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Synthesis, in vitro biological evaluation and in silico studies of novel pyrrolidine derived thiosemicarbazones as dihydrofolate reductase inhibitors†

Hina Aftab,^a Saeed Ullah,^b Ajmal Khan,^{bc} Mariya al-Rashida,^d Talha Islam,^d Abdulrahman Alshammari,^e Norah A. Albekairi,^e Parham Taslimi, ^b Ahmed Al-Harrasi,^{*b} Zahid Shafiq ^b *a and Saeed Alghamdi^g

Dihydrofolate reductase (DHFR) is a crucial enzyme involved in folate metabolism and serves as a prime target for anticancer and antimicrobial therapies. In this study, a series of 4-pyrrolidine-based thiosemicarbazones were synthesized and evaluated for their DHFR inhibitory activity. The synthesis involved a multistep procedure starting from readily available starting materials, leading to the formation of diverse thiosemicarbazone 5(a-r) derivatives. These compounds were then subjected to *in vitro* assays to evaluate their inhibitory potential against DHFR enzyme. The synthesized compounds 5(a-r) exhibited potent inhibition with IC₅₀ values in the range of $12.37 \pm 0.48~\mu M$ to $54.10 \pm 0.72~\mu M$. Among all the derivatives 5d displayed highest inhibitory activity. Furthermore, molecular docking and ADME studies were performed to understand the binding interactions between the synthesized compounds and the active site of DHFR. The *in vitro* and *in silico* data were correlated to identify compounds with promising inhibitory activity and favorable binding modes. This comprehensive study provides insights into the structure–activity relationships of 4-pyrrolidine-based thiosemicarbazones as DHFR inhibitors, offering potential candidates for further optimization towards the development of novel therapeutic agents.

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1. Introduction

Folate metabolism has emerged as a promising and significant target for the creation of anti-cancer, anti-bacterial, and anti-parasitic therapeutic medicines in the last several years. Dihydrofolate reductase (DHFR) plays a crucial role in reducing 7,8-dihydrofolate (DHF) to 5,6,7,8-tetrahydrofolate (THF) in cooperation with NADPH. The production of thymidine (a precursor for DNA replication), purine nucleotides, glycine, methionine,

serine, and N-formyl-methionyl tRNA all rely on this chemical as a precursor to the cofactors needed for these reactions.3 Cell development is restricted due to partial depletion of intracellular reduced folates caused by DHFR inhibition.⁴ Antibacterial DHFR inhibitors work by preventing the production of DNA, RNA, and proteins, which in turn stops the proliferation of cells. Consequently, DHFR becomes a prime target for anticancer and antibacterial drugs. A decrease in DHFR enzyme activity lowers the intracellular THF pool, which impacts the concentration of folate coenzymes and, by extension, the production of purines and pyrimidines.5,6 Since methyl-THF is required for homocysteine remethylation to produce methionine, an essential component of S-adenosylmethionine (SAM) required for the majority of biological methylation reactions, this may also affect homocysteine levels and methylation processes. Higher eukaryotic cells must rely on exogenous folic acid because, unlike prokaryotes and lower eukaryotic cells, they are unable to produce folates de novo. So, to acquire medications to treat infectious disorders caused by protozoan parasites and bacteria, the folic acid production route is an easy target.7 Several DHFR inhibitors, as separate entities, have found clinical utility as antitumor agents.8-10 One of the most potent DHFR inhibitor methotrexate (MTX) is one the first chemotherapeutic agent discovered and is still a support in single agent and combination cancer chemotherapy.11

[&]quot;Institute of Chemical Sciences, Bahauddin Zakariya University, Multan-60800, Pakistan. E-mail: zahidshafiq@bzu.edu.pk

^bNatural and Medical Sciences Research Centre, University of Nizwa, P.O. Box 33, PC 616. Birkat Al Mauz. Nizwa. Sultanate of Oman. E-mail: aharrasi@unizwa.edu.om

Department of Chemical and Biological Engineering, College of Engineering, Korea University, 145 Anam-ro, Seongbuk-gu, Seoul 02841, Republic of Korea

^dDepartment of Chemistry, Forman Christian College (A Chartered University) Lahore, Pakistan

Department of Pharmacology and Toxicology, College of Pharmacy, King Saud University, Post bezBox 2455, Riyadh, 11451, Saudi Arabia

Department of Biotechnology, Faculty of Science, Bartin University, 74100 Bartin, Turkey

^{*}Department of Pharmacy, Riyadh Security Forces Hospital, Ministry of Interior, Kingdom of Saudi Arabia

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One of the nitrogen heterocycles utilized extensively by medicinal chemists to derive molecules for the treatment of human diseases is the five-membered pyrrolidine ring. This saturated scaffold is of great interest for three reasons: (1) the ability to efficiently explore the pharmacophore space via sp³hybridization; (2) the impact on the molecule's stereochemistry; and (3) the increased three-dimensional (3D) coverage caused by the ring's non-planarity; a phenomenon known as "pseudo rotation".12 Additionally, the nitrogen atom of pyrrolidine ring has the capability to engage in hydrogen bonding interactions with amino acids residues present within active site of DHFR,

thereby enhancing the compound's binding affinity. Its size and

shape are well-suited for fitting into the active site pocket of DHFR. This facilitates favorable van der Waals interactions with

nearby amino acids residues, which in turn enhances the

stability of the enzyme-inhibitor complex.13 The incorporation

of heteroatomic fragments into the molecules is a thoughtful

decision, as they serve as valuable tools for adjusting physico-

chemical parameters and achieving optimal ADME outcomes for potential drug candidates.14,15 Thiosemicarbazones are a fascinating class of compounds. Numerous pharmacological and biological features have been discovered including their efficacy as anti-urease, 16,17 chemosensing,18 antibacterial,19 antifungal,20 anticancer,21 antifolate, 22,23 antidiabetic activity 24 and antimicrobial activity. 25 On the other hand, thiosemicarbazones also belong to one of the main families of DHFR (dihydrofolate reductase) inhibitors^{26,27} due to the formation of stable complexes with zinc ions present in the active side of DHFR.28,29 As thiosemicarbazones are known as chelate metal ions, this chelation helps in binding and inhibiting the enzyme. Furthermore, thiosemicarbazones possess a thioamide (-CSNH-) functional group, which may also participate in hydrogen bonding interactions with specific amino acids residues in the DHFR active site, these interactions helps to stabilize the inhibitor in the enzyme's binding pocket, enhancing its potency.28,30 The structural characteristics and redox activities of thiosemicarbazones contribute to their efficacy as anticancer agents, making them promising candidates

Molecular hybridization involves merging different molecules to create new compounds. In our study, we synthesized novel 4-pyrrolidine based thiosemicarbazones by combining 4pyrrolidinyl-benzaldehyde with thiosemicarbazide derivatives. These compounds were then evaluated for their effectiveness against the enzyme DHFR. Additionally, we conducted molecular docking, ADMET studies, and explored structure-activity relationships to understand the properties and potential of these compounds as DHFR inhibitors.

