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*Simulation, and Drug-Likeness Scoring of (E)-1-[2-(3,4-Dimethylphenyl) Diazen-2-Ium-1-Yl] Naphthalen-2-Olate as a Corticosteroid Side-Chain-Isomerase Inhibitor*

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**Abstract:** In this article, the optimized structure and their associated properties of the  $(E)$ -1-[2-(3,4-Dimethylphenyl)diazen-2-ium-1-yl]naphthalen-2-olate compound (EDNO) were obtained and evaluated using The density functional theory DFT at the  $(B3LYP/6-311G++(d,p))$  level in the gas phase, To quantify the intermolecular interactions, Hirshfeld surface(HS) analysis was used, HS and 2D fingerprints indicate H⋅⋅⋅H (52%) and C− H⋅⋅⋅C (26.7%) as the most relevant intermolecular interactions in the crystal packing of EDNO. the reduced density gradient (RDG) method was used to reveal and distinguish between attractive interactions such hydrogen bonds, repulsive interactions and van der Waals interactions. Further, molecular docking, binding free energy calculations, and ADMET prole of the title compound was carried out to determine the binding affinity and toxicity. A 100 ns molecular dynamics (MD) simulation was performed to evaluate the binding stability of the compound EDNO/2WV2 complex using Desmond. Binding free energy of the complex was computed for 100 trajectory frames using the MM-GBSA approach.

*Keywords:* Azo Compounds; Hirshfeld surface analysis; DFT calculations; ADMET; Molecular docking; MD simulation.

# **1. Introduction**

The chemical (E)-1-[2-(3,4-Dimethylphenyl) diazen-2-ium-1-yl].-naphthalen-2-olate belongs to the family of azo compounds, which are distinguished by the presence of an azo group (- N=N-) that connects two aromatic structures. Azo compounds are essential in a variety of disciplines, particularly as dyes, pigments, and coloring agents in the textile and food industries [1,2]. Their chemical structure confers distinctive spectrum features, making them useful as fluorescence probes and pH indicators. Furthermore, because of

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their propensity to undergo reversible photochemical changes, these compounds have sparked interest in their possible use in optoelectronic materials and information storage devices. The growing interest in these compounds arises from their adaptability and the ability to adjust their properties via substitutions on the aromatic rings, paving the path for novel applications in chemistry and biology [3-8].

For instance, in medicinal chemistry, azo compounds have been studied for their antimicrobial and anticancer properties, demonstrating their potential as therapeutic agents [9,10]. Furthermore, their involvement in the development of advanced materials like liquid crystals and polymers highlights their importance in material science [11-14]. Azo compounds are also significant in environmental chemistry, where they are analyzed for their behavior and degradation in both natural and engineered systems. Their persistence and transformation products can affect environmental health, making it essential to understand their fate for effective environmental protection [15-17]. In analytical chemistry, azo compounds are vital as reagents and indicators, aiding in the detection and quantification of various analyses [18-20]. The ability to customize their chemical and physical properties through strategic substitutions enhances their applicability across various scientific domains, showcasing their wideranging and significant impact in modern science and industry.

In this article, we conducted a comprehensive study of the molecule (E)-1-[2-(3,4-Dimethylphenyl) diazen-2-ium-1-yl]-naphthalen-2-olate. Our study began with experimental structural determination using X-ray diffraction, followed by a comparison with the theoretically determined geometry via density functional theory (DFT) using the B3LYP functional with the basis set  $6-311G++(d,p)$ . The agreement between theoretical and experimental results confirmed the reliability of the method used, enabling us to characterize the molecule's properties. We examined the compound's intermolecular interactions using reduced Density Gradient Analysis (RDG). The RDG plot provided insights into steric effects, repulsive interactions, and hydrogen bonding within the compound. Notably, steric repulsion was most apparent in the aromatic and benzene rings, while the presence of green regions indicated hydrogen bonding interactions. Complementing this analysis, we utilized a Hirshfeld (HS) surface analysis to visually illustrate the contributions of intermolecular interactions through a 2D fingerprint plot. Our findings revealed that hydrogen-hydrogen (H---H) interactions were predominant, constituting 52.0% of the total, whereas oxygen-oxygen (O---O) interactions were relatively rare, comprising just 0.2%. This detailed analysis of intermolecular interactions has enabled us to gain a better understanding of protein-ligand interactions, which are essential for biological processes. These interactions also play a crucial role in the design of new drugs based on molecular structure. To explore the structure-activity relationship and study interactions with the appropriate target protein, we carried out a molecular docking study for the compound under study. In addition, we assessed the quality and similarity of the drug. Finally, we reinforced our biological study with a molecular dynamics simulation, confirming the stability of the complex formed between our compound and the protein, thus validating the application of our compound as a potential active ingredient in a drug.

