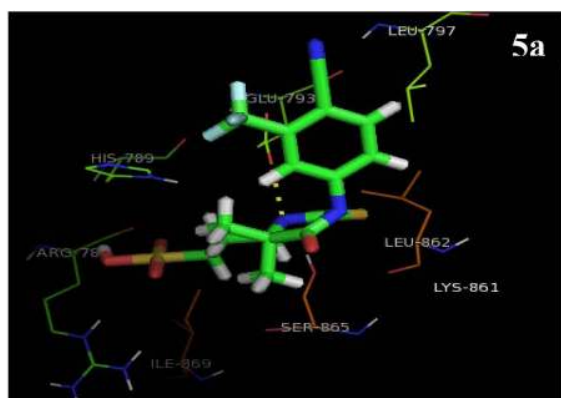
Figure 4. The pockets in androgen receptor (2AX6).

Table 11. Pocket Information for active sites.

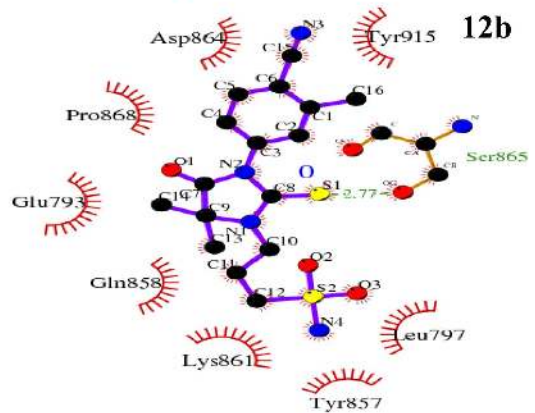
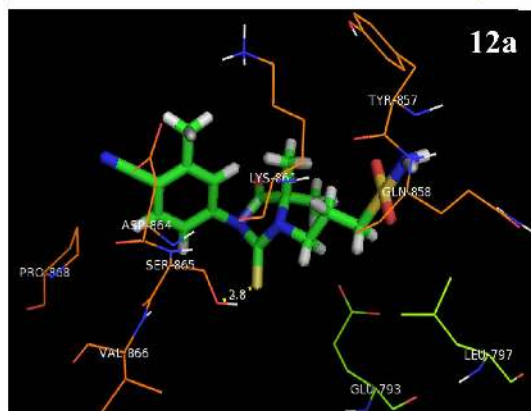
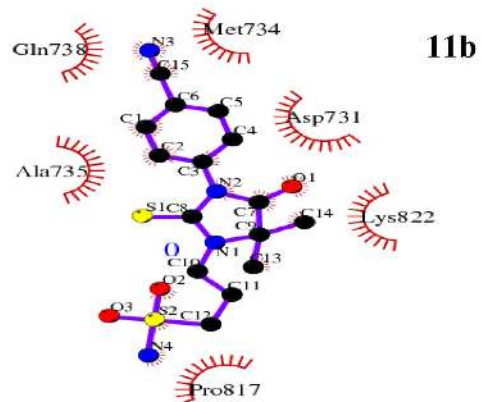
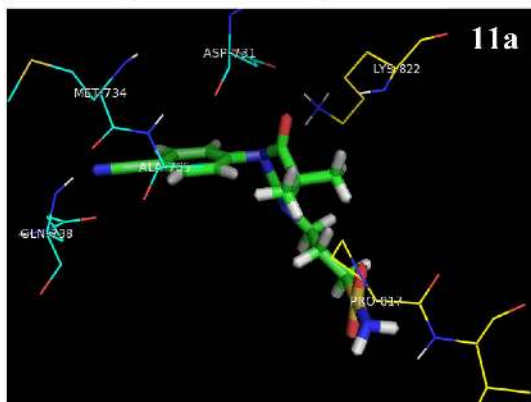
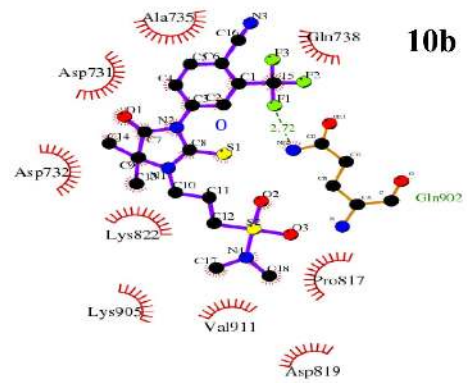
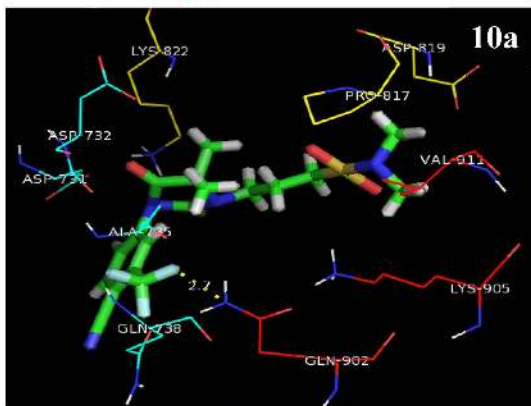
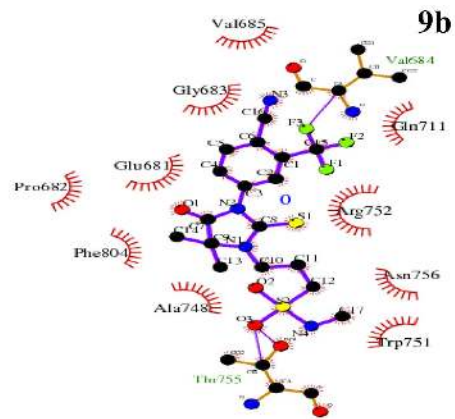
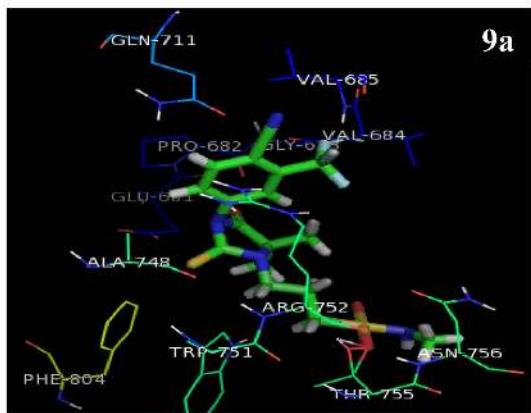
Pocket No	Amino Acid	position	Area	Vol
1	LEU ,MET, GLY, VAL	790, 787, 750, 746	32.00	16.20
2	ASN, ASP, HIS, LEU	823, 732, 729, 729	28.50	14.00
3	MET, TRP, LEU, GLU	895, 741, 712, 709	33.00	17.60
4	LYS, GLN, LYS, TYR, ALA	905, 902, 822, 739, 735	13.20	20.90
5	LEU, PRO, ILE, TYR, VAL	821, 817, 815, 739, 736	35.20	17.80
6	ALA, GLN, VAL, GLY	870, 867, 866,743	28.60	13.70
7	GLU, ILE, ARG, GLN	872, 869, 786, 783	30.40	15.70
8	LEU, LEU , CYS	838,810, 806	29.70	14.50
9	PHE, PHE, LEU, ILE, LEU	916, 856, 838, 835, 810	27.70	13.70
10	ARG, ASP, GLN,ASN, ASP, HIS	774, 695, 693, 691, 690, 689	30.20	15.10
11	SER, ILE, LEU, PHE	900, 882, 881, 878	30.40	16.90
12	PHE, ASN, LEU,ASP, LEU	827, 823, 821, 732, 728	27.90	13.70
13	MET,GLN,VAL,LYS	734,733,730, 720	21.40	14.90
14	PRO, LYS,VAL,ILE,ARG	913, 912, 911, 906, 871	24.50	16.80
15	VAL, ILE, ALA, HIS, MET, TRP,	903, 899, 877, 874, 742, 741	45.40	26.70
16	SER, ARG, LEU, MET, SER, SER	791, 788, 762, 761, 759, 753	35.20	19.30
17	LEU, GLN, GLU, LEU	805, 802, 678, 674	42.60	34.20
18	PHE,ARG, TRP, ALA, GLU	804, 752, 751, 748, 681	28.00	24.50