Structures of some pyrrolidine and thiosemicarbazones based dihydrofolate reductase (DHFR) inhibitors reported in literature are shown in (Fig. 1).32-35

Results and discussion 2.

for further exploration as DHFR inhibitors.31

2.1 Chemistry

Pyrrolidine and thiosemicarbazone moieties have a wide range of biological and pharmacological effects. Therefore, it is reasonable to integrate these two components into a unified molecular framework to create more potent biologically active compounds, such as those depicted in Fig. 1. The desired compounds were synthesized by modifying pyrrolidine moiety through the introduction of 4-fluorobenzaldehyde. This modification primarily involves replacing the NH group of pyrrolidines with 4-fluorobenzaldehyde instead of the hydrogen atom at the N-H position of pyrrolidine, and forming thiosemicarbazones by reacting with thiosemicarbazides, thus enhancing their biological activity. The targeted compounds 5(a-r) were prepared by the substitution at N-H position which was done by the reaction of pyrrolidine (1) with 4-fluorobenzaldehyde (2) using anhydrous K2CO3 and DMF at 90 °C for 12 h, giving 4-(pyrrolidin-1-yl)benzaldehyde (3). The synthesis of thiosemicarbazones was achieved by reaction of compound (3) with thiosemicarbazides 4(a-r). The synthetic route in synthesis of 4-pyrrolidine based thiosemicarbazones 5(a-r) was displayed in Scheme 1.

The structure of synthesized compounds was confirmed using various advanced spectroscopic techniques. In the ¹H-NMR spectra of the pyrrolidine-based thiosemicarbazones, a prominent single peak appeared within the chemical shift range of δ 11.9–10.1 ppm, revealing of NH-protons presence. Furthermore, the protons of N=CH were prominent between δ 7.95 and 9.13 ppm. Notably, C-H protons in the benzene moiety exhibited distinct signals within the range of δ 7.0-8.0 ppm, while those in the pyrrolidine group were identified within δ 1.0–5.0 ppm. Furthermore, negative signals observed in ¹³C-NMR spectra suggested the presence of CH₂ groups in the compounds. Other characteristic signals were detected at δ 23.5, 33.9, 175.8 ppm, corresponding to CH3 and CH, as well as C=S functionalities. In order to determine the purity of compounds, HPLC analysis was carried out using $CH_3CN: H_2O = 80:20$ eluent system with 263 nm wavelength. All the compounds exhibited great than 95% purity.

2.2 Pharmacology

2.2.1 Biological activity. The inhibitory activity of synthesized compounds was evaluated against the dihydrofolate reductase enzyme, utilizing methotrexate as a positive control with an IC₅₀ value of 0.086 \pm 0.07 μ M. The inhibitory activity data, represented as IC₅₀ values, is presented in Table 1. The results indicated that the target compounds displayed IC50 in the range of 12.37 \pm 0.48 μM to 54.10 \pm 0.72 μM . Among all derivatives, 5d and 5l exhibited highest inhibition activity with IC_{50} value 12.37 \pm 0.48 and 12.38 \pm 0.25 $\mu M.$ Among the compounds, 5d, 5e, 5f, 5g, 5l and 5m exhibited good inhibitory potential with IC₅₀ values in the range of 12.38 \pm 0.25 μM to $21.18 \pm 0.44 \,\mu\text{M}$. Derivative 5l exhibited the highest inhibitory activity against dihydrofolate reductase enzyme with IC50 = $12.38 \pm 0.25~\mu\text{M},$ followed by 5d with $IC_{50} = 12.37 \pm 0.48~\mu\text{M},$ 5e $(IC_{50} = 15.30 \pm 0.26 \mu M)$, $5f(IC_{50} = 21.18 \pm 0.44 \mu M)$, $5g(IC_{50} =$ $14.37 \pm 0.29 \ \mu M)$ and $5 \mathbf{m}$ (IC₅₀ = $16.27 \pm 0.26 \ \mu M$). Compound 5a, 5h, 5i, and 5n showed moderate inhibition potency compared to the standard with IC_{50} ranging from 22.06 \pm 0.37 μ M to 29.11 \pm 0.38 μ M. Compounds 5b, 5c, 5j, 5k, 5o, 5p, 5q and

Fig. 1 Reported pyrrolidine and thiosemicarbazone based DHFR Inhibitors.

5r showed moderate to least inhibition activity compared to methotrexate with IC $_{50}$ value as follows.

5b 33.10 ± 0.71

5c 36.20 ± 0.69

50 45.28 ± 0.63

5p 48.15 ± 0.60

structure-activity link. The inhibitory potential of thiosemicarbazone is explored by changing the R group on the

5j 54.10 ± 0.72

 $5k 40.62 \pm 0.59$

5q 35.26 ± 0.59

5r 42.16 ± 0.63

Compound 5j was the least potent member of the series with IC $_{50}$ value of 54.10 \pm 0.72 $\mu M.$

2.3 Structure-activity relationship

Thiosemicarbazones exhibit the remarkable ability to hinder the activity of the DHFR enzyme, thereby disrupting folate metabolism pathways crucial for DNA synthesis and cellular proliferation. This exceptional characteristic renders thiosemicarbazones promising candidates for anti-cancer and anti-microbial therapies. A novel series of thiosemicarbazones, based on 4-pyrrolidinyl-benzaldehyde, has been synthesized and assessed for their efficacy against the dihydrofolate reductase enzyme. The $\rm IC_{50}$ values and percentage inhibition of these synthesized compounds were determined and is presented in Table 1. Thiosemicarbazone and 4-pyrrolidinylbeznaldehyde functioned as significant structural frameworks, and the phenyl or benzyl group bonded to the $\rm N^4$ nitrogen of thiosemicarbazone was crucial in establishing an indisputable

thiosemicarbazides moiety. The R group is varied with aromatic, non-aromatic and aliphatic groups. The compounds 5f and 5p with phenyl and benzyl rings respectively, by comparison, phenyl derivative has more inhibition potential than benzyl derivative. In case of compound 51 and 5d with phenethyl and 4-methoxyphenyl moieties showed maximum inhibition potencies. Compounds 5j and 5o with para and metanitro group on the phenyl ring, among them meta-substitution is slightly active (50, IC₅₀ = 45.28 \pm 0.63 μ M) as compared to para substitution (5j, IC₅₀ = 54.10 \pm 0.72 μ M). Compounds 5d and 5h containing methoxy substitution at the para and meta positions of the phenyl ring respectively, 5d showed much better inhibitory potency with IC₅₀ value 12.37 \pm 0.48 μ M of than **5h** with IC₅₀ value of 23.11 \pm 0.35 μ M. Electron donating substituents, like alkyl groups, have the ability to increase the electron density around the benzene ring through resonance effects. This amplified electron density raises stronger interactions between the inhibitor molecule and the enzymes active

