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Design and evaluation of sulfadiazine derivatives as potent dual inhibitors of EGFR^{WT} and EGFR^{T790M}: integrating biological, molecular docking, and ADMET analysis†

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A series of derivatives (5-14) were synthesized through the diazotization of sulfadiazine with active methylene compounds. The chemical structures of these newly designed compounds were validated through spectral and elemental analysis techniques. The antiproliferative potential of derivatives 5-14 was assessed against three distinct cancer cell lines (A431, A549, and H1975) using the MTT assay. The results revealed that compounds 8, 12, and 14 exhibited the most potent antiproliferative activity, with IC_{50} values ranging from 2.31 to 7.56 μ M. Notably, these values were significantly lower than those of known EGFR inhibitors, including erlotinib, gefitinib, and osimertinib, suggesting the potential of these derivatives as novel antiproliferative agents. Furthermore, compound 12 was identified as the most potent inhibitor of both EGFRWT and EGFRT790M protein kinases, with IC50 values of 14.5 and 35.4 nM, respectively. These results outperformed those of gefitinib and osimertinib, which exhibited IC50 values of 18.2 and 368.2 nM, and 57.8 and 8.5 nM, respectively. Molecular docking studies of compounds 8, 12, and 14 within the ATP-binding sites of both EGFRWT and EGFRT790M corroborated the in vitro results when compared to erlotinib, gefitinib, and osimertinib. The docking results indicated that compound 8 exhibited a favorable binding affinity for both EGFRWT and EGFRT790M, with binding scores of -6.40 kcal mol⁻¹ and -7.53 kcal mol⁻¹, respectively, which were comparable to those of **gefitinib** and osimertinib, with binding scores of -8.01 and -8.72 kcal mol⁻¹, respectively. Furthermore, an assessment of the most promising EGFR inhibitors (8, 12, and 14) using the egg-boiled method for their in silico ADME properties revealed significant lipophilicity, blood-brain barrier (BBB) penetration, and gastrointestinal (GIT) absorption. Collectively, our designed analogs, particularly compounds 8, 12, and 14, exhibit promising dual antiproliferative and EGFRWT and EGFRT790M kinase inhibitory properties, positioning them as efficient candidates for further therapeutic development.

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1. Introduction

Molecular hybridization is a pivotal strategy in drug design,¹⁻⁵ involving the deliberate fusion of two or more biologically relevant pharmacophores, often bridged by a spacer or linker, to yield a novel hybrid entity with significantly enhanced activity and efficacy compared to the parent molecule.⁶⁻⁹ This strategy can also decrease undesirable side effects, thereby improving the overall therapeutic index.¹⁰⁻¹³ Moreover, recent investigations have revealed that sulfonamides possess potent growth inhibitory activity against a diverse range of tumor lines and tyrosine kinase (TK) enzymes, underscoring their potential as promising anticancer agents.

The utilization of pyrimidines as a scaffold for producing new EGFR inhibitors that target multiple pathways, particularly oncogenesis, has been correlated with the significant anticancer

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properties of sulfonamides. These compounds have been shown to exhibit effective activity across a multitude of pathways, including the inhibition of carbonic anhydrase, matrix metalloproteinase, NADPH reductase, histone deacetylase, and PI3K. Notably, compounds **I–III** demonstrated more pronounced activities against both HER2 and EGFR compared to **erlotinib**, with concomitant increases in Bcl-2, Bax, and caspase-3 expression levels and the induction of cell cycle arrest in the G2/M phase^{12–16} (Fig. 1).

EGFR, a receptor TK, plays a significant role in maintaining epithelial tissue homeostasis and is essential for generating and regulating cellular processes. However, the overexpression of the EGFR family members can lead to aberrant cell signaling, contributing to the progression and development of certain solid tumors, including non-small cell lung cancer (NSCLC) and breast cancers. 17,18 The EGFR-TK domain comprises five distinct sites, including the ATP-binding pocket, which is comprised of (1) the adenine binding pocket, where H-bonds are formed with the adenine ring through residue amino acids; (2) the hydrophilic pocket (ribose region); (3) the hydrophobic region I, which is critical for receptor inhibition activity, although it is not targeted by ATP; (4) the hydrophobic region II, which is also essential and frequently targeted by EGFR-TKIs (tyrosine kinase inhibitors) with a Y-shaped structure; and (5) the phosphatebinding region, which influences the pharmacokinetic features of inhibitors. Importantly, most EGFR-TKIs exhibit a Yshaped structure and possess four critical pharmacophoric properties, which are essential for their inhibitory activity. 19-22

The first generation of EGFR-TKIs, comprising **gefitinib** and **erlotinib**, significantly enhanced the therapy landscape for NSCLC, however they were also associated with adverse effects, such as corneal damage, trichomegaly, and occasionally fatal interstitial lung disease (ILD). Furthermore, the development of acquired resistance due to EGFR-TK mutations compromised the antitumor efficacy of these treatments.^{22–25} This type of mutation diminishes the selectivity of ATP-competitive agents for the kinase while increasing EGFR's sensitivity to ATP, thereby facilitating the rapid emergence of drug-resistant mechanisms.²⁶ Particularly, the T790M "gatekeeper" mutation in EGFR is a prevalent mechanism of resistance, frequently observed in patients who have developed resistance to 1st generation EGFR-TKIs.^{27–29}

The development of second-generation EGFR-TKIs, including pelitinib and neratinib, was driven by the need to

address resistance in NSCLC treatment regimens resulting from EGFR-TK mutations.^{30–33} These inhibitors interact with Cys797 near the EGFR's ATP-binding groove *via* electrophilic Michaelacceptor moieties, thereby inactivating the protein. However, their non-selective inhibition of both EGFR^{WT} and EGFR^{T790M} led to a relatively lower maximum tolerated dose (MTD) and adverse clinical outcomes.^{34–36}

Third generation EGFR-TKIs, including olmutinib^{37,38} and **osimertinib** (AZD9291),³⁹ are characterized as irreversible covalent inhibitors with enhanced binding affinity for EGFR^{T790M} and lower affinity for EGFR^{WT}. These inhibitors exhibit sustained sub-nanomolar IC₅₀ values for prevalent activating mutations (del19 and L858R). Due to its favorable biological properties, the FDA approved AZD9291 in 2015 for patients with NSCLC therapeutic regimens with mutant EGFR. However, a 2016 safety study reported two cases of Stevens-Johnson syndrome and severe epidermal necrolysis.⁴⁰⁻⁴² AZD9291 is currently the primary 3rd generation authorized as a first-line therapy for patients with EGFR-positive activating mutations (del19 and L858R), NSCLC, and metastatic EGFR^{T790M}-positive NSCLC, as identified by an approved FDA test

Investigations into the mechanisms underlying resistance to 3rd generation have revealed the presence of both primary and acquired resistant mechanisms.40 In order to inform subsequent treatment strategies, patients experiencing NSCLC progression after 3rd generation therapies should undergo comprehensive next-generation sequencing (NGS) analysis of plasma or tissue samples to elucidate the specific resistance mechanisms employed.35 Further research into the mechanisms of resistance to 3rd generation is imperative for the development of effective therapeutic strategies. This study examines the findings on 3rd generation resistance in aggressive NSCLC, providing a detailed examination of the underlying molecular mechanisms and discussing novel treatment approaches for addressing this complex issue. Mobocertinib (Fig. 2), the latest addition to the 3rd generation family, has been granted accelerated approval by the FDA in 2021 for the treatment of locally advanced or metastatic NSCLC harboring exon 20 insertions, thereby offering a promising therapeutic option for patients with this specific molecular subtype.