19	MET, GLN, ILE,MET, VAL, LEU	894, 738, 737, 734, 716, 712	30.50	25.40
20	ARG, MET, GLU, ASN, ARG	788, 775, 772, 771, 760	20.60	13.60
21	ILE, VAL, GLN, ILE, HIS, TRP,TYR, GLN	906, 903, 902, 898, 874,741, 739, 738	79.60	48.50
22	GLU, ARG, GLN, SER	872, 786, 783, 782	26.60	16.90
23	TRP, GLY, GLN, SER ASN PHE	796, 795, 792, 759, 758, 754	29.50	29.20
24	ASN, LEU, GLU, PHE, ASN	833, 830, 829, 826, 727	34.40	35.30
25	ALA, LYS, LEU, TRP, GLU, LEU	809, 808, 805, 718, 681, 677	73.70	53.80
26	ILE, ARG, GLU, LEU, PHE, PRO	841, 840, 837, 674, 673, 671	57.80	64.70
27	LYS, LEU, ASP, PHE, ARG, PHE	883, 880, 879, 876, 779, 697	85.00	57.80
28	LEU, ALA, SER, ASP, PRO, GLN,ASP, HIS	700, 699, 696, 695, 694, 693, 690, 689	91.00	68.50
29	VAL,LYS, LYS, ASP, VAL, PRO, TYR	911, 910, 905, 819, 818, 817, 739	62.20	79.60
30	PHE, TYR, ILE, GLN, ASP, LEU, ARG, SER, LEU, LEU	916, 915, 914, 867, 864, 863, 831, 814, 811, 810	158.7	114.3
31	TYR, PRO, ILE, HIS, ARG, ALA, PRO, GLN, ILE, SER, LEU, LEU, GLY, SER, TYR	915, 913, 906, 874, 871, 870, 868, 867, 815, 814, 811, 744, 743, 740, 739	246.9	236.7
32	ILE, MET, PHE, LEU, ALA, PHE, LEU, ALA, PHE, LEU, MET, MET, PHE, ARG, MET, VAL, MET, MET, TRP, GLN, GLY, LEU, ASN, LEU, LEU	899, 895, 891, 880, 877, 876, 873, 787, 780, 764, 752, 749, 746, 745, 742, 741, 711, 708, 707, 705, 704, 701	379.7	473.2
33	LYS, PRO, ALA, PHE, TYR, ASN, ARG, ALA, MET, LEU, TRP, VAL, HIS, GLN, VAL, VAL, GLY, PRO, GLU	808, 766, 765, 764, 763, 756, 752, 748, 745, 744, 718, 715, 714, 711, 685, 684, 683, 682, 681	347.3	429.8