	R	Yield	Time,		R	Yield	Time,
		(%)	temperature			(%)	temperature
5a	Cyclohexyl	75	5 h, 70 °C	5j	4-Nitrophenyl	85	4 h, 80 °C
5b	4-Fluorophenyl	81	7 h, 80 °C	5k	4-Ethylmorpholine	89	5 h, 70 °C
5c	2,3-Dichlorophenyl	91	8 h, 80 °C	51	β-Phenethyl	91	5 h, 70 °C
5d	4-Methoxyphenyl	87	5 h, 70 °C	5m	4-Bromophenyl	91	7 h, 80 °C
5e	2,6-Dimethylphenyl	85	8 h, 80 °C	5n	3-Bromophenyl	85	7 h, 80 °C
5f	Phenyl	83	5 h, 70 °C	50	3-Nitrophenyl	85	5 h, 80 °C
5g	4-Methylphenyl	81	5 h, 70 °C	5p	Benzyl	85	5 h, 70 °C
5h	3-Methoxyphenyl	95	5 h, 70 °C	5q	2-Fluorophenyl	87	8 h, 80 °C
5i	4-Chlorobenzyl	79	7 h, 80 °C	5r	4-Methylbenzyl	87	5 h, 70 °C

3 = 5 mmol, 4(a-r) = 5 mmol (1:1)

Scheme 1 Synthetic route for the preparation of thiosemicarbazones.

site, potentially leading to more potent inhibition. Moreover, the positive inductive effect (+I) exerted by alkyl groups on the benzene rings further intensifies its electron density. This phenomenon enhances the inhibitor's capacity to engage with the enzyme's active site. Therefore, with increase in alkyl substitution inhibition potential is increased as we see in compound 5e when compared with 5g and 5r. In case of methyl substitutions at para position of phenyl and benzyl rings of compound 5g and 5r with $IC_{50}=14.37\pm0.29~\mu M$ and $IC_{50}=42.16\pm0.63~\mu M$ respectively, 5g is more potent as compared to 5r. The compound 5e with dimethyl substitution ($IC_{50}=15.30\pm0.26~\mu M$) slightly increases inhibition potency than one methyl substitution. In case of halogens, with the increase in number of substitutions, potency is somewhat decreased, possibly due to electron-withdrawing effect. This effect makes

any carbon center even more electron-deficient than before also causes negative inductive effect (–I). Their activity order is Br > Cl > F. Compounds **5m** and **5n** with bromo substitution at para and meta positions of phenyl ring respectively, among them **5m** (IC $_{50} = 16.27 \pm 0.26 \,\mu\text{M}$) is more potent than **5n** (IC $_{50} = 22.06 \pm 0.37 \,\mu\text{M}$). Compounds **5c**, and **5i** with 2,3-dichlorophenyl and 4-chlorobenzyl exhibited moderate inhibition with IC $_{50} = 36.20 \pm 0.69$, IC $_{50} = 27.46 \pm 0.50$ respectively. While in case of cyclohexyl containing compound **5a** moderate activity against enzyme was observed (IC $_{50} = 29.11 \pm 0.38 \,\mu\text{M}$).

The study reveals a clear correlation between the chemical structure of the compounds and their inhibitory potency, as showed in Fig. 2. Notably, compounds featuring para substitution of the phenyl ring exhibit heightened potency in the series. Among these, compounds **5d** and **5l**, distinguished by 4-

5d

5f

5h

Table 1 $\,$ IC₅₀ values of the synthesized compounds against dihydrofolate reductase

Compounds R		% inhibition (0.5 mM)	$IC_{50} \pm \mu M$ (SEM)
5a	\leftarrow	83.58	29.11 ± 0.38
5b	F	80.64	33.10 ± 0.71
5c	CI	81.29	36.20 ± 0.69

OCH₃
91.14
12.37
$$\pm$$
 0.48

5e
$$90.25$$
 15.30 ± 0.26

ÇH₃

$$86.20 21.18 \pm 0.44$$

$$\mathsf{5g} \qquad \qquad \mathsf{EH}_3 \\ 89.53 \qquad \qquad \mathsf{14.37} \pm 0.29$$

OCH₃
$$86.71$$
 23.11 ± 0.35

5i
$$83.90$$
 27.46 ± 0.50

Table 1 (Contd.)

	\bigcup		
Compound	s R	% inhibition (0.5 mM)	$IC_{50} \pm \mu M$ (SEM)
5j	NO ₂	74.10	54.10 ± 0.72
5k	√ N O	76.90	40.62 ± 0.59
5 l		91.46	12.38 ± 0.25
5 m	Br	90.77	16.27 ± 0.26
5n	Br	88.20	22.06 ± 0.37
50	NO ₂	75.62	45.28 ± 0.63
5 p		74.60	48.15 ± 0.60
5q	F	77.26	35.26 ± 0.59
5r	CH	3 79.83	42.16 ± 0.63
Standard		Methotrexate	$\textbf{0.086} \pm \textbf{0.07}$

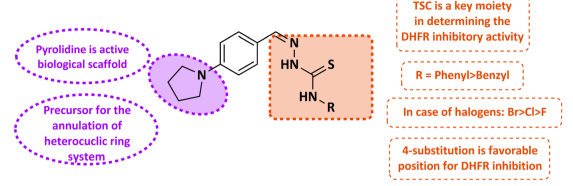


Fig. 2 Structure activity relationship of the synthesized thiosemicarbazones.

methoxyphenyl and phenethyl substitution, emerge as most potent in the series. Conversely, compound 5j, with 4-nitro phenyl substitution, displays diminished activity. It becomes evident that the introduction of electron-withdrawing substituents on the phenyl ring leads to decreased inhibitory potential, whereas electron-donating substitutions enhance activity. This highlights the crucial role of understanding the electronic characteristics of substituents in modulating biological activity, thereby facilitating the rational design of more effective compounds.