## **2. Computational Method**

For the theoretical calculation part ,DFT calculations were carried out using the Gaussian 16 software [21] and the Gauss View 6 program [22] . The hybrid functional B3LYP with the basis set 6- 311G++(d,p) was used in all calculations. The experimental X-ray structure was used as starting geometry for the optimization task. The Hirshfeld surface maps and fingerprint plots were performed using Crystal Explorer 21.5 program [23]. Using the Multiwfn software [24], the scatterplots of the reduced density gradient (RDG) were carried out. Similarly, the 3D isosurfaces are visualized with the use of VMD software [25]. The study of ligandprotein interactions is performed using AutoDock Vina [26] AutoDock Tools [27] and Discovery Studio Visualizer [28] to prepare the target and the ligand, perform molecular docking and visualize the result respectively. The Molecular Dynamics (MD) simulation was conducted using the Desmond Package from the Schrödinger software suite.

#### **3. Results and discussion**

# **3.1. Molecular geometry**

The comparison between experimental (X-ray) and computational (B3LYP/6-311G++(d,p)) results reveals several noteworthy observations regarding the structural properties of the (E)-1-[2-(3,4- Dimethylphenyl)diazen-2-ium-1-yl]-naphthalen-2-

olate compound, The molecular structure with atomic labeling is depicted in Fig.1.

Generally, the computed bond lengths closely match the experimental values, indicating the accuracy of the computational approach in predicting molecular geometry. The C–C distances in both benzene rings, as measured experimentally and calculated, are nearly identical, hovering around 1.38 Å, which aligns well with established values in the literature [29]. The N-N distance is 1.2946 Å (X-ray) and 1.30231 Å (B3LYP). The O-C distance is 1.2873 (X-ray) and 1.25266Å (B3LYP). The bond angles derived from computational methods align well with experimental measurements, such as the bond angle of N2—N1— C9 is 117.66◦ (X-ray) and 120.00◦ (B3LYP), suggesting consistency in the molecular conformation predicted by theory and observed experimentally.

Dihedral angles, crucial for understanding molecular conformations, exhibit good agreement between experimental and computational data, the dihedral angle of C9—N1—N2—C11A is  $-178.13$ ∘ (X-ray) and -179.99◦ (B3LYP), indicating reliable predictions of molecular orientation.

The congruence between experimental and computational results underscores the efficacy of the computational approach (B3LYP/6-311G++(d,p)) in elucidating the structural characteristics of the title compound molecule. These findings provide valuable insights for further investigations into the chemical and physical properties of the title compound and its potential applications.









**Figure 1.** 3D structure of EDNO with atomic labeling.

## **3.2. RDG (Reduced Density Gradient Analysis)**

We conducted the topological analysis to delve into the subtle interactions between various compounds in greater detail. The Atoms in Molecules (AIM) theory is widely employed for this purpose, as it allows us to identify different types of interactions across various molecular systems and analyze the electron density at critical binding points (BCP) [30]. By utilizing topological parameters such as bond energy (E int =  $V(r)/2$ ), kinetic Hamiltonian energy  $(H(r) = G(r) + V(r))$ , potential energy density  $(V(r))$ , Lagrangian kinetic energy  $(G(r))$ , electron density Laplacian ( $\nabla$ 2  $\rho$ (r)), and electron density  $(\rho(r))$ , we were able to observe hydrogen bonding properties among the compound structures. We used an optimized geometric structure to pinpoint critical bond points (BCP) and ring critical points (RCP) within the CEFP crystal structure. The analysis was conducted using the Multiwfn program [24]. Figure 2 displays scatter diagrams and gradient isosurfaces representing the title compound. In Figure 2, we observe the molecular interactions' patterns within the RCP and BCP, as identified by the Multiwfn software. At a ρBCP value of 0.01 and a Laplacian value of 0.03, electron density indicates molecular stability.

