Research article

Design and *in silico* screening of aryl allyl mercaptan analogs as potential histone deacetylases (HDAC) inhibitors

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ABSTRACT

The Zn$^{2+}$ HDACIs show promising anticancer activity. Allyl mercaptan (AM), a metastabilzed monomeric form of diallyl disulphide (DADS) shows better HDACI activity. The present work screens a dataset of aryl AM derivatives 1(a-g) for potential HDACI action via *in silico* models. DFT calculations predicted the geometrical parameters and frontier orbital calculations suggested better chemical reactivity. Negative chemical potential and NBO hyperconjugative interactions predicted their chemical stability. ADME study confirmed favourable drug likeliness. Molecular docked models suggested the formation of coordinate bond between sulphur of allylmercaptan and Zn$^{2+}$ cofactor of HDAC8. Besides, models also predicted the dominance of hydrophobic interactions. The aryl AM analogs docked perfectly with HDAC3 as well. The glide score and S-Zn distance of compounds 1a, 1f and 1g were found to be better than allylmercaptan. Therefore, the designed aryl AM analogs filtered as better HDACIs. These could be further used for design and synthesis of new improved HDACIs.

1. Introduction

HDAC inhibitors have gained the interest of researchers possibly due to their potency and diverse pharmacological applications. Various structurally diverse HDAC inhibitors possess anti-tumor, anti-histaminic, anti-inflammatory and immune modulatory properties. HDACIs have been known to inhibit induction and proliferation, induce differentiation, and influence a variety of processes such as cell cycle arrest, angiogenesis, and apoptosis of tumor cells in culture and in animal models [1, 2].

The Zn$^{2+}$ dependent HDACIs have been widely studied as anticancer drugs to suppress the general action of catalyzing deacetylase activity of HDACs in presence of Zn$^{2+}$ as essential cofactor. Among the four classes of HDACs classes I, II, and IV are inhibited by Zn$^{2+}$ binding HDACIs. However, class III HDACs is structurally homologous with the yeast Sir2 protein and requires NAD$^+$ as a cofactor instead of Zn$^{2+}$ [3, 4]. In class I, HDACs 1, 2, 3 and 8 are mostly preferred for binding studies as they are localized in the nucleus, most abundant and ubiquitously-expressed [5]. The wide active pocket and a larger surface area of HDAC8 is an interesting feature which distinguishes it from other HDAC enzymes [6]. HDAC8 is ubiquitously expressed in all human tissues and organs as reported in experiments performed on total tissue extracts [7, 8, 9]. Thus HDAC8 is the most acceptable receptor for such HDAC inhibition. HDAC3 has a unique recruitment to SMRT complex where it interacts with conserved DAD [10, 11, 12, 13, 14, 15]. The Ser424 residue in the active site of HDAC3 is primarily phosphorylated. Thus dephosphorylation of Ser424 on HDAC3 inhibits HDAC3 activity possibly due to a conformational change that renders it less active. The three FDA approved HDACIs, vorinostat, belinostat and romidepsin have shown considerable anti-cancer activity [16, 17, 18].

The dietary compounds such as sulforane, genistein, tea polyphenol.catechins, curcumin, diallyl disulphide and revestrol play an important role in regulating key molecular targets like HDAC, DNA methyltransferases etc for the treatment of various diseases [19, 20]. Diallyldisulphide (DADS) a dietary organosulphur compound exhibits excellent HDAC inhibitory activity, anti-microbial, anti-inflammatory, anticancer, anti-oxidant and antihistaminic activity [21, 22, 23, 24].
### Table 1. Aryl allyl mercaptan derivatives with electron withdrawing substituents.

| Compound no. | X   | X'   |
|--------------|-----|-----|
| 1a           | H   | NO₂ |
| 1b           | Cl  | H   |
| 1c           | Cl  | Cl  |
| 1d           | Br  | H   |
| 1e           | Br  | Br  |
| 1f           | H   | F   |
| 1g           | H   | CF₃ |