2.4 Molecular docking studies

The crystal structure of human dihydrofolate reductase (DHFR) in complex with the NADP and inhibitor SRI-9439 was downloaded from the PDB (PDB id: 1kms, 1.09 Å). To validate the docking protocol, the co-crystallized inhibitor was re-docked, the docking method was able to successfully reproduce the experimentally observed conformation for the ligand. The NADPH co-factor was retained for the docking studies in accordance with previous studies.³⁶ All compounds (5a–r) were docked and were found to bind in the same area of the active site as the co-crystallized inhibitor (Fig. 3). For reference, molecular docking of methotrexate was also carried out to map

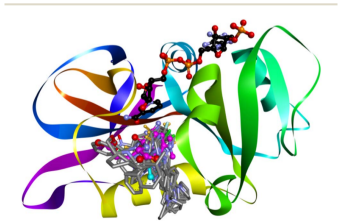


Fig. 3 Overlap of docked conformations of hDHFR inhibitors 5a-r, NADPH is shown in black, the standard inhibitor drug methotrexate is shown in cyan color and the co-crystallized inhibitor is shown in blue.

out important interactions with the binding site amino acids. Details of binding site interactions are given in Table 2.

Compounds **5d**, the most active inhibitor of hDHFR was selected for in depth analysis of binding site interactions (Fig. 4). One of the NH groups was making a hydrogen bond with Ser59. Two carbon hydrogen bonds were observed between methoxy carbon and Val115 and Ile7, the same methoxy carbon was also making alkyl interactions with Val115 and NADP, and pi alkyl interaction with Phe34. The methoxy substituted phenyl ring was making pi alkyl contacts with the NADP and Leu22 and a pi–pi stacked contact with Leu34. While the other phenyl ring was making a pi–pi stacked interaction with Phe31.

2.5 Molecular dynamics simulation studies

Analysis of protein-ligand RMSD show that the complex remained stable throughout the 100 ns simulation time period. The Cα protein RMSD value remained between 1.0 Å to 2.3 Å during the whole simulation mostly around 2.0 Å during the last 50 ns of simulation (Fig. 5). In ligand fit on protein RMSD, the value varied during the first 25 ns up-to 8.0 Å, then remained stable between 5.0 Å to 6.0 Å and stabilized around 5.5 Å towards the end of simulation. In protein RMSF, $C\alpha$ protein all residues showed value less than 3.0 Å (Fig. 6). Protein secondary structure was % total SSE at 43.78, while α -helix was 14.58 and β-strand at 29.21. Ligand RMSF values of fit ligand on protein showed all atoms have RMSF values between 2-3 Å where pyrrolidine moiety and methyl of anisole moiety showed relatively higher RMSF values as compared to other atoms in the ligand (ESI Fig. 1†). Protein-ligand contacts showed GLY 20 showed mostly hydrogen bonds and water bridges interactions with the highest interaction fraction around 0.6. Ligand-protein contacts showed nitrogen from thiosemicarbazide moiety formed interactions with GLY 20 for 55% of simulation time (ESI Fig. 2†). Ligand torsion profile showed most fluctuation and movement in the bonds of pyrrolidine and anisole moieties in the compound while thiosemicarbazide moiety remained relatively stable in bonds torsion (ESI Fig. 3†).

2.6 ADME studies

In silico ADME evaluation of all compounds **5a-r** was carried out using Swiss ADME. Overall, all compounds indicated favorable

Table 2 Binding site interactions of docked conformations of compounds (5a-r)

		Distance (Å)	Hydrophobic					
Code	Hydrogen bond		Alkyl	Pi-alkyl	Pi–pi stacked	Other		
5a	SER59	1.74	ALA9	PHE34				
	ASP21	2.1	NDP202	LEU22				
	SER59	2.1		ILE60				
				NDP202				
5b	VAL115	1.9	PRO61	LEU22	PHE31	VAL8 halogen (fluorine)		
	ASP21	2.67		PRO61	PHE34	GLU30 halogen (fluorine)		
				ILE7		SER59 pi-lone pair		
				ALA9				
5 c	PHE31	2.74	ILE60	PHE34	PHE31			
	PHE31	2.91	VAL115	LEU22				
				ILE60				
5 d	SER59	2.12	VAL115	PHE34	PHE31			
	ILE7	3.06	NDP202	LEU22	PHE34			
	VAL115	2.71		NDP202				
	NDP202	2.97						
5e	LEU22	2.43	PRO26	ILE60	PHE31			
	SER59	1.86	PRO61	PRO61				
	ASP21	1.74		PRO61				
	ASN64	3.03						
5f	SER59	2.14		LEU22	PHE31			
	PHE31	2.58		NDP202				
5g	SER59	2.15	ARG32	PHE34	PHE31			
8			NDP202	LEU22	PHE34			
5h	VAL115	2.1	PRO26	PHE34	PHE31			
			PRO61	LEU22				
			ILE7	ILE60				
				PRO61				
				ALA9				
				NDP202				
5i	VAL115	2.03	ALA9		PHE31			
	ASN64	2.08	LEU22	TRP24	PHE34 (pi–pi T-shaped)			
	ASN64	1.79		TRP24	(r r			
				PHE31				
				PHE31				
				ALA9				
				LEU22				
				NDP202				
5j	THR136	2.31	PRO61	LEU22	PHE34	GLU30 attractive charge		
-,	lig:H33 - VAL115:O	2.16		PRO61		SER59 pi-lone pair		
	PHE34:HA:O25	2.96		ILE7		sanos prione pan		
	11120111111020	2.50		ALA9				
5k	ASP21	1.99	LEU67	ILE60		ASP21 salt bridge; attractive charge		
	SER59	2.17	LLC0.	12200		1251 21 Sait Strage, accraetive charg		
	ASP21	3.01						
	SER59	1.94						
	ASP21	2.37						
5 l	SER59	1.66		ILE60	PHE31			
01	ASP21	1.64		PRO61	111201			
	7101 21	1.01		PRO26				
				PRO61				
5m	VAL115	1.9	ALA9	LEU22	PHE31	SER59 pi-lone pair		
JIII	ASP21	2.6	PRO61	PRO61	PHE34	SER39 prione pan		
	1 1.J1 2/1	2.0	ILE7	ILE7	111101			
			iuli/	ALA9				
5n	SED50	1.76	LEU22	PHE31:Br24	PHE31			
5n	SER59 PHE31	2.33	LEUZZ	ILE60	PHE31 PHE34 (pi–pi T-shaped)			
	LUESI	4.33		LEU22	rinesa (hi-hi i-siiahea)			
Eo.	ACD21	1.04	37AT 44F	NDP202				
50	ASP21	1.94	VAL115	PHE34				
	VAL115 LEU22	2.68 3.17	NDP202	LEU22 ILE60				

Table 2 (Contd.)