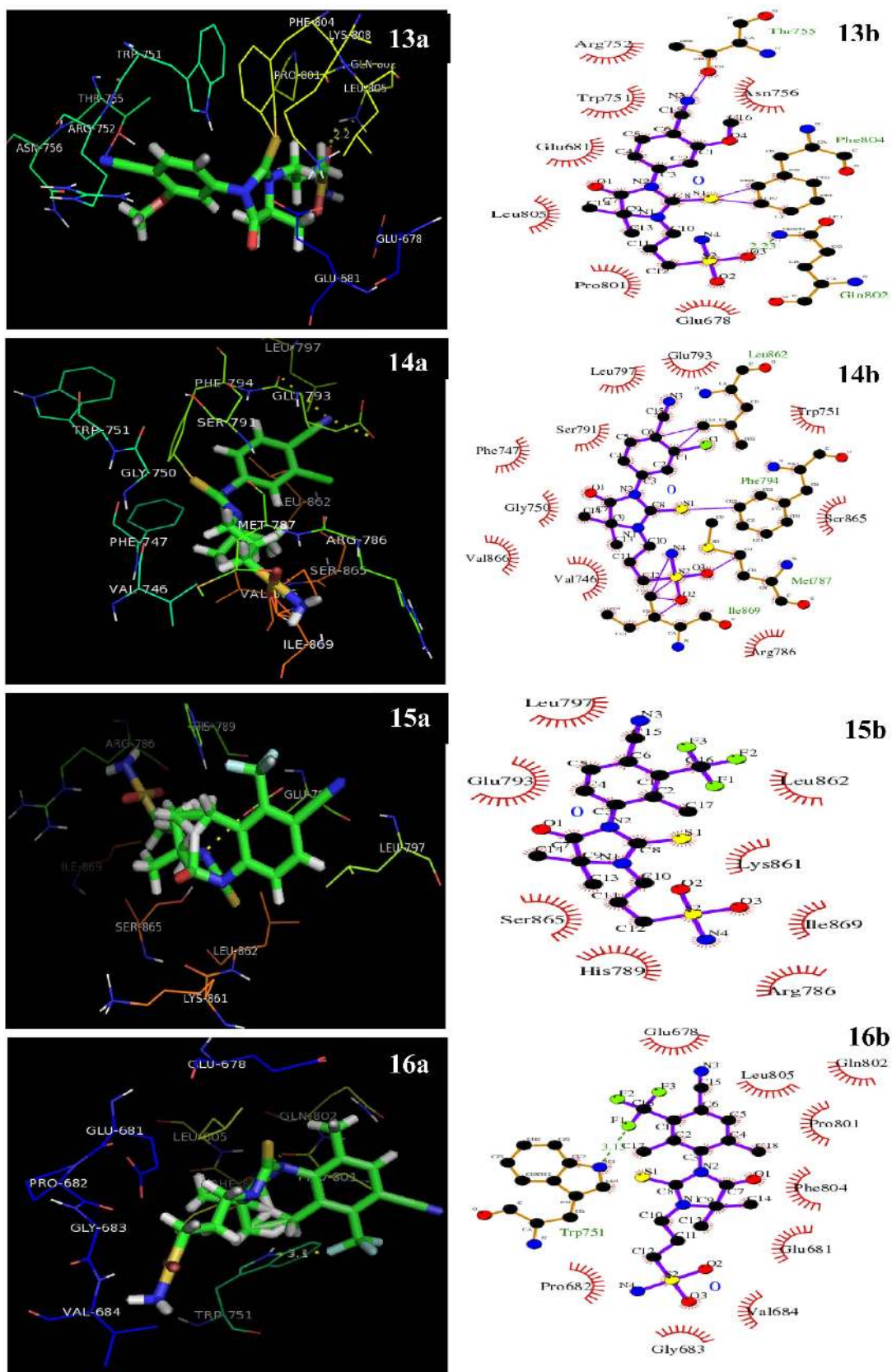
(a)