Molecular stability hinges on various interactions, including both intra- and intermolecular forces. These interactions can be discerned through reduced density gradient (RDG) analysis, a method rooted in the examination of non-covalent interactions (NCI) within the system [31]. In Figure 2 (a), we present the NCI-RDG analysis conducted with an isosurface value of 0.5. The reduced density gradient (RDG) encompasses fundamental aspects, involving density and its first derivative, as outlined in prior literature [32]. The RDG isosurface surrounding specific regions delineates the interactions among molecules, as illustrated in Figure 2 (b). The dotted red areas signify steric effects, indicating repulsive interactions. Notably, the most pronounced steric repulsion occurs at the centers of two aromatic rings engaged in  $\pi$ -π stacking interactions [33], as well as at the benzene ring, highlighted in the RDG chart as the red zone. The overlapping red and green regions around the (O, H) of the two benzene rings and (C, C) of the methyl group denote repulsive areas. Conversely, the green regions denote hydrogen-bonding interactions, which contribute to the stability of the title compound's molecular system.



**Figure 2.** (a) the gradient isosurfaces and (b) the scatter diagram of the title compound



**Figure 3**. Intermolecular interactions represented in red circle point collapsing on the *d* norm area in baxis for the crystal structure of the title compound.



**Figure 4**. Hirshfeld surface analysis of the title compound with d e, d i, d norm, shape index, curvedness, and fragment patch

#### **3.3. Hirshfeld Surface (HS) analysis**

Hirshfeld Surface (HS) analysis is a technique that divides molecular crystal space into discrete, nonoverlapping molecular entities. With the development of HS fingerprint plots [34], which provide an easily understood representation of intermolecular contacts in crystals, significant advancements occurred. Further developments include the use of decoration, a mapping method that applies a color scale onto the HS to graphically represent scalar properties and quickly obtain insights into molecular environments in the crystalline state [35]. Feature decomposition is another innovation that helps identify specific interactions. The Crystal Explorer 21.5 [36] package was utilized to examine Hirshfeld surfaces and their corresponding fingerprint plots. The crystallographic information file (CIF) was used to enter the required data for this analysis [37]. Fig. 3 shows the three-dimensional Hirshfeld surface plot on dnorm. The visualization illustrates the interactions between hydrogen and carbon atoms by revealing four different red dots, the distance between nearby molecules is 2.513 Å. Fig. 3, illustrates the Hirshfeld surface analysis (HSA) conducted on the title molecule, presenting results via parameters such as d norm, d e, d i, shape index, fragment patch, and curvedness. Notably, the shape

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index map reveals the presence of  $\pi$ - $\pi$  stacking interactions among molecules, as indicated by adjacent blue and red triangles. Fig. 4 displays the fingerprint plot (100%) of the title compound, specifically plotting d<sup>i</sup> against de. This plot illustrates the distances between nuclei starting from the surface and the nearest nuclei within the molecule's surface, spanning from 0.9050 to 2.5474 Å. Fig. 4, displays by illustration how the Hirshfeld surface (HS) contributes to the overall molecular surface. It also highlights the specific proportions of different types of intermolecular contacts within the molecule. Essentially, it visually breaks down the contributions of intermolecular interactions using 2D fingerprint plotting. The most prevalent interactions involve hydrogen (H···H) interactions, constituting 52.0% of the total. This dominance is attributed to the abundance of hydrogen atoms on the surface of CEFP. Following closely are carbonhydrogen (C···H/H···C) interactions, contributing 26.7%. Additionally, carbon-carbon  $(C^{\cdots}C)$ interactions make up 5.8% of the total. Furthermore, nitrogen-hydrogen/hydrogennitrogen  $(N''H/H''N)$  interactions account for 4.5%, while oxygen-carbon/carbon-oxygen  $(O^{\cdots}C/C^{\cdots}O)$  interactions represent 0.3%. A minimal contribution is seen from oxygen-oxygen  $(O \cdots O)$  interactions, constituting only 0.2%. These proportions are visualized in Figure 5.



**Figure 5.** 2D fingerprint graphs depicting the total contributions to the total HS surface region and the single percentages of the different intermolecular interactions.