Code	Hydrogen bond	Distance (Å)	Hydropho	bic		<u></u>		
			Alkyl	Pi-alkyl	Pi–pi stacked	Other		
				NDP202				
				LEU22				
				PRO26				
5 p	ASP21	1.94	VAL115	PHE34				
•	VAL115	2.68	NDP202	LEU22				
	LEU22	3.17		ILE60				
				NDP202				
				LEU22				
				PRO26				
5q	SER59	1.73		ILE60	PHE31	THR56 halogen (fluorine)		
•	ILE60	1.92		LEU22				
	PHE31	2.85		ILE60				
				NDP202				
5r	PRO61	2.72	LEU67	PHE34				
			ILE60	ILE60				
				LEU22				
				ILE60				
				NDP202				

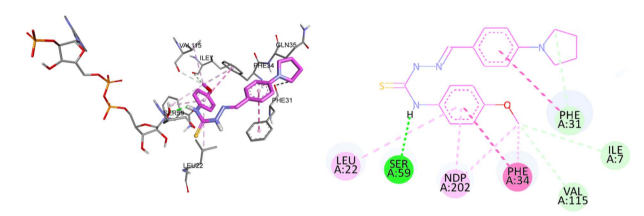


Fig. 4 3D and 2D binding site interactions of most hDHFR inhibitor 5d

ADME profile as indicated by several physicochemical parameters given in (Table 3). Oral bioavailability of a compound is represented in the form of a bioavailability radar diagram based on parameters such lipophilicity, size, polarity, solubility, saturation rate and molecular flexibility. All compounds were predicted to have good oral bioavailability, as a selected example radar diagram of most active compound 5d is shown in (Fig. 7). All compounds were predicted to have good to moderate water solubility, and with high gastrointestinal (GI) absorption. Interestingly most of the compounds were predicted to cross the blood–brain barrier (BBB). This ability to cross BBB, coupled with potent DHFR inhibition activity can be tailored for the design of novel anticancer drugs targeting brain tumors.

3. Experimental

3.1 Chemistry

All of the chemicals needed to synthesize 4-(pyrrolidin-1-yl) benzaldehyde-based thiosemicarbazones were purchased from

Sigma-Aldrich. Petroleum ether, ethyl acetate, glacial acetic acid, ethanol, and methanol were among the chemicals and solvents purchased from Merck and utilized in their original forms. Aluminum-backed silica gel plates were used to monitor the start and finish of the reaction. ^1H and $^{13}\text{C-NMR}$ spectra of were obtained at 25 $^{\circ}\text{C}$ using a Bruker Ascend 600 MHz NMR spectrometer in deuterated solvent such as DMSO-d₆ (600 MHz for ^{1}H and 151 MHz for ^{13}C). Chemical shifts (ppm) were used to portray NMR spectra, and coupling constants (f) were shown in hertz (Hz) to show signal multiplicity HPLC was carried out on Agilent, Germany (liquid chromatographic column 150 mm \times 4.6 mm (id) packed with 5-micron C18; 263 nm).

3.2 General method for the synthesis of 4-(pyrrolidin-1-yl) benzaldehyde (3)

Compound (3) was synthesized using a method that had been previously reported.³⁷ To summarize, pyrrolidine (1 mmol) was dissolved in 4.0 ml of DMF. After that, the solution was heated

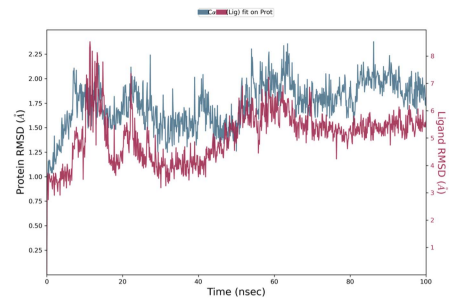


Fig. 5 RMSD graph of 5d-hDHFR protein ligand complex for 100 ns simulation run

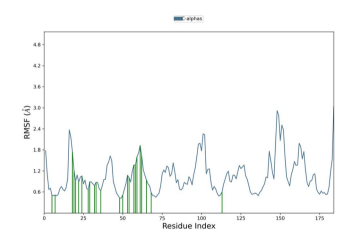


Fig. 6 Protein RMSF graph of **5d** complexed with hDHFR. Residues that interact with the ligand are shown in green vertical bars.

to 80 °C, while being stirred, anhydrous K_2CO_3 (2 mmol) was added. After 30 minutes, 4-fluorobenzaldehyde (1 mmol) was added in reaction mixture. The heating was continued for 12 hours. Once the reaction was finished, the mixture was cooled to room temperature and added to the ice water drop by drop. The product was precipitated out, filtered and dried. The formed product was pure and used without additional purification.

3.3 General method for the synthesis and characterization of compounds 5(a-r)

A previously reported³⁸ one-step condensation method was employed to synthesize thiosemicarbazones, with some modifications to the solvent, in a tropical setting. First, thiosemicarbazides (4a-r) (5 mmol) and compound (3) (0.1 g, 5 mmol) were mixed in a 20 ml ethanol solution in an oven-dried,

single-neck round bottom flask. As a catalyst, a small amount (three to four drops) of glacial acetic acid was added. After that, the mixture was refluxed for four to eight hours while thin layer chromatography (TLC) was used for monitoring the reaction's development. The resultant solid product was filtered and dried once the reaction was finished. The residue was then crystallized using ethanol to obtain the required thiosemicarbazones 5(a-r). Characterization data of each synthesized compound is given below.

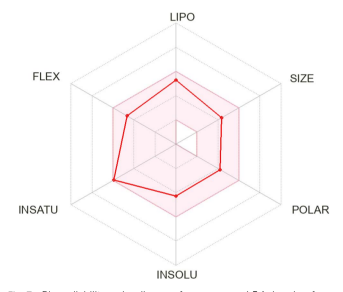
3.3.1 *N*-Cyclohexyl-2-[4-(pyrrolidin-1-yl)benzylidene] hydrazinecarbothioamide (5a). Color: red, yield: 75%, melting point: 234–236 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.14 (1 H, s), 7.93 (1 H, s), 7.79 (1 H, d, J 8.6), 7.56 (2 H, d, J 8.3), 6.54 (2 H, d, J 8.4), 4.18 (1 H, ddt, J 11.2, 6.9, 3.8), 3.28 (4 H, d, J 6.3), 2.01–1.91 (4 H, m), 1.88 (2 H, dd, J 12.5, 4.2), 1.72 (2 H, dt, J 13.2, 3.7), 1.60 (1 H, dt, J 12.9, 3.9), 1.42 (2 H, qd, J 12.2, 3.5), 1.29 (2 H, qt, J 12.8, 3.4), 1.15 (1 H, ddd, J 15.9, 8.0, 3.5); ¹³C NMR (151 MHz, DMSO) δ 175.37, 149.25, 143.91, 129.26, 121.07, 111.91, 52.77, 47.70, 32.45, 25.63, 25.42, 25.38, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.164 min, purity: 98.7%, $C_{18}H_{26}N_4$ S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 331.1956, found: 331.1956.