(b)



(c)



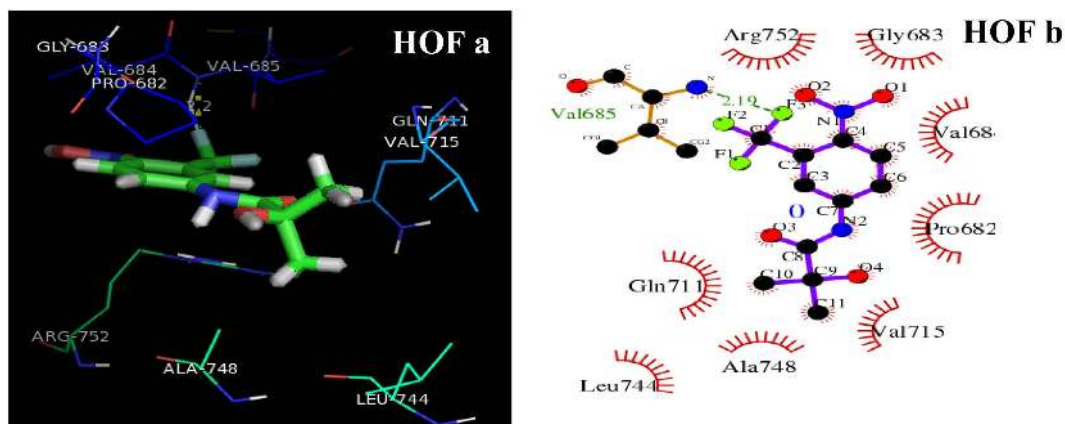


Figure 5. (a) 1a-4a) binding modes of compound 1 - 4 visualized using pymol software respectively, and 1b-4b) Ligplot + results showing the interactions of compounds 1 - 4 respectively with 2AX6; (b) 5a-8a) binding modes of compound 5 - 8 visualized using pymol software respectively, and 5b-8b) Ligplot + results showing the interactions of compounds 5 - 8 respectively with 2AX6; (c) 9a-12a) binding modes of compound 9 - 12 visualized using pymol software respectively, and 9b-12b) Ligplot + results showing the interactions of compounds 9 - 12 respectively with 2AX6; (d) 13a-16a) binding modes of compound 13 - 16 visualized using pymol software respectively, and 13b-16b) Ligplot + results showing the interactions of compounds 13 - 16 respectively with 2AX6; (e) HOF a) binding modes of hydroxyflutamide visualized using pymol software and HOF b) Ligplot + results showing the interactions of hydroxyflutamide with 2AX6.