### Table 2. Designed aryl allyl mercaptan compounds with electron withdrawing substituents.

| Property | Compounds | 1a | 1b | 1c | 1d | 1e | 1f | 1g | AM |
|----------|-----------|----|----|----|----|----|----|----|----|
| Log P    |           | 2.94 | 3.44 | 4.09 | 3.57 | 4.35 | 3.15 | 3.59 | 1.28 |
| TPSA     |           | 45.82 | 38.80 | 38.80 | 38.80 | 38.80 | 38.80 | 38.80 | 38.80 |
| Natoms   |           | 13 | 11 | 12 | 11 | 12 | 11 | 14 | 4 |
| MW       |           | 195.24 | 184.69 | 219.13 | 215.11 | 308.03 | 168.23 | 218.24 | 74.15 |
| nOH      |           | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| nOHNH    |           | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| Nrotb    |           | 3 | 2 | 2 | 2 | 2 | 3 | 1 |    |
| Volume   |           | 169.26 | 159.46 | 172.99 | 163.81 | 181.69 | 150.85 | 174.38 | 74.83 |

### Table 3. Designed aryl allyl mercaptan compounds with electron withdrawing substituents.

| Property | Compounds | 1a | 1b | 1c | 1d | 1e | 1f | 1g | AM |
|----------|-----------|----|----|----|----|----|----|----|----|
| BBB      |           | 1.96 | 1.36 | 1.02 | 1.30 | 0.98 | 1.65 | 1.19 | 1.19 |
| Caco2    |           | 18.56 | 58.14 | 58.48 | 57.75 | 58.51 | 55.51 | 54.24 | 54.78 |
| HIA      |           | 84.18 | 98.11 | 98.05 | 98.03 | 98.27 | 98.46 | 98.36 | 96.86 |
| MDCK     |           | 35.75 | 42.13 | 11.28 | 0.14 | 0.18 | 132.15 | 36.60 | 137.61 |
| PPB      |           | 83.66 | 73.28 | 90.85 | 78.18 | 91.69 | 75.74 | 87.65 | 50.56 |
| SP       |           | -2.21 | -1.02 | -0.99 | -0.93 | -0.80 | -1.20 | -0.77 | -1.93 |

BBB: Blood Brain Barrier, HIA: Human Intestinal Absorption, PPB: Plasma Protein Binding, SP: Skin Permeability.

*Figure 1. Ground state optimized structure of aryl AM 1a at B3LYP/6-311G**(d,p) level.*
DADS has been reported to metabolize to allylmercaptan (AM) within 30 min of administration. The inhibitory HDAC action of AM was reported better compared to its precursors Diallyldisulphide (DADS) and S-allylmercaptocysteine (SAMC) under cell-free conditions [25, 26, 27]. Compounds containing thiol moiety such as allyl alcohol and mercaptoethanol have been found to abolish and decrease HDAC inhibitory activity respectively in contrast to compounds containing sulfhydryl group. Therefore the sulfhydryl group of AM plays a major role in HDAC inhibition. Various compounds prepared using synthetic strategies with -(CH)2-SH group have been reported with strong HDAC inhibitory action majorly due to -S-Zn binding within the active site [28, 29].

Owing to the fact that diallyl disulphide metabolizes to AM within few minutes of administration, in the present study the allyl mercaptan moiety of the pre-synthesized DADS derivatives by Rai et al. [30] have been screened for their pharmacokinetics, drug likeliness and potential HDAC inhibitory action via in silico DFT, ADME and molecular docking studies.

2. Methodology

A preliminary dataset of 7 aryl AM derivatives (Table1) was selected with electron withdrawing substituents. The derivatives were screened for their drug likeliness and non-toxicity via in silico studies. The most reactive positions in these compounds were screened using DFT (Frontier orbital and NBO) calculations. The compounds were further screened for their potential HDAC inhibitory activity via Molecular docking study.