# **3.4. Prediction of Activity Spectra for Substances (PASS) analyses**

PASS (Prediction of Activity Spectra for Substances) is a crucial computational tool in cheminformatics and drug discovery. It predicts the biological activities of chemical compounds by analyzing their structural formulas and drawing from a knowledge base of structure-activity relationships (SAR) derived from experimental data [38]. This tool is invaluable in early drug development stages, assisting researchers in narrowing down the pool of compounds for further

investigation. Available at http://way2drug.com/PassOnline, the PASS Software can predict over 4000 types of biological activity, including pharmacological effects, mechanisms of action, toxic and adverse effects, and interactions with metabolic enzymes and transporters [39]. It claims an impressive 90% validity rate. The outcomes of PASS calculations are expressed as Pa (probability of active molecule) and (probability of inactive molecule) [40]. A Pa value of 0.7 or higher indicates a high likelihood of experimental pharmacological activity, while

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values between 0.5 and 0.7 suggest a lower likelihood. A Pa value below 0.5 indicates a lower likelihood of discovering activity through experimentation but may also imply a chance of discovering a novel chemical [41]. These parameters guide researchers in prioritizing compounds for further experimental investigation. The results presented in Table 2 were interpreted and applied flexibly. (EDNO) compound exhibited the highest Pa value (0.731) and a Pi value of 0.007 for its activity as a corticosteroid side-chain isomerase inhibitor. This type of medication functions by inhibiting the enzyme corticosteroid side-chain isomerase [42]. This enzyme plays a crucial role in the synthesis of corticosteroids, hormones that are involved in various physiological processes, including immune response, metabolism, and stress regulation [43, 44]. One example of a corticosteroid side-chain isomerase inhibitor is aminoglutethimide, which was historically used to treat Cushing's syndrome [45]. Another example of a corticosteroid side-chain isomerase inhibitor is fluconazole. Although primarily used as an antifungal medication, fluconazole can also inhibit the enzyme 14-alphademethylase, which plays a role in cortisol synthesis [46].

Pa	Pi	<b>Activity</b>
0,731	0,007	Corticosteroid side-chain-isomerase inhibitor
0,642	0,007	Histamine release inhibitor
0,643	0,014	<b>Platelet aggregation stimulant</b>
0,691	0,072	Ubiquinol-cytochrome-c reductase inhibitor
0,631	0,015	Aspartate-phenylpyruvate transaminase inhibitor
0,636	0,025	2-Dehydropantoate 2-reductase inhibitor
0,636	0,032	<b>Glucose oxidase inhibitor</b>
0,612	0,014	Amine dehydrogenase inhibitor
0,621	0,026	Alkane 1-monooxygenase inhibitor
0,665	0,075	Aspulvinone dimethylallyltransferase inhibitor

**Table 3.** Physicochemical and Pharmacokinetic properties of the title molecule.



#### **3.5. Druglikeness and ADMET analysis**

Pharmaceutical companies invest heavily in drug discovery, but clinical trial failures due to pharmacokinetic issues or toxicities are costly. AIbased computational methods, such as machine learning, leverage data to predict these properties, including absorption, distribution, metabolism, excretion, and toxicity (ADMET), potentially reducing risks and accelerating drug development [47]. ADMET profiling is a critical step in drug development, involving the assessment of a compound's Absorption, Distribution, Metabolism, Excretion, and Toxicity. These properties determine a drug's efficacy and safety [48]. The SWISS ADME tool is a widely used web-based application that evaluates the physicochemical properties of drug candidates. This tool provides reliable predictive testing for several key parameters: Absorption evaluates how well a compound is absorbed into the bloodstream. Distribution predicts the distribution of the compound throughout the body and its potential to reach target tissues. Metabolism assesses the compound's metabolic stability and the likelihood of it being broken down by the body. Excretion predicts the routes and rates at which the compound is excreted from the body. Toxicity assesses the potential toxic effects of the compound [49]. Using the SWISS ADME tool involves inputting the molecular structure of a compound to receive predictions on its ADMET properties. In silico ADMET analysis utilizes computational models, such as Lipinski's Rule of Five, which considers factors like molecular weight, the number of hydrogen bond acceptors and donors, and the logP value. Compounds that do not meet these criteria are likely to have poor bioavailability [50]. Recent advancements in machine learning have improved the accuracy of ADMET predictions by analyzing large datasets, significantly reducing the time and costs in drug development [51]. The drug-likeness parameters of the little compound are a reported in Table 3 and were expressed by the Lipinski's rule of five. According to this guideline, oral medications must meet the following requirements: POLAR (Polarity): 20Å 2 < TPSA < 130Å2, LIPO [Lipophility]:  $-0.7 < XLOGP3 < +5.0$ , INSOLU [Insolubility]:  $-6 <$  Log S (ESOL)  $< 0$ , FLEX [Flexibility]: 0< Num. rotatable bonds < 9, size:

150g.mol-1< MV < 500g.mol-1, INSATU [Insaturation]:  $0.25 <$  Fraction Csp3  $<$  1 [52]. The results indicate that the title compound aligns well with Lipinski's Rule of Five, suggesting favorable bioavailability. Additionally, rule-based filters from Lipinski, Ghose, Veber, and Egan were employed to evaluate the predicted drug-like properties of the compound. The compound has two rotatable bonds, which is a crucial parameter for molecular flexibility and should be less than 10. The TPSA is within the acceptable range, indicating that the compound has suitable physicochemical properties to be used as a pharmaceutical drug candidate. The BOILED-Egg model classifies molecules based on their likelihood of intestinal absorption (HIA) and blood-brain barrier (BBB) penetration. The compound is predicted to have good permeability through the blood-brain barrier, excellent intestinal absorption, and it is not a substrate for P-glycoprotein, suggesting favorable pharmacokinetic properties for oral administration and central nervous system targeting. This comprehensive analysis indicates that the compound meets all the criteria for drug-likeness according to Lipinski's Rule of Five and is wellsuited for further drug development and testing.



**Figure 6.** a) Oral bioavailability radar and b) bloodbrain barrier (BBB) permeability [BOILED-Egg] diagram of title compound.

## **3.6. Molecular docking against targeted receptor**

In recent years, Molecular Docking has become integral to in-silico drug development. This method predicts how small molecules interact with proteins at the atomic level, aiding in understanding biochemical processes [53]. It relies on highresolution 3D representations of target proteins obtained through techniques like X-ray crystallography. Molecular Docking facilitates the study of small molecule behavior within protein binding sites, including nutraceuticals. Its structurebased approach is essential for accurate predictions. [54] According to the results of PASS prediction, the target protein is identified with PDB ID: 2WV2 (Method: X-ray diffraction, Resolution: 2.70 Å) was downloaded from protein database website (https://www.rcsb.org) [55]. The PDB ID: 2WV2 represents the crystal structure of the enzyme responsible for cholesterol biosynthesis, human lanosterol 14-alpha demethylase (CYP51). [56]. The CYP51 gene is highly conserved throughout different species, and it performs a comparable function in the manufacture of sterols in both fungi and humans [57]. The structural similarities between CYP51 in humans and fungi makes it a crucial target for antifungal medications, as inhibitors can impact both types of enzymes [58]. Comprehending the mechanism of action of 2WV2 aids in the development of specific antifungal drugs that specifically target fungal CYP51 while minimizing any adverse effects on the human enzyme.

## **3.7. Protein preparation**

The protein preparation was conducted using BIOVIA Discovery Studio Visualizer [28]. This procedure involved selecting the target protein, removing water molecules, heteroatoms, and coligands from the active site, and then adding polar hydrogen atoms [59]. The final protein structure was saved in the .pdb format. The molecular docking of the little compounds was performed using AutoDock vina program [26].

## **3.8. Docking and interactions study**

Docking simulations were conducted, which involved creating a 3D grid box with the following dimensions: center x=-48.422, center y=11.915, center  $z=1.617$ , and grid cells of  $28x28x28 \text{ Å}^3$ . The

selection of this grid design was purposely made to thoroughly cover the active site pocket of the target molecules. AutoDock is used to assess the binding affinity of the ligand, which helps in identifying the most favorable ligand structure and rotation in relation to the receptor (protein). Table 4 provides the binding energies and RMSD values for the nine potential conformations of the synthesized ligand in (PDB ID: 2WV2) .Based on the findings, it can be concluded that the optimal docking position is in mode 1, with a binding energy of -10.2 kcal/mol. The binding energy suggests a docking position that is highly favorable, similar to the highest scores shown in molecular docking studies for strong interactions between a ligand and receptor [57]. Mode 1 has both the highest binding affinity and an RMSD value of 0.000, indicating a flawless match with no deviation. This highlights its stability and optimal interaction inside the binding pocket. The compound was found to interact with various amino acid residues via hydrogen bonding and hydrophobic interactions. It was further revealed that the hydrophobic interactions with various amino acid residues of CYP51 occur through the three aromatic rings in the compound. The detailed description of the types of interactions and the interacting amino acid residues is presented in Table 5. Two amino acid residues, namely SER296 and GLY428, interacted to form conventional hydrogen bonds with distances of 2.04184 Å and 3.22652 Å, respectively. This compound showed Pi-alkyl interactions with PRO355, LEU356, CYS422, and ALA291. The alkyl interactions of the compound occurred with ALA288, ALA291, and LEU134. Pi-Pi T-shaped interactions were formed when two aromatic rings were oriented perpendicular to each other, resembling a "T" shape, with one aromatic ring's edge facing the plane of the other aromatic ring, involving PHE415. Additionally, the compound also showed Pi-Sigma and amide-Pi stacked interactions with THR295 and ALA291, respectively. These interactions contribute to the stability, good affinity, and highest activity. Fig. 7 provides a visual representation of the intramolecular interactions. Part (a) shows the schematic diagram of the interactions including van der Waals, hydrogen bonds, and Pi interactions. Part (b) illustrates the 3D molecular docking pose within the binding pocket. Parts (c) and (d) offer additional perspectives, highlighting the spatial