Our study aimed to design novel EGFR-TKI derivatives, incorporating the four primary pharmacophoric properties of EGFR-TKIs, with the goal of enhancing their efficacy and

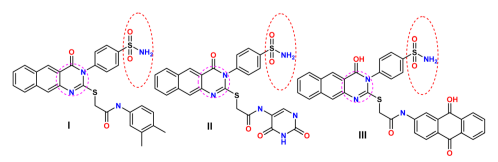


Fig. 1 Examples of sulfonamide-based derivatives as EGFR inhibitors.

First generation of EGFR

Gefitinib (IV)

Second generation of EGFR

Afatinib (VI)

Pelitinib (VII)

Third generation of EGFR

Fig. 2 The chemical structures of reported EGFR-TKIs

selectivity. To achieve this, we synthesized a series of derivatives by structurally modifying **erlotinib** at four distinct positions (Fig. 3). Building upon our previous research focused on the scope of azole and pyrimidine derivative-based scaffolds, which have demonstrated promising antiproliferative activity against both EGFR^{WT} and EGFR^{T790M}, we intended to further investigate the potential of these scaffolds in the development of nextgeneration EGFR-TKIs.

Osimertinib (VIII)

2. Results and dissection

2.1. Chemistry

In the present study, we initiated the synthesis by reacting 4-acetamidobenzene sulfonyl chloride (1) with 2-amino-pyrimidine (2), resulting in the formation of N-(4-(N-(pyrimidin-2-yl)sulfamoyl)phenyl)acetamid (3). Subsequently, compound (3) was subjected to hydrolysis, yielding sulfadiazine (4). The next step involved the diazotization of sulfadiazine, which was then coupled with a diverse array of compounds, including ethyl acetoacetate, acetylacetone, ethyl benzoylacetate, benzoyl acetone, ethyl cyanoacetate, cyanothioacetamide, a dimer of

malononitrile, diethyl malonate, N-[4-amino-5-cyano-6-(methylthio)pyridin-2-yl]-3-oxobutanamide, and 4,6-diamino-2-(methylthio)nicotinonitrile, in the presence of pyridine as a solvent. This reaction sequence led to the successful synthesis of hydrazone derivatives 5–14, as illustrated in Scheme 1.

Mobocertinib (IX)

The chemical identities of the synthesized derivatives 5-14 were unequivocally confirmed through a combination of various spectral analyses. The IR spectra revealed the presence of NH₂ and NH groups, which were identified by absorption bands in the range of 3457-3103 cm⁻¹. Additionally, the IR spectra showed characteristic absorption bands for CN groups between 2215-2211 cm⁻¹ and C=O groups between 1725-1655 cm⁻¹. The ¹H NMR spectra of 5-14 displayed distinct signals for NH groups, which resonated between δ 13.65-11.34 ppm. Furthermore, the ¹H NMR spectra of derivatives 5, 7, 9, and 12 exhibited quartet and triplet signals for ester groups, which appeared between δ 4.37–4.23 and δ 1.29–1.24 ppm, respectively. Especially, the SCH3 group in compound 14 was identified as a singlet at δ 2.52 ppm. The ¹³C NMR spectra provided additional confirmation of the structures of all compounds. For example, the 13 C NMR spectrum of 13 showed

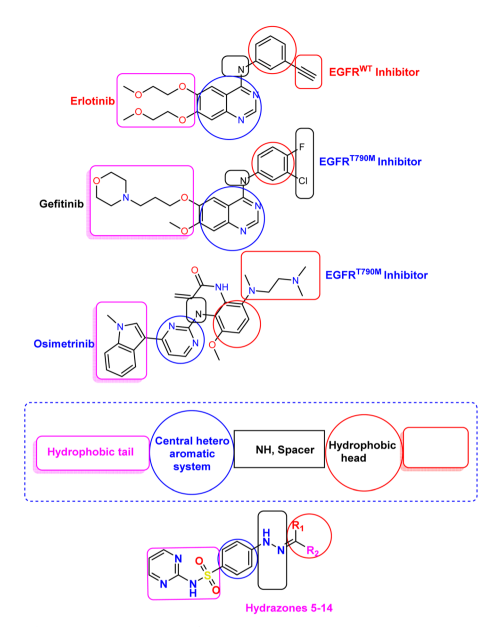


Fig. 3 Rationale design of the target derivatives utilizing gefitinib, osimertinib, and erlotinib

a signal at δ 12.87 ppm, which was attributed to the SCH₃ group, and a signal at δ 79.92 ppm, which was assigned to the C-CN group. Similarly, the 13C NMR spectrum of 12 revealed two distinct signals for C=O groups at δ 195.22 and 196.65 ppm, as well as a signal for a CH₃ group at δ 25.39 ppm. The elemental analysis data further verified the proposed structures of the sulfadiazine derivatives, providing a comprehensive confirmation of their chemical identities.

2.2. Biological evaluation

2.2.1. In vitro anti-proliferative activities. The antiproliferative activity of the synthesized derivatives was assessed toward NSCLC cells (A549 and H1975) and human epidermoid carcinoma cells (A431) using the MTT assay method.37-39,43-45 Notably, A431 and A549 cells overexpress EGFRWT, whereas H1975 cells harbor the EGFRT790M

mutation.45 The results, presented in Table 1, reveal the IC50 values (µM) of the investigated derivatives, with erlotinib, gefitinib, and osimertinib serving as reference standards. The data indicate that compounds 8, 12, and 14 exhibited the most potent anti-proliferative activity, surpassing that of gefitinib and erlotinib (IC₅₀ = 15.75 and 17.50 μ M, respectively), with IC₅₀ values ranging from 4.78 to 11.09 μM. In contrast, the remaining derivatives displayed comparable or reduced activity relative to gefitinib and erlotinib. Specifically, compounds 8, 12, and 14 demonstrated significant anti-proliferative activity against A431 cells, with IC50 values ranging from 4.78 to 7.56 μ M, which exceeded the potency of **gefitinib** (IC₅₀ = 9.60 μ M). Particularly, compound 8 exhibited more potent antiproliferative activity than osimertinib; however, its effect on the H1975 cell line was less pronounced than that of osimerti**nib** ($IC_{50} = 0.93 \mu M$).

Comp.	\mathbf{R}_1	\mathbb{R}_2
5	· vvv	NA.
6	No.	NA.
7		good of the same o
8	g de la companya de l	The state of the s
9		N N
10	H ₂ N s	N N
11	H ₂ N	N N
12		
13	HN N S NH ₂	-
	مم	H ₂ N S

Scheme 1 Synthesis of sulfadiazine-based derivatives 5-14.