### Table 4. Theoretical optimized geometrical properties of aryl AM 1a at B3LYP/6-311G++(d,p) level.

| Bond length (Å) | Theoretical calculated | Bond angle (°) | Theoretical calculated | Bond dihedral (°) | Theoretical calculated |
|-----------------|------------------------|----------------|------------------------|------------------|------------------------|
| C1-C2           | 1.387                  |                |                        |                  |                        |
| C2-C3           | 1.399                  | C1-C2-C3       | 120.44                 |                  |                        |
| C3-C4           | 1.398                  | C2-C3-C4       | 119.19                 | C1-C2-C3-C4      | -0.05                  |
| C4-C5           | 1.386                  | C3-C4-C5       | 120.71                 | C2-C3-C4-C5      | 0.04                   |
| C5-C6           | 1.400                  | C4-C5-C6       | 119.86                 | C3-C4-C5-C6      | 0.03                   |
| C7-C1           | 1.099                  | C7-C1-C2       | 119.31                 | C7-C1-C2-C3      | -179.96                |
| C8-C2           | 1.096                  | C8-C2-C1       | 119.61                 | C8-C2-C1-C6      | 179.93                 |
| C9-C4           | 1.102                  | C9-C4-C3       | 118.65                 | C9-C4-C3-C2      | -179.82                |
| C10-C5          | 1.099                  | C10-C5-C4      | 119.38                 | C10-C5-C4-C3     | -179.94                |
| C11-C3          | 1.459                  | C11-C3-C2      | 119.61                 | C11-C3-C2-C1     | 179.98                 |
| C12-C11         | 1.339                  | C12-C11-C3     | 122.41                 | C12-C11-C3-C2    | -179.07                |
| C13-C11         | 1.098                  | C13-C11-C3     | 116.34                 | C13-C11-C3-C2    | 0.95                   |
| C14-C12         | 1.104                  | C14-C12-C11    | 120.08                 | C14-C12-C11-C3   | -0.58                  |
| C15-C12         | 1.479                  | C15-C12-C11    | 122.99                 | C15-C12-C11-C3   | 179.14                 |
| C16-C15         | 1.105                  | C16-C15-C12    | 111.93                 | C16-C15-C12-C1   | 5.79                   |
| C17-C15         | 1.106                  | C17-C15-C12    | 108.90                 | C17-C15-C12-C1   | -111.29                |
| C18-C15         | 1.827                  | C18-C15-C12    | 108.06                 | C18-C15-C12-C1   | 128.09                 |
| C19-H19         | 1.307                  | C19-H18-C15    | 99.02                  | C19-H18-C15-C1   | 174.53                 |
| N20-C6          | 1.495                  | N20-C6-C5      | 120.15                 | N20-C6-C5-C4     | 179.92                 |
| O21-N20         | 1.215                  | O21-N20-C6     | 119.48                 | O21-N20-C6-C5    | -0.78                  |
| O22-N20         | 1.215                  | O22-N20-C6     | 119.48                 | O22-N20-C6-C5    | 179.29                 |

![Figure 2. Optimized ground state geometries of conformers of aryl AM 1a at B3LYP/6-311G++(d,p) level with their relative Gibbs free energies.](image)

### Table 5. Representative Relative Gibbs free energies of 1(b-g) conformers indicating the most stable conformation.

| CONFORMERS | RELATIVE GIBBS FREE ENERGIES (HARTREES) |
|------------|----------------------------------------|
|            | 1b          | 1c          | 1d          | 1e          | 1f          | 1g          |
| 1           | 0           | 0           | 0           | 0           | 0           | 0           |
| 2           | 0.0002      | 18.7145     | 2.7603      | 0.0001      | 0.0001      | 0.0005      |
| 3           | 0.0003      | 18.7154     | 2.7614      | 0.0002      | 0.0002      | 0.0009      |
| 4           | 0.0034      | 18.7155     | 2.7615      | 0.0010      | 0.0002      | 0.0027      |
| 5           | 0.0035      | 18.7180     | 36.9312     | 0.0011      | 0.0009      | 0.3430      |
| 6           | 0.0048      | 18.7181     | 36.9374     | 0.0014      | 0.0026      | 4.0966      |

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Figure 3. Optimized ground state geometries of most stable conformers of i) 1b, ii) 1c, iii) 1d, iv) 1e, v) 1f and vi) 1g at B3LYP/6-311G++(d,p) level with their relative Gibbs free energies (kcal/mol).