3.3.2 *N*-(4-Fluorophenyl)-2-[4-(pyrrolidin-1-yl)benzylidene] hydrazinecarbothioamide (5b). Color: brown, yield: 81%, melting point: 249–251 °C, $\delta_{\rm H}$ (600 MHz, DMSO- $d_{\rm 6}$) 11.59 (1 H, s), 9.92 (1 H, s), 8.03 (1 H, s), 7.68 (2 H, d, J 8.3), 7.58 (2 H, dd, J 8.7, 5.0), 7.19 (2 H, t, J 8.6), 6.55 (2 H, d, J 8.4), 3.28 (4 H, d, J 6.5), 1.97 (4 H, d, J 5.9); ¹³C NMR (151 MHz, DMSO) δ 175.61, 160.74, 159.13, 149.41, 144.78, 136.08, 129.67, 128.27, 128.22, 120.94, 115.13, 114.98, 111.88, 47.71, 25.43, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.088 min, purity: 98.2%, C₁₈H₁₉FN₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 343.1392, found: 343.1392.

3.3.3 *N*-(2,3-Dichlorophenyl)-2-[4-(pyrrolidin-1-yl) benzylidene]hydrazinecarbothioamide (5c). Color: brown, yield: 91%, melting point: 301–303 °C, $\delta_{\rm H}$ (600 MHz, DMSO- $d_{\rm 6}$) 11.85

Table 3 In silico ADME properties of compounds (5a-r)

Code	MW	RB	HBA	HBD	TPSA	$\log P$	$\log S$	GI	BBB	B score
5a	330.49	6	1	2	71.75	3.53	-4.60	High	Yes	0.55
5 b	343.43	6	2	2	71.75	3.70	-5.95	High	No	0.55
5 c	393.33	6	1	2	71.75	4.45	-6.87	High	Yes	0.55
5d	354.47	7	2	2	80.98	3.32	-5.79	High	No	0.55
5e	352.50	6	1	2	71.75	4.01	-5.73	High	Yes	0.55
5f	324.44	6	1	2	71.75	3.38	-5.68	High	Yes	0.55
5g	338.47	6	1	2	71.75	3.72	-6.06	High	Yes	0.55
5h	354.47	7	2	2	80.98	3.38	-5.79	High	No	0.55
5i	372.91	7	1	2	71.75	3.95	-6.67	High	Yes	0.55
5j	369.44	7	3	2	117.57	2.98	-5.51	High	No	0.55
5k	361.50	8	3	2	84.22	1.99	-4.14	High	No	0.55
5l	352.50	8	1	2	71.75	3.70	-6.48	High	Yes	0.55
5m	403.34	6	1	2	71.75	4.02	-6.48	High	Yes	0.55
5n	403.34	6	1	2	111.66	4.01	-6.48	High	Yes	0.55
50	369.44	7	3	2	117.57	2.98	-5.51	High	No	0.55
5 p	338.47	7	1	2	71.75	3.42	-6.08	High	Yes	0.55
5q	342.43	6	2	2	71.75	3.73	-7.59	High	Yes	0.55
5r	352.50	7	1	2	71.75	3.74	-6.46	High	Yes	0.55



 $\label{eq:Fig.7} \textbf{Fig. 7} \quad \text{Bioavailability radar diagram for compound } \textbf{5d} \text{ showing favorable oral bioavailability profile.}$

(1 H, s), 10.02 (1 H, s), 8.05 (1 H, s), 7.84 (1 H, d, J 8.0), 7.64 (2 H, d, J 8.3), 7.54 (1 H, d, J 8.1), 7.39 (1 H, t, J 8.1), 6.56 (2 H, d, J 8.4), 3.29 (4 H, d, J 6.4), 2.01–1.90 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 175.64, 149.52, 145.07, 139.17, 132.00, 129.59, 129.12, 128.22, 128.12, 127.96, 120.73, 111.98, 47.72, 25.43, HPLC: CH₃CN: H₂O = 80:20; tR: 3.956 min, purity: 98.3%, C₁₈H₁₈Cl₂N₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 393.0707, found: 393.0706.

3.3.4 *N*-(4-Methoxyphenyl)-2-[4-(pyrrolidin-1-yl)

benzylidene]hydrazinecarbothioamide (**5d**). Color: light yellow, yield: 87%, melting point: 232–234 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.49 (1 H, s), 9.80 (1 H, s), 8.03 (1 H, s), 7.67 (2 H, d, J 8.3), 7.43 (2 H, d, J 8.4), 6.92 (2 H, d, J 8.5), 6.55 (2 H, d, J 8.4), 3.77 (3 H, s), 3.28 (4 H, d, J 6.4), 2.03–1.85 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 175.70, 157.20, 149.34, 144.40, 132.62, 129.59, 127.74,

121.07, 113.64, 111.88, 55.69, 47.71, 25.43, HPLC: $CH_3CN : H_2O = 80 : 20$; tR: 1.977 min, purity: 98.3%, $C_{19}H_{22}N_4OS$, QTOF MS ES+ (m/z): $[M + H]^+$, calcd: 355.1592, found: 355.1582.

3.3.5 N-(2,6-Dimethylphenyl)-2-[4-(pyrrolidin-1-yl)

benzylidene]hydrazinecarbothioamide (5e). Color: yellow, yield: 85%, melting point: 218–220 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.46 (1 H, s), 9.64 (1 H, s), 8.01 (1 H, s), 7.68 (2 H, d, J 8.4), 7.11 (3 H, q, J 5.1), 6.54 (2 H, d, J 8.4), 3.29 (4 H, d, J 6.4), 2.20 (6 H, s), 2.02–1.91 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 176.37, 149.25, 143.92, 137.89, 137.01, 129.50, 127.97, 127.18, 121.34, 111.86, 47.72, 25.43, 18.63, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.300 min, purity: 97.9%, C₂₀H₂₄N₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 353.1799, found: 353.1789.

3.3.6 *N*-Phenyl-2-[4-(pyrrolidin-1-yl)benzylidene]

hydrazinecarbothioamide (5f). Color: yellow, yield: 83%, melting point: 270–272 °C, δ_H (600 MHz, DMSO-d₆) 11.58 (1 H, s), 9.91 (1 H, s), 8.04 (1 H, s), 7.75–7.63 (2 H, m), 7.66–7.56 (2 H, m), 7.36 (2 H, t, J7.9), 7.19 (1 H, dd, J8.1, 6.6), 6.55 (2 H, d, J8.6), 3.33–3.24 (4 H, m), 2.00–1.91 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 175.23, 149.41, 144.69, 139.71, 129.66, 128.45, 125.88, 125.41, 120.95, 111.89, 47.72, 25.43, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.150 min, purity: 97.5%, C₁₈H₂₀N₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 325.1486, found: 325.1463.