2.2.2. EGFR^{WT} and EGFR^{T790M} kinase inhibitory activities. The sulfadiazine derivatives 5–14 were examined for their ability to inhibit the EGFR^{WT} and EGFR^{T790M} kinase enzymes employing the homogeneous time-resolved fluorescence (HTRF) KinEASE-TK assay, $^{40-42,46-50}$ with osimertinib and gefitinib as references (Table 2). The outcomes showed that the majority of the derivatives exhibited moderate inhibitory effects against EGFR^{WT}, with IC₅₀ values ranging from 14.5 to 52.6 nM. Particularly, sulfadiazines 12 and 14 demonstrated significantly greater inhibitory activity against EGFR^{WT} compared to gefitinib. In contrast, all of the sulfadiazine derivatives 5–14 displayed improved inhibitory effects against the mutant EGFR

Table 1 Anti-proliferative activity of the final compounds *versus* A549, A431, and H1975

	$IC_{50} (\mu M)^a$							
Comp.	A549	A431	H1975					
5	19.24 ± 3.09	18.21 ± 0.46	20.12 ± 2.21					
6	16.60 ± 0.92	17.21 ± 0.52	15.70 ± 0.21					
7	11.25 ± 1.09	13.89 ± 0.80	10.32 ± 1.80					
8	4.78 ± 0.12	4.12 ± 0.54	2.31 ± 0.23					
9	14.98 ± 1.27	14.70 ± 0.18	13.14 ± 0.67					
10	19.67 ± 0.34	18.23 ± 2.09	15.80 ± 2.48					
11	13.56 ± 1.32	15.09 ± 0.58	14.13 ± 1.23					
12	5.15 ± 0.30	6.38 ± 0.12	6.12 ± 0.24					
13	15.45 ± 1.34	20.31 ± 2.12	17.09 ± 1.62					
14	7.56 ± 0.27	6.15 ± 0.23	4.59 ± 0.43					
Erlotinib	17.50 ± 2.34	11.09 ± 0.67	12.86 ± 1.23					
Gefitinib	15.75 ± 1.09	9.60 ± 0.43	10.15 ± 0.39					
Osimertinib	_	5.96 ± 0.63	0.93 ± 0.09					

^a $IC_{50} = Mean \pm SD, n = 3.$

(EGFR^{T790M}) compared to **gefitinib**. Derivative **8**, in particular, exhibited exceptional inhibitory activity, with an IC_{50} value of 9.6 nM, which was 38-fold more potent than **gefitinib**. Furthermore, derivative **8** demonstrated a comparable inhibitory effect against EGFR^{T790M} to **osimertinib**, which may account for its substantial antiproliferative effect on the H1975 cell line.

3. In silico studies

3.1. Molecular docking study

In silico simulation tests were conducted on derivatives **8**, **12**, and **14** at the active ATP binding sites of both EGFR^{WT} and EGFR^{T790M} protein kinases (PDB: 4HJO and PDB: 3W2O, respectively). The protein models were obtained from the PDB website using the MOE 2019.01 program. The results showed that the newly designed compounds docked against the EGFR^{WT} kinase with good binding affinity scores, ranging from -6.40 to -7.10 kcal mol⁻¹ (Table 3). The outcomes of this

Table 2 In vitro enzymatic inhibitory impacts of scaffolds **5–14** versus EGFR^{WT} and EGFR^{T790M}

	IC ₅₀ (nM)		
Comp.	$\mathrm{EGFR}^{\mathrm{WT}}$	$\mathrm{EGFR}^{\mathrm{T790M}}$	
5	49.2 ± 6.7	93.6 ± 8.5	
6	52.6 ± 9.3	66.8 ± 10.4	
7	38.9 ± 8.3	114.6 ± 9.6	
8	23.2 ± 0.1	9.6 ± 3.2	
9	43.5 ± 0.2	17.1 ± 0.1	
10	31.5 ± 5.8	60.8 ± 6.2	
11	47.7 ± 11.4	132.8 ± 16.2	
12	14.5 ± 7.5	35.4 ± 8.6	
13	41.7 ± 8.2	214.9 ± 15.2	
14	15.3 ± 3.1	24.3 ± 3.4	
Gefitinib	18.2 ± 2.2	368.2 ± 25.6	
Osimertinib	57.8 ± 10.2	8.5 ± 2.7	

simulation approach were elucidated secondarily by docking erlotinib and TAK-285 as the co-localized ligands at the active sites of both EGFRWT and EGFRT790M. The root mean square deviation (RMSD) values of erlotinib and TAK-285 secondly docked analogs and co-localized conformers, respectively, were 1.4 and 1.85 Å, respectively, illustrating the rationality of this docking technique (Fig. 1S and 5S, respectively, in ESI†).

To investigate the potential dual EGFR inhibitory activities of gefitinib and osimertinib, these reversible first- and irreversible third-generation EGFR inhibitors were docked against wild-type and mutant EGFR, respectively. Additionally, erlotinib, the colocalized ligand, was re-docked into its corresponding cocrystalized protein model (PDB code: 4HJO) to evaluate the ability of MOE to replicate the native ligand superimposition to the wild-type EGFR protein active site (Fig. 1S and 2S,† and Table 3).

The re-docked erlotinib structure exhibited a high degree of similarity with the native ligand, with an energy score (ΔG) of -6.83 kcal mol⁻¹ and a good RMSD value of 1.4 Å. Particularly, the re-docked erlotinib structure formed one H-bond (hydrogen bond) with Met769 and two H2O-dependent H-bonds with Gln767 and Thr766, highlighting the accuracy of the docking protocol.

Osimertinib and gefitinib, their binding interactions with the active EGFRWT binding site were investigated through molecular docking studies. Osimertinib was found to form one π -H bond with Val702, and one π -cation contact with Lys692 AAs (amino acids), resulting in an energy score (ΔG) of -7.01 kcal mol⁻¹ and 2.38 Å as its RMSD value (Fig. 3S† and Table 3). However, these interactions were not sufficient to inhibit the EGFRWT protein kinase, consistent with previous literature data indicating that osimertinib has a higher affinity

for the mutant EGFR^{T790M} protein kinase and no activity against the EGFRWT.

In contrast, gefitinib was found to form hydrogen bonds with Met769 and H₂O-assisted H-bonds with Thr766 and Gln767 amino acids, similar to erlotinib. Additionally, gefitinib formed two hydrophobic bonds with Leu694 and Leu820 AAs within the EGFR binding site, resulting in an energy score (ΔG) of -8.01 kcal mol⁻¹ and an RMSD value of 1.33 Å (Fig. 4S† and Table 3). These interactions suggest that gefitinib can afford good inhibitory activity within the EGFRWT protein active site, consistent with its use as reference drug in the in vitro experiments.