Table 6. The electronic properties of compounds calculated at DFT-B3LYP/6-311G++(d,p) level.

| Compound | $E_{\text{HOMO}}$(KJ/mol) | $E_{\text{LUMO}}$(KJ/mol) | $E_{\text{L}}-E_{\text{H}}$ (KJ/mol) | $\eta$(KJ/mol) | $\mu$(KJ/mol) |
|----------|----------------|----------------|-------------------------------|----------------|--------------|
| 1a       | -0.1283        | 0.0167         | 0.145                         | 0.0725         | -0.1116      |
| 1b       | -0.2452        | 0.2071         | 0.452                         | 0.4523         | -0.0381      |
| 1c       | -0.2515        | 0.1901         | 0.441                         | 0.4416         | -0.0614      |
| 1d       | -0.1211        | 0.0487         | 0.169                         | 0.1698         | -0.0724      |
| 1e       | -0.1278        | 0.0513         | 0.171                         | 0.1734         | -0.0803      |
| 1f       | -0.1165        | 0.0599         | 0.176                         | 0.1764         | -0.0566      |
| 1g       | -0.1216        | 0.0486         | 0.170                         | 0.1702         | -0.0730      |
| AM       | -0.2642        | 0.0276         | 0.236                         | 0.1459         | -0.1183      |
2.1. Calculation of druglikeliness properties

The molecular properties of compounds 1a-g were calculated using the Molinspiration cheminformatics software (www.molinspiration.com). The major druglikeliness parameters calculated include number of atoms, molecular weight, partition coefficient (LogP), hydrogen bond acceptors and donors, topological surface area (TPSA), and Lipinski's rule violations [31].

2.2. Pharmacokinetic and toxicity parameters

PreADMET online server 2.0 version [32] was used to calculate adsorption, distribution, metabolism and excretion (ADME) parameters. The properties calculated were plasma protein binding (PPB), human intestinal absorption (HIA), logKp (degree of skin permeation), Caco2 cell lines and MDCK cell lines barrier penetration. These predicted the oral absorption of the compounds [33].

2.3. Geometry optimization using density functional theory (DFT)

DFT study illustrates the electronic structural properties of the molecules. The DFT calculations were done using Gauss View 5.0 [34] molecular visualization program and Gaussian 09 program [35]. The molecular structures of the aryl AM compounds in the ground state were optimized at B3LYP/6-311G++(d,p) level of computation [36].

The optimized ground state geometry of the most stable conformer was used to predict different structural parameters like bond lengths, bond angles, and dihedral angles.

Figure 4. Frontier orbital diagram of 1a using DFT-B3LYP/6-311G++(d,p) method with frontier orbital gap value of 0.145 kcal/mol.

Figure 5. Frontier orbital diagram a) 1b with frontier orbital gap of 0.452, b) 1c with frontier orbital gap value of 0.441, c) 1d with frontier orbital gap of 0.169, d) 1e with frontier orbital gap of 0.171, e) 1f with frontier orbital gap of 0.176, f) 1g with frontier orbital gap of 0.170.
2.4. Frontier molecular orbital calculations

The electronic properties like frontier molecular orbital, HOMO (highest occupied molecular orbital) and LUMO (lowest unoccupied molecular orbital) energies are important for defining reactivity of a chemical species \(^{37}\). The global reactivity descriptors, chemical potential (\(\mu\)) and hardness (\(\eta\)) described by Parr and Pearson were calculated using equations \(^{38}\):
\[ \mu = -\frac{I + A}{2} \]  

\[ n = \frac{(I - A)}{2} \] respectively

where I is the ionization potential \((-E_{\text{HOMO}})\) and A is the electron affinity \((-E_{\text{LUMO}})\) as correlated using the Koopman’s theorem \([39, 40]\).