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arrangement and interaction details of the compound within the protein structure. The study's comprehensive approach, including both quantitative (binding affinities, RMSD values) and qualitative (interaction types and distances) data, offers a thorough analysis of the docking results.

The compound shows a strong binding affinity and stability through various interactions, making it a promising candidate for further studies.

**Table 4** AutoDock Vina results of the binding affinity and RMSD values of different poses in little compound inhibitor of 2WV2.

<b>Mode</b>	<b>Affinity (kcal/mol)</b>	<b>RMSD l.b</b>	RMSD u.b.
	$-10.2$	0.000	0.000
	$-10.0$	1.622	7.210
	$-9.3$	3.484	7.066
4	$-8.8$	2.012	7.013
	$-8.6$	2.109	7.852
6	$-8.5$	3.714	5.702
	$-8.4$	5.581	9.773
8	$-8.4$	5.474	7.419
9	$-8.4$	3.866	5.402

**Table 5.** Distances. Types and location of intermolecular interactions formed from the residues of (PDB ID: 2WV2) and the molecule.







**Figure 7.** Intramolecular interactions that connect the structure to the 2wv2 protein



**Figure 8.** Validation of docking procedure through RMSD calculation of superimposed ligand Poses: retrieved and re-docked co-ligand in protein x-ray crystal structure (ID:2WV2).

### **3.9. Docking validation**

The docking process was validated through removing the present co-ligand inside the protein active site (ID: 2WV2), re-docking it using the same protocol including the grid parameters was unchanged in the process. It was done to ensure the inhibitor binds exactly to the active site cleft and must show less deviation compared to the actual cocrystallized complex. The re-docked complex was then superimposed on to the reference cocrystallized complex using PyMOL 2.3 and the root mean square deviation (RMSD) was calculated (Fig. 8). The co-ligand bound exactly to the active site with good binding energy of −8.12 kcal/mol and RMSD of 0.12  $A^{\circ}$  (<2 $A^{\circ}$ ).

#### **3.10. Methodology of molecular dynamics simulation**

The Molecular Dynamics (MD) simulation was conducted using the Desmond Package from the Schrödinger software suite [60], leveraging CUDA support on Nvidia RTX 3060Ti GPU machines. For this study, the top-docking scored model of the studied compound, Mol, along with the co-ligand

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Fluconazole (CID: 3365) as a reference, were selected as the starting coordinates. Before solvation, the complex was optimized, and its energy minimized using the OPLS\_2005 force field, which is known for its accuracy in predicting molecular geometries, conformational energies, and non-bonded interactions for proteins and small molecules [61]. Each system was then solvated using a three-point (TIP3P) water model within cubic periodic boundary conditions measuring 100  $\times$  100  $\times$  100 Å, ensuring at least 10 Å of space between the protein and the edges of the simulation box. Simulations were performed under physiological conditions at pH 7.0 in an NPT ensemble (constant number of particles, pressure, and temperature) with settings of 1 atm and 303.15 K, considering the expected ionization states of protein residues and utilizing periodic boundary conditions.

# **3.11. Results of molecular dynamics simulations**

Molecular dynamics simulations were performed for 100 nanoseconds on both the studied ligand Mol and the reference co-ligand STD, in complex with the target protein sterol  $14α$ -demethylase, also known as CYP51 (PDB ID: 2WV2). The primary objective was to evaluate the stability of the ligand Mol in comparison to the co-ligand STD within the protein's catalytic site. To achieve this, various parameters were analyzed, including Root Mean Square Deviations (RMSD) to measure the structural stability over time, and Root Mean

Square Fluctuations (RMSF) to assess the flexibility of the protein backbone. Additionally, solvent-accessible surface area (SASA) and radius of gyration (Rg) were calculated to evaluate the compactness and exposure of the protein. Principal Component Analysis (PCA) and free-energy landscape (FEL) plots were used to investigate the conformational changes and energy states of the system.