Derivatives 5, 12, and 14, their binding interactions with the active EGFRWT binding site (PDB code: 4HJO) were investigated. Derivative 5 was able to bind to the EGFRWT active site with a satisfactory interaction affinity, exhibiting an energy score (ΔG) of -6.40 kcal mol⁻¹ and 1.99 Å as its RMSD value. The binding mode of derivative 5 involved one H bond with Lys721 AA and one π -H bond with the major amino acid Met769 (Fig. 4 and Table 3). Derivative 12 demonstrated a good interaction affinity relative to the reference erlotinib, with an energy score (ΔG) of -7.10 kcal mol⁻¹ and 1.05 Å as its RMSD value (Fig. 5 and Table 3). The binding mode of derivative 12 involved hydrogen bonds with Met769, Lys692, Lys704, and H2O-assisted H-bonds with Thr766 and Gln767 amino acids, similar to erlotinib. Additionally, derivative 12 formed two hydrophobic bonds with Val702 amino acid within the EGFR binding site. Derivative 14 exhibited a different binding mode than erlotinib, with an acceptable interaction affinity relative to erlotinib, characterized by an energy score (ΔG) of -6.53 kcal mol⁻¹ and 2.14 Å as its RMSD value (Fig. 6 and Table 3). The binding interaction mode of derivative 14 involved a H-bond with the

Table 3 Molecular docking results for 8, 12, 14, erlotinib, gefitinib, and osimertinib at the main binding site of EGFRWT

Comp.	ΔG score kcal mol ⁻¹	RMSD_refine (Å)	Amino acid/bond	Distance (Å)	E (kcal mol ⁻¹)
8	-6.40	1.99	Met769/pi–H	3.98	-0.5
			Lys721/H acceptor	3.39	-2.5
12	-7.10	1.05	Met769/H acceptor	3.34	-1.5
			Lys692/H acceptor	3.21	-1.4
			Lys704/H acceptor	2.95	-8.3
			Val702/pi–H	4.11	-0.5
			Val702/pi–H	4.24	-0.7
			HOH Thr766/H acceptor	3.17	-0.7
			HOH Gln767/H acceptor	3.17	-0.7
14	-6.53	2.14	Met769/H donor	2.98	-0.6
			Lys721/H acceptor	2.91	-8.1
			Leu694/pi–H	3.91	-0.7
			Val702/pi–H	3.84	-0.8
Erlotinib	-6.83	1.4	Met769/H acceptor	2.95	-4.4
			HOH Thr766/H donor	3.03	-1.6
			HOH Gln767/H donor		-1.1
Gefitinib	-8.01	1.33	Met769/H acceptor	3.03	-3.3
			HOH Thr766/H donor	3.10	-1.5
			HOH Gln767/H donor	3.10	-1.5
			Leu694/pi-H	4.3	-0.8
			Leu820/pi–H	4.56	-0.7
Osimertinib	-7.01	2.38	Lys692/pi–cation	4.38	-0.7
			Val702/pi–H	3.77	-0.7

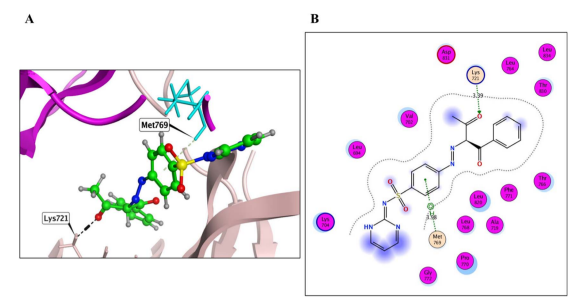


Fig. 4 3D (A) and 2D images (B) of 8 (green sticks) within the wild EGFR kinase.

major amino acid Met769 and another hydrogen bond with Lys721 AA inside the EGFR^{WT} active site. Furthermore, derivative 14 formed two hydrophobic interactions with Leu694 and Val702 AAs.

The binding interactions of derivatives **8**, **12**, and **14** with the EGFR^{T790M} mutant model (PDB code: 3W2O) were investigated and the outcomes illustrated that these derivatives displayed good binding affinities *versus* EGFR^{T790M}, with energy scores ranging from -6.33 to -7.53 kcal mol⁻¹ (Table 4). Relying on previous literature data about the EGFR^{T790M} mutant model, Met790 amino acid in the EGFR^{T790M} mutant model can adopt specific conformations to fit with various inhibitors, while the R858 residue can afford conformational activation loop variants. These outcomes provide a deeper understanding of the SARs that can facilitate the development of more efficient inhibitory agents against sensitive and resistant drugdependent EGFR mutations.⁵¹ Regarding the validation of the

docking protocol of the co-crystallized ligand **TAK-285**, we redocked it into its corresponding EGFR^{T790M} adenine binding pocket of the co-crystal protein model (PDB code: 3W2O). The results showed that the re-docked **TAK-285** binding score (ΔG) was -7.35 kcal mol⁻¹, with a good RMSD value of 3.01 Å (Fig. 5S† and Table 4). The binding mode of **TAK-285** involved one H-bond with each Met793, Asp800, Leu788, and Lys745 amino acids. Additionally, the pyrimidine ring of **TAK-285** formed one H₂O-assisted H-bond with Met790, with 3.01 Å as the bond length value.

The binding interactions of **gefitinib**, **osimertinib**, and **erlotinib** with the EGFR^{T790M} mutant model (PDB code: 3W2O) were investigated as well. **Gefitinib**, utilized as a reference in the *in vitro* experiments, was found to form only one significant hydrogen bond with Met790 AA, with 3.62 Å a distance value and an energy score of -5.98 kcal mol⁻¹ (Fig. 6S† and Table 4). In contrast, **osimertinib** was found to bind to the EGFR^{T790M}

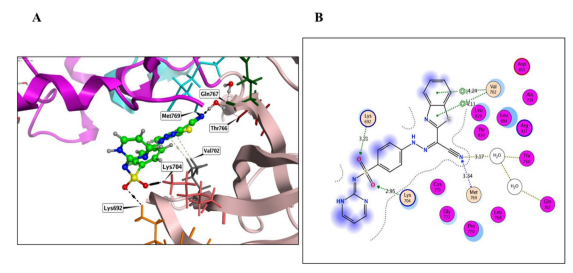


Fig. 5 3D (A) and 2D images (B) of 12 (green sticks) within the wild EGFR kinase.

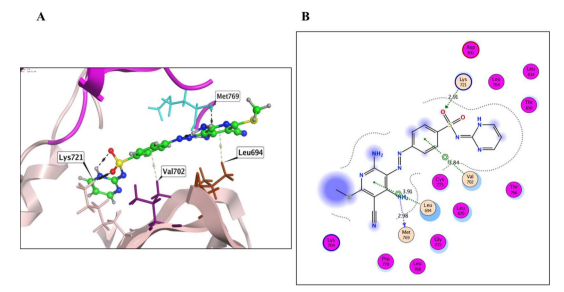


Fig. 6 3D (A) and 2D images (B) of 14 (green sticks) within the wild EGFR kinase.

mutant model in a more potent manner, with a binding score (ΔG) of -8.72 kcal mol⁻¹ and an RMSD value of 1.53 Å (Fig. 7S[†] and Table 4). The binding mode of osimertinib involved two Hbonds with Met790, one H-bond with Glu762, two ionic bonds with Asp855, and three hydrophobic bonds with Leu718 AA within the mutant EGFR^{T790M} binding site. These results suggest that osimertinib can afford potent inhibitory activity within the mutant EGFR^{T790M} protein active site, consistent with its auspicious IC₅₀ value in the in vitro EGFR^{T790M} biological investigation. Erlotinib was also re-docked against the mutant EGFR^{T790M} binding site to evaluate its affinity. The results elucidated that erlotinib made H-bonds with only two non-essential amino acids, Met769 and Lys745, with distances of 3.84 and 3.04 Å, respectively (energy score = $-6.54 \text{ kcal mol}^{-1}$) (Fig. 8S[†] and Table 4). These findings are consistent with literature data, which suggest that erlotinib has

Table 4 Molecular modelling results for 8, 12, 14, gefitinib, Tak-285, erlotinib, and osimertinib at the active binding site of mutant EGFR^{T790M}