2.5. Natural bond order analysis (NBO)

Reed and Weinhold performed the natural bond orbital (NBO) calculations \([41]\) which depict the second order interactions between the filled Lewis orbital to empty non-Lewis orbital of a sub-system. These interactions determine the intermolecular delocalization or hyperconjugation within a system. Considering the title compound the second
order perturbation theory analysis for Fock matrix in NBO was carried out to evaluate donor acceptor interactions. They result in a loss of occupancy from localized NBO of Lewis structure to empty non-Lewis orbital [33]. The stabilization energy $E(2)$, for each donor (i) and acceptor (j) is given by the formula:

$$E(2) = \Delta E_{ij} = (q_i \times F(i,j))^2/(e_j - e_i)$$

where $q_i$ is the donor orbital occupancy, $e_i$ and $e_j$ are diagonal elements and $F(i,j)$ is the off diagonal NBO Fock matrix element. These calculations allow us to analyse the probable charge-transfers and the intermolecular bond paths.

### 2.6. Molecular docking

The three dimensional target receptor HDAC8 (PDB ID: 1T67) and HDAC3 (PDB ID: 4A69) structures were retrieved from the Protein Data Bank (www.rcsb.org). The Schrödinger suite was used to refine protein structure via protein preparation wizard. Prime (Prime v2.0, 2012) was used to add missing side chains and hydrogens to the raw protein structure. The crystallized water molecules in the active site present beyond 5 Å were removed using an all atom force field. In order to remove potential steric clashes minimizations on the protein structure were done until the RMSD value (root mean square deviation) reached 0.3 Å for non-hydrogen atoms using OPLS-2005 force field.

The lead targets were prepared using Ligprep function of the Schrödinger suite in which the energy minimization was performed using OPLS-2005 force field. A grid was generated around the receptor using Grid generation program and Glide Docking program respectively. The XP (Xtra Precision) mode of docking was selected in the present case [42, 43].

### 3. Results and discussion

#### 3.1. Drug likeness properties

The compounds 1a-g obeyed the Lipinski’s rule of five (i.e. MW < 500, Log P < 5, number of H bond donors up to 5 and H bond acceptors up to 10) [31](Table 2). Thus these designed derivatives depicted drug like properties.
3.2. ADME parameters

The ADME parameters for the compounds 1a-g have been given in Table 3. The human intestinal (HIA) values indicate good oral absorption as the values were above 90%. The negative Kp values indicate poor skin permeation which results in good oral absorption. The PPB values except 1e were less than 90%. The Gaco 2 values were within range of 4–70, indicative of moderate permeability. MDCK values were greater than 25 except for compounds 1c-1e, indicative of good absorption. Apart from these factors various other parameters need to be considered. The BBB values are less than two which indicates non-neurotoxicity of these compounds. Thus these screened compounds depicted good pharmacokinetics and bioavailability.

3.3. Geometry optimization

The ground state optimized geometry of aryl AM 1a with NH2 substituents is given in Figure 1. The structural parameters like bond lengths, bond angles and dihedral angles for the title molecule are given in Table 4. The C-C bond lengths of the phenyl ring and allyl moiety are shorter than other C-C bonds verifying the double bond character of the ring and allyl C-C bonds. The average C-C bond length is 1.39 Å which is almost identical to that of diamond, and the C-H distances lie in the range 1.096–1.106 Å. The S-H bond length in aryl AM 1a is calculated to be 1.307 Å. The preferred bond angle value between successive carbon chain bonds in phenyl ring system is around 119°. Also, the dihedral angles should be nearly 0° or 180° conferring the planarity of the compound. The major dihedral angles which depict the orientation of AM with respect to the phenyl ring via S atom, S18-C15-C12-C11, H16-C15-C12-C11, H19-S18-C15-C12 with the corresponding values 128.09°, 5.79°, 174.53° respectively, revealed the deviation of planarity by sulphur atom that leads to different conformations of the compound. The different conformations particularly arise due to the non-planarity of sulphur atom as the mercaptan bond can undergo free rotation.