3.3.7 2-[4-(Pyrrolidin-1-yl)benzylidene]-*N*-(*p*-tolyl)hydrazinecarbothioamide (5g). Color: yellow, yield: 81%, melting point: 230–232 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.53 (1 H, s), 9.83 (1 H, s), 8.03 (1 H, s), 7.68 (2 H, d, *J* 8.7), 7.52–7.42 (2 H, m), 7.16 (2 H, d, *J* 8.1), 6.55 (2 H, d, *J* 8.7), 3.32–3.23 (4 H, m), 2.31 (3 H, s), 2.01–1.91 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 175.32, 149.37, 144.53, 137.14, 134.54, 129.62, 128.92, 125.88, 121.00, 111.89, 47.71, 25.43, 21.06, HPLC: CH₃CN: H₂O = 80: 20; *t*R: 2.459 min, purity: 97.7%, C₁₉H₂₂N₄S, QTOF MS ES+ (*m*/*z*): [M + H]⁺, calcd: 339.1643, found: 339.1630.

3.3.8 N-(3-Methoxyphenyl)-2-[4-(pyrrolidin-1-yl)

benzylidene]hydrazinecarbothioamide (5h). Color: shine yellow, yield: 95%, melting point: 220–222 °C, $\delta_{\rm H}$ (600 MHz,

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DMSO- d_6) 11.60 (1 H, s), 9.86 (1 H, s), 8.04 (1 H, s), 7.74–7.64 (2 H, m), 7.36 (1 H, t, J 2.1), 7.30–7.19 (2 H, m), 6.76 (1 H, dt, J 7.5, 2.1), 6.60–6.50 (2 H, m), 3.77 (3 H, s), 3.28 (4 H, q, J 4.7, 3.2), 2.01–1.91 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 174.90, 159.44, 149.43, 144.77, 140.81, 129.69, 129.14, 120.88, 117.71, 112.09, 111.90, 111.17, 110.83, 55.60, 47.71, 25.43, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.166 min, purity: 97.1%, C₁₉H₂₂N₄OS, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 355.1592, found: 355.1582.

3.3.9 *N*-(4-Chlorobenzyl)-2-[4-(pyrrolidin-1-yl)benzylidene] hydrazinecarbothioamide (5i). Color: brown, yield: 79%, melting point: 234–236 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.36 (1 H, s), 8.89 (1 H, t, *J* 6.3), 7.97 (1 H, s), 7.59 (2 H, d, *J* 8.6), 7.38 (4 H, s), 6.53 (2 H, dd, *J* 8.6, 2.0), 4.81 (2 H, d, *J* 6.3), 3.32–3.22 (4 H, m), 1.99–1.92 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 177.21, 149.26, 144.13, 139.30, 131.64, 129.62, 129.56, 129.30, 129.07, 128.54, 121.15, 111.87, 47.70, 46.27, 25.42, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.558 min, purity: 96.3%, C₁₉H₂₁ClN₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 373.1253, found: 373.1253.

3.3.10 *N*-(4-Nitrophenyl)-2-[4-(pyrrolidin-1-yl)benzylidene] hydrazinecarbothioamide (5j). Color: orange, yield: 85%, melting point: 250–252 °C, $\delta_{\rm H}$ (600 MHz, DMSO- $d_{\rm 6}$) 11.95 (1 H, s), 10.28 (1 H, s), 8.26–8.21 (2 H, m), 8.15–8.10 (2 H, m), 8.08 (1 H, s), 7.72–7.67 (2 H, m), 6.61–6.52 (2 H, m), 3.32–3.25 (4 H, m), 2.00–1.93 (4 H, m); $^{13}{\rm C}$ NMR (151 MHz, DMSO) δ 174.30, 149.62, 146.12, 145.93, 143.53, 129.95, 124.21, 124.19, 120.52, 111.91, 47.72, 25.42, HPLC: CH₃CN:H₂O = 80:20; tR: 2.344 min, purity: 98.8%, $C_{18}H_{19}N_5O_2S$, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 370.1337, found: 370.1337.

3.3.11 *N*-(2-Morpholinoethyl)-2-[4-(pyrrolidin-1-yl) benzylidene]hydrazinecarbothioamide (5k). Color: off-white, yield: 89%, melting point: 227–229 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.25 (1 H, s), 8.26 (1 H, t, *J* 5.5), 7.93 (1 H, s), 7.60–7.51 (2 H, m), 6.62–6.51 (2 H, m), 3.65 (2 H, q, *J* 6.4), 3.60 (4 H, t, *J* 4.6), 3.32–3.25 (4 H, m), 2.53 (2 H, t, *J* 6.7), 2.44 (4 H, s), 2.00–1.92 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 176.54, 149.28, 143.67, 129.02, 121.13, 111.95, 66.83, 57.09, 53.65, 47.72, 25.43, HPLC: CH₃CN: H₂O = 80: 20; tR: 1.610 min, purity: 98.9%, C₁₈H₂₇N₅OS, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 362.2014, found: 362.2014.

3.3.12 *N*-Phenethyl-2-[4-(pyrrolidin-1-yl)benzylidene] hydrazinecarbothioamide (5l). Color: off-white, yield: 91%, melting point: 237–239 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.25 (1 H, s), 8.31 (1 H, t, *J* 5.9), 7.93 (1 H, s), 7.60–7.52 (2 H, m), 7.33 (2 H, t, *J* 7.5), 7.31–7.26 (2 H, m), 7.26–7.21 (1 H, m), 6.58–6.52 (2 H, m), 3.76 (2 H, ddd, *J* 9.4, 7.7, 5.9), 3.33–3.23 (4 H, m), 2.96–2.87 (2 H, m), 2.01–1.90 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 176.61, 149.26, 143.73, 139.82, 129.15, 129.09, 128.94, 126.66, 121.17, 111.91, 47.72, 45.37, 35.51, 25.43, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.608 min, purity: 98.4%, C₂₀H₂₄N₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 353.1799, found: 353.1799.