	ΔG	RMSD-refine		Distance	E (kcal mol ⁻¹)	
Comp.	score kcal mol ⁻¹	(Å)	Amino acid/bond	(Å)		
8	-7.53	2.03	Met790/H donor	3.56	-0.9	
			Gly796/H acceptor	3.38	-0.5	
			Gly796/H acceptor	3.46	-0.5	
12	-6.62	1.73	Met790/H donor	4.15	-0.5	
			Ile752/H donor	4.30	-0.7	
			Val726/pi–H	4.37	-0.7	
			HOH Met790/H acceptor	2.93	-0.8	
14	-6.33	1.06	HOH Met790/H acceptor	2.79	-0.8	
			Asp855/H donor	2.99	-1.3	
Gefitinib	-5.98	1.25	Met790/H donor	3.62	-0.5	
TAK-285	-7.35	1.85	HOH Met790/H acceptor	3.01	-0.8	
			Met793/H acceptor	3.25	-3.1	
			Asp800/H donor	3.10	-2.3	
			Leu788/H donor	3.39	-1.3	
			Lys745/H acceptor	3.17	-0.8	
Osimertinib	-8.72	1.53	Met790/H donor	3.81	-1.1	
			HOH Met790/H acceptor	3.31	-0.9	
			Glu762/H donor	3.24	-0.7	
			Asp855/ionic	3.90	-0.7	
			Asp855/ionic	3.81	-0.9	
			Leu718/pi–H	4.39	-0.5	
			Leu718/pi–H	4.34	-0.5	
			Leu718/pi–H	3.96	-1.0	
Erlotinib	-6.54	2.06	Met769/H donor	3.84	-0.5	
			Lys745/H acceptor	3.04	-4.9	

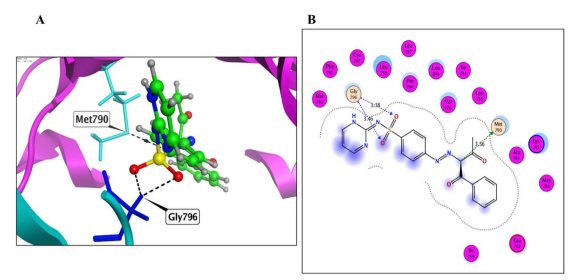


Fig. 7 3D (A) and 2D images (B) of 8 (green sticks) within the mutant EGFR^{T790M} kinase.

affinity only for the wild-type EGFR protein tyrosine kinase, not the mutant EGFR $^{\rm T790M~50}.$

Derivative **8** was found to bind to the active EGFR^{T790M} site *via* three main H-bonds: one with Met790 and two with Gly796 AAs, with distance values of 3.56, 3.38, and 3.46 Å, respectively (energy score = -7.53 kcal mol⁻¹) (Fig. 7 and Table 4). This binding mode suggests that derivative **8** has a strong affinity for the EGFR^{T790M} mutant model. Derivative **12** was found to bind to the EGFR^{T790M} mutant model in a similar manner to **TAK-285**, with a binding mechanism that involved hydrogen bonding with only two essential amino acids, Met790 and Ile752, with distance values of 4.15 and 4.30 Å (energy score = -6.62 kcal mol⁻¹) (Fig. 8 and Table 4). Additionally, derivative **12** formed one hydrophobic interaction with Val726. Derivative **14** was found to bind to the EGFR^{T790M} mutant model *via* two fundamental H-bonds with Met790 and Asp855, with distance

values of 2.79 and 2.99 Å, respectively (energy score = $-5.51 \text{ kcal mol}^{-1}$) (Fig. 9 and Table 4). These binding modes highlight the significance of the hydrogen bond donating and accepting moieties in the designed ligands to act as efficient dual EGFR^{WT} and EGFR^{T790M} inhibitors. The results suggest that the designed ligands have the potential to dually inhibit EGFR^{WT} and EGFR^{T790M} protein tyrosine kinases, which is a critical aspect of cancer therapy.

3.2. Structure-activity relationship (SAR) study

Our rationalized drug design approach involved a comprehensive study of SAR of the target derivatives as potential antiproliferative agents. Specifically, we investigated the impact of R_1 and R_2 substituents on the biological activity of the scaffolds (Fig. 10). The results of our study revealed that compounds **8**, **12**, and **14** exhibited highly potent IC_{50} values, indicating significant *in vitro*

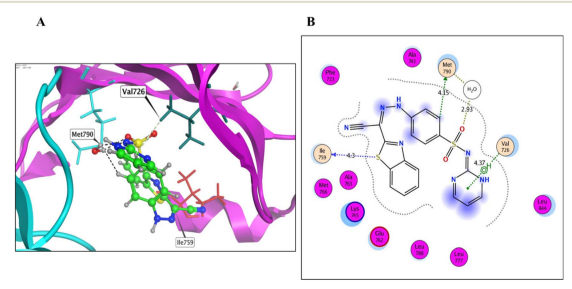
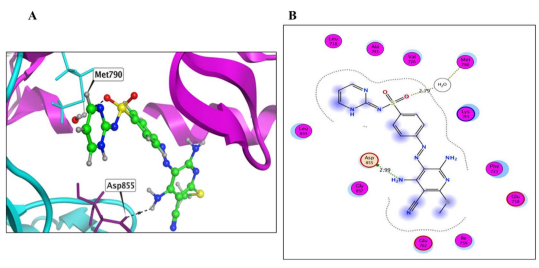


Fig. 8 3D (A) and 2D images (B) of 12 (green sticks) within the mutant EGFR^{T790M} kinase.



3D (A) and 2D images (B) of 14 (green sticks) within the mutant EGFR^{T790M} kinase.

antiproliferative activities and dual EGFRWT and EGFRT790M inhibitory activity. Furthermore, molecular docking studies demonstrated that these compounds formed strong interactions with the target protein, which correlated with their high biological activity. A key finding of our SAR analysis was the importance of hydrogen bond acceptor (HBA) functional groups in achieving effective dual EGFR inhibition. The presence of HBA groups in compounds 8, 12, and 14 was found to be crucial for their

outstanding potency and selectivity towards both EGFRWT and EGFR^{T790M}. These results afford valuable information that greatly helps in the design of novel EGFRIs and illustrate the importance of careful consideration of the SAR of target compounds in the production of promising cancer therapies.

Lapatinib has been shown to interact with the inactive tyrosine kinase domain of EGFR due to its large, sterically hindered [(3-fluorobenzyl)oxy] core scaffold on the peripheral aniline

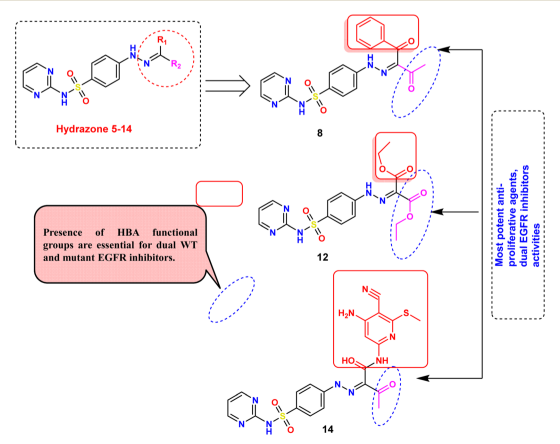


Fig. 10 SAR study of sulfadiazine-based hydrazones 8, 12, and 14.