## **3.12. Root Mean Square Deviation studies (RMSD)**

The resulting MD trajectory was analyzed to compute the root mean square deviation (RMSD) of the protein's backbone atoms in complex with both the studied ligand Mol and the co-ligand STD, as well as the RMSD of the ligands when fitted onto the protein. This analysis highlights the ligands' movement or shift from their initial binding site in the protein's active site. Fig. 9 shows the RMSD plot for the biomolecular systems Mol-2WV2 and STD-2WV2, providing insights into their dynamic behavior. The RMSD analysis of the protein backbone revealed average deviations of 3.17 Å for Mol and 2.49 Å for STD, indicating comparable stability of the protein with both the studied ligand and the co-ligand. The RMSD values for the ligands fitted onto the protein were also analyzed, with average deviations of 2.47 Å for Mol and 1.9 Å for STD within the protein's catalytic site. Both ligands demonstrated significant stability, as indicated by their lower deviations (Fig. 9).



**Figure 9.** Root Mean Square Deviation plots of studied complex 2WV2-mol in contrast to the standard complex 2WV2-STD with the sterol  $14\alpha$ -demethylase, also known as CYP51 protein.



**Figure 10.** Root Mean Square Fluctuation plots comparing the studied complex 2WV2-Mol to the standard complex 2WV2-STD with the sterol  $14\alpha$ -demethylase, also known as the CYP51 protein.



**Figure 11.** Solvent Accessible Surface Area plots comparing the studied complex 2WV2-Mol to the standard complex 2WV2-STD with the sterol 14α-demethylase, also known as the CYP51 protein.

## **3.13. Root Mean Square Fluctuatons analysis (RMSF)**

Analysing atomic fluctuations through Root Mean Square Fluctuation (RMSF) in Molecular Dynamics (MD) simulations is pivotal for comprehending biomolecular dynamics. A higher RMSF value signifies increased flexibility within the biomolecule's region, while a lower RMSF value indicates rigidity. The RMSF profiles were derived from the fluctuations of alpha carbon atoms

in amino acid residues, visually depicted in Fig. 10. These residue-specific fluctuations were meticulously examined for the two complexes mol-2WV2 and STD-2WV2. RMSF average of recorded values was 1.18A° and 1.21A° for mol-2WV2 and STD-2WV2 complexes respectively, hinting at the stabilizing effect of studied compound on the protein residues.

#### **3.14. Solvent accessible surface area and radius of gyration analysis**

In our investigation, Solvent Accessible Surface Area (SASA) serves to quantify the portion of the protein that interacts with surrounding solvents within the simulation box. The SASA plot (Fig. 11) illustrates variations that cluster around a relatively stable mean value. Across the complexes 2WV2- Mol and 2WV2-STD, SASA averages were recorded as  $3366.52 \text{ A}^2$  and  $3304.65 \text{ A}^2$ respectively. Despite these fluctuations, there is no significant deviation from the mean SASA value across the studied compound complex simulations in contrast to the standard ligand complex, indicating a stable interaction between the protein and its surrounding solvent molecules.

Additionally, a radius of gyration (Rg) analysis was conducted to assess the structural attributes of the protein within the investigated system, in comparison to the co-ligand complex (Fig. 12). Generally, Rg changes were consistent across the studied system named 2WV2-mol, indicating stable structural characteristics. The recorded average Rg values for the protein structures were 22.1 Å and 22.3 Å for 2WV2-mol and 2WV2-STD, respectively. These minor discrepancies imply that the protein structures retained a similar compactness in both scenarios.



**Fig. 12** Radius of gyration plots comparing the studied complex 2WV2-Mol to the standard complex  $2WV2$ -STD with the sterol 14 $\alpha$ -demethylase, also known as the CYP51 protein

**Table 6** Percentage of Protein Secondary Structure Elements during 100ns MD Simulation across the studied system 2WV2-mol and co-ligand System 2WV2-STD.