The conformational behavior of the molecule provides useful information regarding drug actions. A detailed conformational theoretical analysis has been carried out in order to understand the conformational properties of the molecules. The presence of double bond on the AM results in the generation of cis and trans isomers. The trans isomer was predicted to be more stable than the cis form as predicted by the B3LYP/6-311G results in the generation of cis and trans isomers. The trans isomer was predicted to be more stable than the cis form as predicted by the B3LYP/6-311G calculations.

The relative stability of sub-conformations is not affected by different orientations of the phenyl rings in the main configuration (maximum: 0.25 kcal/mol) [44]. Thus the order of decreasing stability of conformers in compound 1a is i (0.0 kcal/mol) >ii (0.0025 kcal/mol) >iii (11.5137 kcal/mol).

The first conformation is the most stable form as the mercaptansulphur is eclipsed with hydrogens and methyl. For propionaldehyde the carbonyl oxygen is eclipsed with hydrogens and methyl and it has been shown by 1H NMR spectra that such types are the preferred conformers in order to minimize the steric effect [45]. The conformational free energy of compound 1a is approximately 11.5 kcal/mol in favor of the conformer i in which the mercaptansulphur is nearly eclipsed with methyl group. Similarly, for the compounds 1d, 1e, 1f, and 1g the lowest energy conformer as determined from the lowest Gibbs free energy values (Table 5) was found to be the one in which sulphur atom was eclipsing with methyl group (Figure 3).

3.4. Frontier molecular orbitals

The calculated electronic properties are listed in Table 6. The compound 1a has the lowest frontier orbital gap pertaining to its higher chemical reactivity (Figure 4). The negative chemical potential values for compounds 1(a-g) indicate non-spontaneous decomposition which is a prerequisite of this study. The global hardness (n) depicts the resistance towards electron cloud deformation under perturbation observed in a chemical process [47,49].

Similarly, on the basis of frontier orbital gap, chemical potential and global hardness the compounds 1d, 1e, 1f and 1g had lesser frontier orbital gap and negative chemical potential compared to compounds 1b and 1c implying that compounds 1b and 1c are less reactive as compared to other compounds. For the compounds 1d, 1e, 1f and 1g the HOMO of sigma nature is localized over the sulphur atom while in LUMO the electron density is delocalized over the phenyl ring implying electron density transfer from the sulphur atom to the phenyl ring (Figure 5). This study provides an insight into the binding interactions which could occur majorly via the sulphur atom.

3.5. NBO analysis

The second order perturbation stabilization energy values E(2) (given in Table 7) revealed significant interactions between Lewis and non-Lewis NBO orbitals for the compound 1a. The transfer of electron density from oxygen atom lone pair (O22) in antibonding orbitals p(N20-O22) resulted in strong interaction with high stabilization energy 197.72 kcal/mol. Other important interactions include overlap of bonding p(C3-C4) with antibonding orbitals p(C11-C12), p(C5-C6) and p(C1-C2) with corresponding stabilization energies 38.85, 25 and 20.51 kcal/mol respectively, bonding p(C5-C6) with antibonding orbitals p(C1-C2) and p(C3-C4) with corresponding stabilization energies 21.67 and 20.41 kcal/mol respectively, and bonding p(C1-C2) with antibonding orbitals p(C3-C4) and p(C5-C6) with stabilization energies 20.32 and 21.63 respectively. These interactions result in pronounced decrease in C3/C4, C5/C6 and C1/C2 orbital’s occupancy (0.36548, 1.6525 and 1.67642 respectively). It also indicates a possibility of hyperconjugation within the phenyl ring and between the ring and allyl carbons. The lone pair interactions include overlap of lone pair (LP2) of oxygen atom (O21) to antibonding orbitals p(N20-O22) with stabilization energy 15.94 kcal/mol. Similarly, the interaction of lone pair (LP2) of oxygen atom (O21) with p(C5-C6) has stabilization energy of 14.39 kcal/mol.