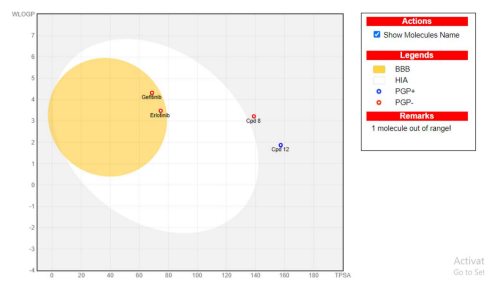


Fig. 11 The boiled-egg diagram for 8, 12, 14, erlotinib, and gefitinib

residue. This prevents it from fitting into the small-sized ATPbinding site, instead allowing it to reach and fit into the allosteric-binding site that is re-opened by the dislocation of the αC helix in the inactive conformation model.⁵² Similarly, the large-sized R₁ and R₂ side chains in compounds 8, 12, and 14, particularly in compound 12, prevent them from fitting into the small ATP-binding pocket. As a result, compound 12 selectively interacts with the inactive EGFR conformation, but not the active one. Compound 12 was detected to be the most active inhibitor of both EGFRWT and EGFRT790M kinases among all tested derivatives. This is attributed to its binding mode, dimensions, and stable docking scores, which are superior to those of the reference inhibitors erlotinib and gefitinib for wild-type EGFR, and TAK-285 and osimertinib for mutant EGFR^{T790M}. These outcomes are coherent with the in vitro inhibition of compound 12 against both EGFRWT and EGFRT790M protein kinases, demonstrating its potential as a potent and selective EGFR inhibitor.

3.3. Pharmacokinetic characteristics and ADMET predict

To further investigate the potential of the synthesized scaffolds as orally bioavailable candidates,⁵² their absorption, distribution, metabolism, and excretion (ADME) properties were evaluated using the ADME Swiss prediction free online tool.^{51–56} The boiled egg diagram of the WLOGP *versus* TPSA (Topological Polar Surface Area) of the uploaded derivatives 8 and 12 revealed that both compounds have low GI absorption and are unable to pass through the BBB relative to the reference

compounds erlotinib and gefitinib (Fig. 11). This suggests that these compounds are unlikely to cause central nervous system (CNS) negative impacts due to their low BBB permeability. In contrast, scaffold 14, a highly polar compound, was found to be out of range and vanished from the boiled egg diagram, highlighting the need for optimization to make it an effective orally bioavailable candidate. The ADME Swiss prediction tool also suggested that derivative 8 is unlikely to be a P-glycoprotein (PGP) substrate, which means that it may not be exposed to drug resistance mechanisms. However, scaffold 12 is predicted to be a PGP substrate, indicating a possibility of exposure to drug resistance mechanisms. The TPSA values of the designed scaffolds 8, 12, and 14 ranged from 138 to 216.04 Å², while their log P values were between 1.85 and 3.93 (Table 5). These values provide further insight into the lipophilicity and bio-molecular polarity of the designed scaffolds.

The bioavailability diagram for scaffolds 8, 12, and 14 is displayed in Fig. 12(A-E). This radar plot consists of six axes representing six critical oral bioavailability characteristics: solubility (INSOLU), flexibility (FLEX), size (SIZE), lipophilicity (LIPO), saturation (INSATU), and polarity (POLAR) (Table S1†). The pink area in the radar plot represents the optimum property parameters for oral bioavailability. The red colored lines represent scaffolds 8 (Fig. 12A), 12 (Fig. 12B), and 14 (Fig. 12C), which are nearly within the pink area relative to erlotinib (Fig. 12D) and gefitinib (Fig. 12E). This suggests that the designed scaffolds have acceptable assumed oral

Table 5 ADMET outcomes of 8, 12, 14, and erlotinib and gefitinib via Swiss ADME webserver

Comp.	M. wt	Fraction Csp ³	RB	НВА	HBD	MR	TPSA	$\operatorname{Log} P$	GI absorption	BBB perm.	Lip. V.	Bio. Sc.
8	423.45	0.05	8	7	2	111.67	138.86	3.22	Low	No	Yes	0.55
12	435.48	0.00	6	7	2	114.36	169.64	3.93	Low	No	Yes	0.55
14	441.49	0.06	6	8	4	119.04	216.04	1.85	Low	No	Yes	0.55
Erlotinib	393.44	0.27	10	6	1	111.40	74.73	3.07	High	Yes	Yes	0.55
Gefitinib	446.90	0.36	8	7	1	121.66	68.74	4.32	High	Yes	Yes	0.55

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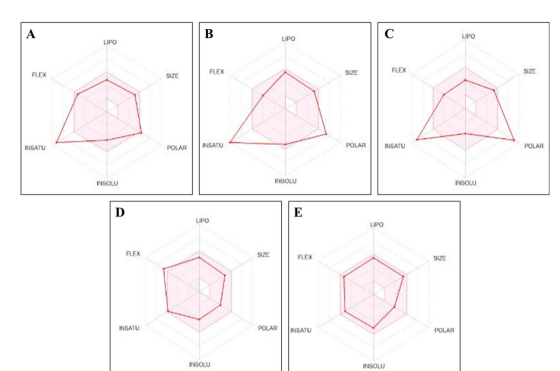


Fig. 12 Pink zoned oral bioavailability radars for 8 (A), 12 (B), 14 (C), erlotinib (D), and gefitinib (E)

bioavailability. Scaffolds 8, 12, and 14 are predicted to afford a suitable bioavailability with a score value equal to 0.55, similar to the reference compounds erlotinib and gefitinib. Additionally, scaffolds 8 and 12 follow the Lipinski rule of 5 with no detected violations, similar to the reference drugs. However, derivative 14 obeys the Lipinski rule of 5 with only one detected violation (number of polar atoms; nitrogen or oxygen atoms higher than 10) compared to the reference compounds (Table S1†). These results demonstrate that the designed compounds have promising pharmacokinetic and drug-like properties. As shown in Fig. 12(A-E) and Table 5, analogs 8 and 12 have good lipophilic characters compared to the reference compounds, unlike compound 14, which has lower lipophilicity. The unsaturation property of scaffolds 8, 12, and 14 is nearly out of the tinted area, indicating that these compounds may require optimization to improve their saturation characteristics. The fraction Csp³ values of scaffolds 8, 12, and 14 were calculated at 0.06, indicating that these compounds have a low ratio of sp³ hybridized carbons over all the carbon numbers. For saturation character, the ratio of sp³ hybridized carbons over all the carbon numbers of the analog (fraction Csp³) should not be lower than 0.25.