<b>Complex</b>	Alpha-Helices	<b>Beta-Strands</b>	SSE%	Loops $\%$
2WY2-mol	32.27	8.16	40.43	58.57
2WY2-STD	36.59	7.91	44.50	55.50

### **3.15. Protein Secondary Structure analysis**

The protein's secondary structure elements, specifically alpha helices and beta strands, are vital in assessing its stability during the 100ns molecular dynamics simulation. A high percentage of secondary structure indicates that the residues are more rigid, whereas the presence of unstructured loop regions suggests flexibility and instability in the protein structure. Fig.13, illustrates the distribution of secondary structure elements by residue index throughout the protein structure. In the case of the molecule of interest the total SSE% of the protein is 40.43 while it is 44.50. These outcomes suggest that the residues of the sterol  $14\alpha$ -

demethylase, also known as the CYP51 protein maintain a stable, rigid structure predominantly in both systems which hint at the stability of formation of the complex with the protein of the studied ligand.



**Figure 13.** Monitoring Protein Secondary Structure Elements across 100ns MD Simulation: Comparison of systems 2WV2-mol and co-ligand system 2WV2-STD; Alpha-helices highlighted in orange, Betastrands in blue, and Loops in white.

## **3.16. Principle Component Analysis and Freeenergy Lanscape plots**

Principal Component Analysis (PCA) is a statistical method used to simplify protein dynamics data by reducing its dimensionality. This technique highlights the most important motions by sorting them from the largest to the smallest spatial scales. PCA achieves this by performing a linear transformation on a covariance or correlation matrix, which is created from the atomic coordinates that depict the protein's available degrees of freedom (DOF). These coordinates usually comprise Cartesian coordinates that describe atomic movements within trajectory conformations [62].

The PCA data calculated for 2WV2-mol and 2WV2-STD was utilized to create the free energy landscape (FEL) profiles of the systems (Fig. 14). The FEL derived from the PCA data reveals several energy transitions and metastable states, with a few notably large energy wells identified for both systems. However, longer simulations are needed to fully explore the global minima and stable conformations of the structures. The presence of an intense blue area in the last 200 frames of the 2WV2-mol, compared to 2WV2-STD, within the FEL plots indicates that the ligand plays a role in lowering the system's energy relative to the coligand system. This underscores the ligand's stabilizing effect over an extended simulation period (Fig. 15).

#### **3.17. MMGBSA calculations**

The Prime MMGBSA [63,64] method was employed to assess the relative binding-free energy (ΔG bind) of ligand molecule, along with its individual energy contribution, throughout the 100ns MD simulation. In each step, 100 frames were captured, and the findings are detailed in Table 7 and depicted in Fig. 16. Prime MM-GBSA calculates the energy of optimized free receptors, free ligands, and their complexes. Among the two compounds evaluated, the compound of interest

Mol within the complex 2WV2-mol demonstrated the most favorable binding energy of -63.19 kcal/mol, surpassing the co-ligand present in 2WV2-STD, which exhibited a binding energy of - 10.09 kcal/mol.



**Figure 14.** PCA and FEL plots of the protein Cα atoms dynamics evolution 2WV2-STD complex.



**Figure 15.** PCA and FEL plots of the protein Cα atoms dynamics evolution 2WV2-mol complex.

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**Figure 16** Comparative Analysis of Energy Levels: Binding Energy, Electrostatic Energy, Covalent Binding Energy, Hydrogen Bonding, and Van Der Waals Interactions for the tested complex 2WV2-mol the co-ligand complex 2WV2-STD.

#### **4. Conclusions**

Theoretical methods accurately reproduced X-ray geometrical coordinates and quantum chemical parameters for  $(E)$ -1-[2-(3,4-Dimethylphenyl)diazen-2-ium-1-yl]naphthalen-2 olate compound (EDNO). Hirshfeld surfaces were employed to confirm and measure the occurrence of intermolecular interactions inside the crystal stack. The molecular docking simulation illustrated the manner in which the (EDNO) molecule interacts with specific residues (amino acids) that are crucial for its inhibitory impact. The connection facilitates the movement of the compound into active protein sites with a binding energy of -10.2 kcal/mol. Our investigation into molecular docking reveals that establishing hydrogen bonds with key amino acids (SER296) in the target protein is crucial for stabilizing ligands inside the active site of the 2WV2 Protein and enhancing their inhibitory effectiveness. A molecular dynamics simulation took place on the docked molecule, which demonstrated significant hydrogen bonding with crucial amino acids of the target protein throughout the simulation. The thermodynamic stability of the molecule was assessed through running the MD simulation for 100 ns. The molecule's thermodynamic stability was evaluated by conducting a 100 ns molecular dynamics (MD) simulation. Consequently, in a biological context, they will probably attach to the target protein and

remain stable. The study's findings clearly suggest that the mentioned compound shows promise as an inhibitor of Corticosteroid side-chain-isomerase

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