The NBO analysis predicts the stability of molecules arising from hyperconjugative interactions and charge delocalization. The results clearly show that electron density in anti-bonding orbitals and second order delocalization energies confirm the occurrence of inter-molecular charge transfer within the molecules [50].

These interactions indicate that the addition of electron withdrawing substituent to aromatic ring results in further molecular stabilization of the compound. Similarly, hyperconjugative interactions within the phenyl ring and between the ring and allyl carbons were seen for other compounds along with other interactions resulting in stabilization of the compounds 1(b-g).

3.6. Molecular docking

The overlapping of docked compounds 1(a-g) with AM based 3D HDAC8 and HDAC3 binding model is depicted in Figure 6.

The conserved interacting amino acid residues were Tyr 306, His 180, Asp 267, Phe 152, Asp 178, His 142 and His 143. Docking investigations revealed the formation of metal coordinate bond between S and Zn at a distance of 2.348 Å, 2.358 Å and 2.362 Å for compounds 1a, 1f and 1g.
respectively. The value for reference AM was 2.56 Å which indicates better binding affinity of compounds 1a, 1f and 1g (Figure 7).

The glide score, gibbs energy values and Zn-S bond distances for the compounds are listed in Table 8. The hydrophobic interactions were seen with residues Trp 141, Tyr 306, Met 274, Phe 208 and Phe 152 for all the compounds 1(a-g) (Figure 8).

Polar interactions were observed with residues His 180 and His 142. There is preponderance of hydrophobic interactions on the linker chain which indicates that binding majorly takes place in the hydrophobic pocket of the receptor (Figure 9). It is well known through various published works and our own experiences that both hydrophobic as well as hydrogen binding interactions play pivotal role in complexation of ligands with proteins [45, 46, 47]. It has been reported that the benzyl mercaptan molecule bulkier than AM inhibited HDAC8 better than Hela nuclear extracts primarily containing HDAC3 indicating specific HDAC8 inhibition [29]. Watson et al. [48] predicted that the binding site of ligand in HDAC3 is far away from Zn atom a possible reason for selective binding to HDAC8. In the present study the designed aryl AM compounds bearing electron withdrawing groups 1(a-g) were docked with HDAC3 also. Figure 6 (II) depicts docked aryl AM compounds 1(a-g) with HDAC3.

The docking scores and Gibbs energy values were found to be good and comparable to the values as obtained for HDAC8 (Table 8). The non-covalent interactions like hydrophobic and Vander Waal interactions dominated the binding site. The results clearly state that these new designed aryl AM compounds bearing electron withdrawing substituents show general HDACI action and could be modified accordingly to treat multiple diseases.

4. Conclusion

A comprehensive in silico investigation of designed aryl AM compounds 1(a-g) bearing electron withdrawing substituents was done. The most stable conformation calculated at DFT B3LYP/6-311G++(d,p) level of computation had methyl carbon eclipsed with mercaptansulphur. The structural parameters determined the planarity of the compounds except for sulphur atom which lies slightly above the plane. The electronic properties suggested high chemical reactivity. The NBO results indicated hyperconjunction within the phenyl ring and the ring with allyl carbons. The drug likeness was confirmed viaADME. Docking data suggested the importance of hydrophobic interactions and metal coordination between the ligands 1(a-g) and receptor. It supports the fact that substitution of aryl AM moiety with electron withdrawing substituents makes aryl AM better general HDACs. Thus these potential HDACs candidates could further help in design, development and screening of new pharmaca-phores which could be modified for diversified pharmacological actions.

Declarations

Author contribution statement

Himanshu Ojha, Malika Pathak, Paban K Agrawala: Conceived and designed the experiments; Analyzed and interpreted the data; Wrote the paper.

Sugandha Singhal: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.

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Competing interest statement

The authors declare no conflict of interest.

Additional information

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