the protein, but got the hydrophobic interaction with Glu793, Lys861, Leu862, Arg786, Ile869, His789, Ser865 and Leu797. Also, compound **6** had no hydrogen bond interaction with the protein, but got the hydrophobic interaction with Glu793, Lys861, His789, Ile869, Arg786, Ser865, Leu862 and Leu797. However, compound **7** was found to show three hydrogen bonds interaction with Ala735, Gln902 and Lys822 a distance of 2.61, 3.19 and 2.85 respectively, and the hydrophobic interaction with Tyr739, Lys910, Pro817, Asp732, Asp731, Asp819 and Gln738. Moreover, compound **8** had no hydrogen bond interaction with the protein, but got the hydrophobic interaction with Lys717, Val713, Met734, Glu893, Glu897, Gln738, Met894, Leu712 and Val716. However, compound **9** had no hydrogen bond interaction with the protein but got two external bonds with Val 684 and Thr755. Furthermore, this compound got hydrophobic interaction with Trp751, Asn756, Arg752, Gln711, Val685, Gly683, Glu681, Pro682, Phe804 and Ala748. On the other hand, compound **10**, was found to show one hydrogen bond interaction with Gln902 a distance of 2.72 and the hydrophobic interaction with Gln738, Ala735, Asp731, Asp732, Lys822, Lys905, Val911, Asp819 and Pro817. Compound **11** had no hydrogen bond interaction with the protein, but got the hydrophobic interaction with Pro817, Lys822, Asp731, Met734, Gln738 and Ala735. On the other hand, compound **12**, was found to show one hydrogen bond interaction with Ser865 a distance of 2.77 and the hydrophobic interaction with Tyr915, Asp864, Pro868, Glu793, Glu793, Gln858, Lys861, Tyr857 and Leu797. Compound **13**, was found to show one hydrogen bond interaction with Gln802 a distance of 2.23, but got two external bonds with Phe804 and Thr755, and the hydrophobic interaction with Asn756, Arg752, Trp751, Glu681, Leu805, Pro801 and Glu678. However, compound **14** had no hydrogen bond interaction with the protein, but got four external bonds with Ile869, Met787, Phe794 and Leu862, also this compound got hydrophobic interaction with Arg786, Ser865, Glu793, Leu797, Ser791, Phe747, Gly 750 and Val746. However, compound **15** had no hydrogen bonds interaction with the protein, but got hydrophobic interaction with Arg786, Ser865, Glu793, Leu797, His789, Ile869, Lys861, and Leu862. On the other hand, compound **16** was found to show one hydrogen bond interaction with Trp751 a distance of 3.13, but got the hydrophobic interaction with Pro682, Gly683, Val684, Glu681, Phe804, Pro801, Leu805, Gln802 and Glu678. The interactions of hydroxyflutamide with the functional residues of 2AX6 presented in figure in (**Figure 5(e)**), it shows one hydrogen bond interaction With Val 685 a distance of 2.19, but got the hydrophobic interaction with Arg752, Gly683, Val684, Pro682, Val715, Ala748, Leu744 and Gln711.

3.11. ADMET

As discussed earlier, physicochemical properties such as molecular weight MiLogP and TPSA of all a title com-

pounds follow Lipinski's Rule of Five. Furthermore, online software admetSAR (<http://www.admetexp.org/predict/>), were used to generate in silico pharmacokinetics parameters for all compounds in order to estimate their drug-like-ness properties. Various ADMET parameters are characterized for compounds **4** and hydroxyflotamide using in silico module admetSAR. The results of the analysis can be seen in **Table 12**. The analysis includes human intestinal absorption, blood–brain barrier penetration, Caco-2 permeability, P-glycoprotein substrate and inhibitor, CYP450 substrate and inhibitor (CYP1A2, 2C9, 2D6, 2C19, and 3A4), hERG inhibitors, AMES mutagenicity, carcinogens, fathead minnow toxicity, honey bee toxicity, aqueous solubility and Tetrahymena pyriformis toxicity. As per **Table 12**, hydroxyflotamide is a carcinogen as compared to the non-carcinogenic nature of the compound **4**. The major limitation of hydroxyflotamide is its CYP450 3A4 substrate nature, lead to high drug–drug interaction and interruption in the metabolism of the drug combination. The compound **4** overcomes that limitation. Furthermore, the compound **4** is AMES non-toxic and non-carcinogens with compared to standard drug hydroxyflotamide.

Table 12. Prediction of ADMET profiles of the most active compound (4) and droxyflutamide.