Hydrazones **8**, **12**, and **14** exhibited low saturation character, with fraction Csp³ values of 0.05, 0.00, and 0.06, respectively. Conversely, their hydrophilicity ranged from moderately soluble to very soluble, relative to the reference compounds **erlotinib** and **gefitinib**, which had solubility ranging from moderately soluble to poorly soluble. Unlike **erlotinib** and **gefitinib**, which showed inhibitory activity against CYP2C9, CYP2C19, CYP2D6, and CYP3A4, hydrazones **8**, **12**, and **14** did not exhibit inhibitory activity against CYP1A2 and CYP2D6. Additionally, scaffold **14** did not show inhibitory activity against CYP2C9 and CYP2C19,

unlike derivatives 8 and 12. According to PAINS and Brenk filters, hydrazones 8, 12, and 14 are suitable for biological experiments, exhibiting potent chemical reactivity, low hazardous doses, and adequate metabolic stability. Scaffolds 8, 12, and 14 demonstrated lead-likeness, unlike erlotinib and gefitinib, and their synthetic procedures were categorized as accessible. Compounds 8, 12, and 14 had better safety profiles than the reference compounds erlotinib and gefitinib (Table S1†). These compounds are classified as non-AMES toxic, hERG I inhibitors, and skin sensitization agents. Hydrazones 8, 12, and 14 exhibited hepatotoxicity and parameters indicating carcinogenic agents, which may cause severe hepatic side effects, similar to the reference compounds erlotinib and gefitinib. Compounds 8 and 12 demonstrated hERG II K⁺ channel inhibitory activity, similar to the reference compounds erlotinib and gefitinib. In contrast, scaffold 14 did not exhibit hERG II K+ channel inhibitory activity (Table S1†).

4. Experimental part

4.1. General chemistry

All general information were provided in the ESI.†

4.2. Synthesis of sulfadiazine 4

A mixture of an equimolar amount of 4-acetamidobenzene sulfonyl chloride (1) with 2-aminopyrimidine (2) was refluxed in pyridine for 1 hour, then the mixture was added to 5 mL of $\rm H_2O/HCl$ in ratio 2:1, then filtered the mixture, washed with $\rm H_2O$ several times, dried, and recrystallized from EtOH, m.p. 251–253 °C (reported m.p. 252–256 °C).³⁷

4.3. Synthesis of sulfadiazine 5-14

A solution of sodium nitrite (0.1 g, 1.3 mmol) in water (2 mL) was gradually added to a solution of sulfadiazine (0.25 g, 1 mmol) in conc. HCl (3 mL) at 0–5 °C. The formed diazonium was mixed under stirring with an ice-cooled solution of appropriate coupling compounds (1 mmol), namely, ethyl acetoacetate, acetylacetone, ethyl benzoylacetate, benzoyl acetone, ethyl cyanoacetate, cyanothioacetamide, dimer of malononitrile, diethyl malonate, *N*-[4-amino-5-cyano-6-(methylthio)pyridin-2-yl]-3-oxobutanamide and 4,6-diamino-2-(methylthio)nicotinonitrile in pyridine (20 mL). The mixture was kept for 30 minutes in an ice bath and then mixed into 100 mL of cold water. The product was filtered off, washed with distilled water, dried, and recrystallized from EtOH.

4.3.1. 4-(2-(2,4-Dioxohexan-3-ylidene)hydrazineyl)-*N*-(pyrimidin-2-yl)benzenesulfonamide (5).

Brownish yellow solid, yield (91%), m.p. 160–162 °C. IR (KBr) ν : 3404, 3256 (2NH), 3089, 3039 (C–H Aromatic), 2974, 2938, 2874, 2811, 2734 (C–H aliphatic), 1690 (br, C=O), 1584 (C=N); ¹H NMR δ: 1.25–1.29 (t, 3H, J=8 Hz, CH_{3 ester}), 2.41 (s, 3H, CH₃), 4.29–4.34 (q, 2H, J=8 Hz, CH₂), 7.03–7.06 (t, 1H, J=6 Hz, CH_{arom}), 7.54–7.56 (d, 2H, J=8 Hz, CH_{arom}), 7.97–8.00 (d, 2H, J=12 Hz, CH_{arom}), 8.50–8.51 (d, 2H, J=4 Hz, CH_{arom}), 11.59 (s, 1H, NH, D₂O exchangeable), 11.76 (br, 1H, NH, D₂O exchangeable); 13°C NMR δ: 14.28 (CH_{3 ester}), 25.76 (CH₃), 61.84 (CH₂), 115.06, 116.21, 129.97, 134.22, 134.62, 146.55, 157.44, 158.75, 162.60 (C=O), 194.31 (C=O); anal. calcd for C₁₆H₁₇N₅O₅S (391.40): C, 49.10; H, 4.38; N, 17.89. Found: C, 49.22; H, 4.26; N, 17.75.

4.3.2. 4-(2-(2,4-Dioxopentan-3-ylidene)hydrazineyl)-*N*-(pyrimidin-2-yl)benzenesulfonamide (6).

Yellow solid, yield (80%), m.p. 228–230 °C. IR ν : 3422, 3312 (2NH), 3082, 3033 (C–H aromatic), 2936, 2865, 2808, 2729 (C–H aliphatic), 1677 (C=O), 1585 (C=N); 1 H NMR δ 2.42 (s, 3H, CH₃), 2.47 (s, 3H, CH₃), 7.04–7.05 (m, 1H, CH_{arom}), 7.68–7.70 (d, 2H, J = 8 Hz, CH_{arom}), 8.00–8.02 (d, 2H, J = 8 Hz, CH_{arom}), 8.50 (s, 2H, CH_{arom}), 11.70 (br, 1H, NH, D₂O exchangeable), 13.51 (s, 1H, NH, D₂O exchangeable); 13 C NMR δ : 26.75 (CH₃), 31.70 (CH₃), 116.25, 129.88, 135.83, 135.95, 145.95, 157.34, 158.83, 159.04, 196.92 (C=O), 197.92 (C=O); anal. calcd for C₁₅H₁₅N₅O₄S (361.37): C, 49.85; H, 4.18; N, 19.38. Found: C, 49.73; H, 4.27; N, 19.51.

4.3.3. Ethyl-3-benzoyl-2-(4-(*N*-(pyrimidin-2-yl)sulfa) hydrazineylidene)propanoate (7).

Yellow solid, yield (76%), m.p. 205–207 °C; IR ν : 3224, 3172 (2NH), 3083, 3039 (C–H aromatic), 2984, 2942, 2874, 2813, 2738 (C–H aliphatic), 1684, 1662 (C=O), 1582 (C=N); ¹H NMR δ : 1.24–1.27 (t, 3H, J = 6 Hz, CH₃), 4.32–4.37 (q, 2H, J = 6 Hz, CH₂), 7.01–7.04 (m, 1H, CH_{arom}), 7.39–7.41 (d, 2H, J = 8 Hz, CH_{arom}), 7.53–7.60 (m, 3H, CH_{arom}), 7.65–7.72 (m, 1H, CH_{arom}), 7.91–7.98 (m, 3H, CH_{arom}), 7.49–8.51 (m, 2H, CH_{arom}), 11.66 (br, 1H, NH), 11.99 (br, 1H, NH); ¹³C NMR δ : 14.27 (CH₃ ester), 61.91 (CH₂), 115.12, 116.20, 128.28, 129.56, 129.97, 130.37, 133.43, 134.73, 137.04, 146.44, 157.41, 158.76, 162.43 (C=O), 192.49 (C=O); anal. calcd for C₂₁H₁₉N₅O₅S (453.47): C, 55.62; H, 4.22; N, 15.44. Found: C, 55.47; H, 4.38; N, 15.32.

4.3.4. 4-[2-(1-Benzoyl-2-oxopropylidene)hydrazino]-*N*-pyrimidin-2-ylbenzenesulfonamide (8).