3.3.13 *N*-(4-Bromophenyl)-2-[4-(pyrrolidin-1-yl) benzylidene]hydrazinecarbothioamide (5m). Color: off-white, yield: 91%, melting point: 228–230 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.67 (1 H, s), 9.95 (1 H, s), 8.04 (1 H, s), 7.70–7.66 (2 H, m), 7.64–7.59 (2 H, m), 7.56–7.51 (2 H, m), 6.61–6.51 (2 H, m), 3.32–3.25 (4 H, m), 1.99–1.92 (4 H, m); $^{13}{\rm C}$ NMR (151 MHz, DMSO) δ 175.07, 149.46, 145.04, 139.15, 131.23, 129.73, 127.80, 120.84,

117.55, 111.89, 47.72, 25.43, HPLC: $CH_3CN: H_2O = 80: 20; tR: 3.025$ min, purity: 97.4%, $C_{18}H_{19}BrN_4S$, QTOF MS ES+ $(m/z): [M + H]^+$, calcd: 403.0592, found: 403.0503.

3.3.14 *N*-(3-Bromophenyl)-2-[4-(pyrrolidin-1-yl)

benzylidene]hydrazinecarbothioamide (5n). Color: green, yield: 85%, melting point: 223–225 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.71 (1 H, s), 8.05 (1 H, s), 7.95 (1 H, t, J 2.0), 7.75–7.66 (3 H, m), 7.36 (1 H, dt, J 8.0, 1.4), 7.32 (1 H, t, J 8.0), 6.60–6.52 (2 H, m), 3.34–3.24 (4 H, m), 2.01–1.90 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 174.93, 149.48, 145.18, 141.36, 130.25, 129.78, 127.94, 127.88, 124.58, 120.91, 120.79, 112.06, 111.88, 47.72, 25.43, HPLC: CH₃CN: H₂O = 80:20; tR: 3.022 min, purity: 97.2%, C₁₈H₁₉BrN₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 403.0592, found: 403.0522.

3.3.15 *N*-(3-Nitrophenyl)-2-[4-(pyrrolidin-1-yl)benzylidene] hydrazinecarbothioamide (5o). Color: yellow, yield: 85%, melting point: 220–222 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.85 (1 H, s), 10.24 (1 H, s), 8.71 (1 H, t, *J* 2.2), 8.21–8.13 (1 H, m), 8.07 (1 H, s), 8.02 (1 H, ddd, *J* 8.2, 2.4, 1.0), 7.77–7.66 (2 H, m), 7.64 (1 H, t, *J* 8.1), 6.62–6.51 (2 H, m), 3.33–3.25 (4 H, m), 2.00–1.92 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 174.96, 149.55, 147.69, 145.63, 140.99, 131.83, 129.86, 129.56, 120.68, 119.71, 119.69, 111.89, 47.72, 25.43, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.207 min, purity: 97.9%, C₁₈H₁₉N₅O₂S, QTOF MS ES+ (m/z): [M + H][†], calcd: 370.1337, found: 370.1337.

3.3.16 *N*-Benzyl-2-[4-(pyrrolidin-1-yl)benzylidene]

hydrazinecarbothioamide (**5p**). Color: off-white, yield: 85%, melting point: 181–183 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.33 (1 H, s), 8.85 (1 H, t, J 6.3), 7.97 (1 H, s), 7.64–7.55 (2 H, m), 7.38–7.35 (2 H, m), 7.35–7.30 (2 H, m), 7.27–7.20 (1 H, m), 6.58–6.48 (2 H, m), 4.84 (2 H, d, J 6.3), 3.32–3.22 (4 H, m), 2.02–1.90 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 177.19, 149.25, 144.00, 140.21, 129.27, 128.60, 127.73, 127.13, 121.19, 111.88, 47.70, 46.93, 25.42, HPLC: CH₃CN: H₂O = 80: 20; tR: 2.202 min, purity: 98.1%, C₁₉H₂₂N₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 339.1643, found: 339.1643.

3.3.17 N-(2-Fluorophenyl)-2-[4-(pyrrolidin-1-yl)

benzylidene]hydrazinecarbothioamide (**5q**). Color: green, yield: 87%, melting point: 228–230 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.60 (1 H, s), 9.92 (1 H, s), 8.03 (1 H, s), 7.74–7.64 (2 H, m), 7.65–7.53 (2 H, m), 7.26–7.13 (2 H, m), 6.62–6.51 (2 H, m), 3.28 (4 H, d, J 6.3), 2.00–1.91 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 175.61, 160.73, 159.13, 149.41, 144.77, 136.09, 136.07, 129.67, 128.27, 128.21, 120.95, 115.13, 114.98, 111.88, 47.71, 25.43, HPLC: CH₃CN: H₂O = 80:20; tR: 2.164 min, purity: 98.6%, C₁₈H₁₉FN₄S, QTOF MS ES+ (m/z): [M + H]⁺, calcd: 343.1392, found: 343.1395.

3.3.18 *N*-(4-Methylbenzyl)-2-[4-(pyrrolidin-1-yl)

benzylidene]hydrazinecarbothioamide (5**r**). Color: off-white, yield: 87%, melting point: 205–207 °C, $\delta_{\rm H}$ (600 MHz, DMSO- d_6) 11.30 (1 H, s), 8.78 (1 H, t, J 6.3), 7.96 (1 H, s), 7.63–7.53 (2 H, m), 7.32–7.21 (2 H, m), 7.13 (2 H, d, J 7.8), 6.59–6.47 (2 H, m), 4.79 (2 H, d, J 6.2), 3.30–3.20 (4 H, m), 2.28 (3 H, s), 2.00–1.89 (4 H, m); ¹³C NMR (151 MHz, DMSO) δ 177.08, 149.24, 143.94, 137.12, 136.15, 129.25, 129.14, 127.77, 121.20, 111.88, 47.70, 46.70, 25.42, 21.18, HPLC: CH₃CN : H₂O = 80 : 20; tR: 2.603 min,

purity: 98.3%, $C_{20}H_{24}N_4S$, QTOF MS ES+ (m/z): $[M + H]^+$, calcd: 353.1799, found: 353.1799.

4. Conclusion

In conclusion, the comprehensive study into the potential of 4pyrrolidine-based thiosemicarbazones 5(a-r) as DHFR inhibitors through in vitro and in silico studies sheds light on their therapeutic applications. The synthesized promising compounds 5(a-r) demonstrated notable inhibitory activity against DHFR with IC50 value in the range of 12.37 \pm 0.48 μM to $54.10 \pm 0.72~\mu M$, showcasing their potential as effective agents against various diseases, including cancer and microbial infections. Amongst all derivatives, 5d is more potent towards DHFR enzyme. Furthermore, the molecular docking simulations provided valuable insights into the binding interactions between the compounds 5(a-r) and the target enzyme, elucidating their mechanism of action at the molecular level. These findings underscore the significance of further exploration and optimization of 4-pyrrolidine-based thiosemicarbazones for the development of novel DHFR inhibitors with enhanced efficacy and reduced toxicity, thereby offering new avenues for drug discovery and therapeutic invention.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors have declared no conflict of interest.

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