Parameter	Compound 11 (Most active)		Hydroxyflotamide	
Absorption	Result	Probability	Result	Probability
Blood-Brain Barrier	BBB+	0.8199	<u>BBB-</u>	0.8396
Human Intestinal Absorption	HIA+	0.9623	<u>HIA+</u>	0.9625
Caco-2 Permeability	Caco2-	0.6938	<u>Caco2-</u>	0.5898
P-glycoprotein Substrate	Non-substrate	0.5229	<u>Non-substrate</u>	0.6835
P-glycoprotein Inhibitor	Non-inhibitor	0.6193	<u>Non-inhibitor</u>	0.6634
	Non-inhibitor	0.9403	<u>Non-inhibitor</u>	0.8954
Renal Organic Cation Transporter	Non-inhibitor	0.8429	<u>Non-inhibitor</u>	0.9684
Distribution Metabolism				
CYP450 2C9 Substrate	Non-substrate	0.7697	<u>Non-substrate</u>	0.8100
CYP450 2D6 Substrate	Non-substrate	0.8041	<u>Non-substrate</u>	0.8308
CYP450 3A4 Substrate	Non-substrate	0.5490	<u>Substrate</u>	0.5225
CYP450 1A2 Inhibitor	Non-inhibitor	0.8211	<u>Non-inhibitor</u>	0.6559
CYP450 2C9 Inhibitor	Non-inhibitor	0.6976	<u>Inhibitor</u>	0.5266
CYP450 2D6 Inhibitor	Non-inhibitor	0.7479	<u>Non-inhibitor</u>	0.8475
CYP450 2C19 Inhibitor	Non-inhibitor	0.6359	<u>Inhibitor</u>	0.5260
CYP450 3A4 Inhibitor	Non-inhibitor	0.8317	<u>Inhibitor</u>	0.5374
CYP Inhibitory Promiscuity	Low CYP Inhibitory Promiscuity	0.5981	<u>High CYP Inhibitory Promiscuity</u>	0.5357
Excretion Toxicity				
Human Ether-a-go-go-Related Gene Inhibition	Weak inhibitor	0.9390	<u>Weak inhibitor</u>	0.9957
AMES Toxicity	Non AMES toxic	0.5689	<u>AMES toxic</u>	0.5150
Carcinogens	Non-carcinogens	0.6999	<u>Carcinogens</u>	0.5530
Fish Toxicity	High FHMT	0.9702	<u>High FHMT</u>	0.9984
Tetrahymena Pyriformis Toxicity	High TPT	0.8713	<u>High TPT</u>	0.9757
Honey Bee Toxicity	Low HBT	0.8241	<u>Low HBT</u>	0.8084
Biodegradation	Not ready biodegradable	0.9876	<u>Not ready biodegradable</u>	1.0000
Acute Oral Toxicity	III	0.5839	<u>III</u>	0.5581
Carcinogenicity (Three-class)	Non-required	0.5815	<u>Non-required</u>	0.4472

Acute Oral Toxicity: Category III includes compounds with LD50 values greater than 500 mg/kg but less than 5000 mg/kg. Carcinogenicity (three-class): Carcinogenic compounds with TD50 (tumorigenic dose rate 50) B10 mg/kg body wt/day were assigned as "Danger," those with TD50 [10 mg/kg body wt/day were assigned as "Warning," and non-carcinogenic chemicals were assigned as "Non-required." Probability indicates scale between 0 and 1. Parameters indicating difference between the most active and the least compound have been highlighted in bold letters.

4. Conclusion

The *in silico* study of 5,5-dimethylthiohydantoin derivatives gives promise results for using these compounds as an androgen antagonist. The title compounds bind with more competence to the binding site of similar to hydroxyflutamide. Our study has chosen 16 molecules, which demonstrate the better result in silico analysis with better binding efficiency (in terms of docking score) towards androgen receptor than that of hydroxyflutamide. Hydroxyflutamide is a carcinogen as compared to the non-carcinogenic nature of the compound **4**. On the other hand, the compound **4** is AMES non-toxic and non-carcinogens with compared to standard drug hydroxyflutamide. Hence, it has been predicted that all the title compounds can possibly act as new leads for the treatment of prostate cancer as they possess androgen antagonist activity. These results may be used in future experiments to investigate the interactions of 5,5-dimethylthiohydantoin derivatives with the androgen receptor, or may be used *in vivo* experiments to test their effects on the abilities of treatment of prostate cancer.

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