Orange solid, yield (75%), m.p. 235–237 °C. IR ν : 3268, 3113 (2NH), 3084, 3038 (C–H aromatic), 2943, 2873, 2813, 2739 (C–H aliphatic), 1655 (C=O), 1582 (C=N); ¹H NMR δ 2.51 (s, 3H, CH₃), 7.03–7.05 (t, 1H, J=4 Hz, CH_{arom}), 7.50–7.58 (m, 4H, CH_{arom}), 7.69–7.72 (t, 1H, J=6 Hz, CH_{arom}), 7.80–7.82 (d, 2H, J=8 Hz, CH_{arom}), 7.95–7.97 (d, 2H, J=8 Hz, CH_{arom}), 8.50–8.51 (d, 2H, J=4 Hz, CH_{arom}), 11.34 (s, 1H, NH), 13.65 (s, 1H, NH); ¹³C NMR δ : 25.39 (CH₃), 114.58, 116.22, 129.21, 129.54, 129.94, 133.84, 134.94, 135.85, 141.45, 147.22, 157.22, 158.77, 195.22 (C=O), 196.65 (C=O); anal. calcd for C₂₀H₁₇N₅O₄S (423.44): C, 56.73; H, 4.05; N, 16.54. Found: C, 56.57; H, 4.21; N, 16.47.

4.3.5. Ethyl (Z)-2-cyano-2-(2-(4-(*N*-(pyrimidin-2-yl) sulfamoyl)phenyl)hydrazineylidene)acetate (9).

Yellow solid, yield (78%), m.p. 225–227 °C; IR ν : 3387, 3218 (2NH), 3036, 3032 (C–H aromatic), 2972, 2935, 2866, 2806 (C–H aliphatic), 2214 (C \equiv N), 1725 (C \equiv O), 1582 (C \equiv N); ¹H NMR δ : 1.27–1.30 (t, 3H, J=6 Hz, C $\underline{\rm H}_3$), 4.27–4.32 (q, 2H, J=6 Hz, C $\underline{\rm H}_2$), 7.03–7.06 (t, 1H, J=6 Hz, C $\underline{\rm H}_{\rm arom}$), 7.59–7.61 (d, 2H, J=8 Hz, CH $_{\rm arom}$), 8.01–8.03 (d, 2H, J=8 Hz, C $\underline{\rm H}_{\rm arom}$), 8.50–8.51 (d, 2H, J=4 Hz, C $\underline{\rm H}_{\rm arom}$), 11.84 (br, 1H, N $\underline{\rm H}$, D $_2$ O exchangeable) 12.45 (br,

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1H, NH, D_2O exchangeable); ¹³C NMR δ : 14.35 (CH₃), 62.29 (CH₂), 106.78 (C≡N), 111.49, 116.20, 116.64, 129.95, 135.97, 145.81,

157.31, 158.81, 160.89 (C=O); anal. calcd for C₁₅H₁₄N₆O₄S (374.37): C, 48.12; H, 3.77; N, 22.45. Found: C, 48.28; H, 3.63; N, 22.57.

4.3.6. 2-Cyano-(2-(4-(N-(pyrimidin-2-yl)sulfamoyl)phenyl) hydrazono)ethanethioamide (10).

Reddish brown solid, yield (69%), m.p. 275-277 °C. IR ν: 3360, 3228, 3188 (NH + NH₂), 3041 (C-H aromatic), 2211 (C \equiv N), 1581 (C=N); ¹H NMR δ : 7.06–7.08 (t, 1H, J = 4 Hz, CH_{arom}), 7.86–7.88 $(d, 2H, J = 8 Hz, CH_{arom}), 7.93-7.95 (d, 2H, J = 8 Hz, CH_{arom}),$ 8.51-8.52 (d, 2H, J = 4 Hz, $C\underline{H}_{arom}$), 9.50 (s, 1H, $N\underline{H}_{2}$, $D_{2}O$ exchangeable), 9.87 (s, 1H, NH₂, D₂O exchangeable), 11.82 (br, 1H, NH, D₂O exchangeable); ¹³C NMR δ : 111.30 (C \equiv N), 115.28, 116.31, 116.52, 129.49, 135.25, 146.12, 157.36, 158.86, 188.03 (C=S); anal. calcd for $C_{13}H_{11}N_7O_2S_2$ (361.40): C, 43.20; H, 3.07; N, 27.13. Found: C, 43.32; H, 3.19; N, 27.02.

4.3.7. 2-Amino-3,3-dicyano-*N*-(4-(*N*-(pyrimidin-2-yl) sulfamoyl)phenyl)acrylohydrazonoyl cyanide (11).

Yellow solid, yield (77%), m.p. 295-297 °C; IR v: 3370, 3303, 3214 $(NH + NH_2)$, 3088, 3050 (C-H aromatic), 2215 (br, C=N), 1634 (C=N); ¹H NMR δ : 7.06–7.09 (t, 1H, J = 4 Hz, C \underline{H}_{arom}), 7.53 (br, 1H, N<u>H</u>), 7.76–7.78 (d, 2H, J = 8 Hz, C<u>H</u>_{arom}), 8.22–8.24 (d, 2H, J =8 Hz, C_{Harom} , 8.54–8.55 (d, 2H, J = 4 Hz, C_{Harom}), 10.04 (br, 3H, NH + NH₂); ¹³C NMR δ: 79.43 (C-(CN)₂), 112.91 (C \equiv N), 114.48 $(C \equiv N)$, 116.02, 117.29 $(C \equiv N)$, 127.46, 129.05, 141.51, 144.12, 148.59, 152.58, 157.35, 158.78; anal. calcd for C₁₆H₁₁N₉O₂S (393.38): C, 48.85; H, 2.82; N, 32.05. Found: C, 48.72; H, 2.67; N, 32.22.

4.3.8. Diethyl 2-(2-(4-(N-(pyrimidin-2-yl)sulfamoyl)phenyl) hydrazineylidene)malonate (12).

Brownish red solid, yield (72%), m.p. 148-150 °C. IR ν: 3225, 3103 (2NH), 3077, 3034 (C-H aromatic), 2981, 2934, 2870, 2806, 2732 (C-H aliphatic), 1722, 1681 (C=O), 1578 (C=N); ¹H NMR δ : 1.25–1.30 (m, 6H, 2CH₃), 4.23–4.35 (qq, 4H, J = 6.6 Hz, 2CH₂), 7.02–7.04 (t, 1H, J = 4 Hz, $C\underline{H}_{arom}$), 7.50–7.52 (d, 2H, J = 8 Hz, CH_{arom}), 7.98-8.00 (d, 2H, J = 8 Hz, CH_{arom}), 8.50-8.51 (d, 2H, J $= 4 \text{ Hz}, \text{CH}_{arom}$, 11.71 (s, 1H, N<u>H</u>), 11.89 (s, 1H, N<u>H</u>); ¹³C NMR δ: 14.33 (CH₃), 14.46 (CH₃), 61.54 (CH₂), 62.01 (CH₂), 115.06, 116.20, 125.15, 129.98, 134.75, 146.34, 157.40, 158.75, 161.69 (C=O), 162.52 (C=O); anal. calcd for $C_{17}H_{19}N_5O_6S$ (421.42): C, 48.45; H, 4.54; N, 16.62. Found: C, 48.59; H, 4.41; N, 16.53.

4.3.9. 4-{-2-[4-Amino-5-cyano-2-imino-6-(methylthio)pyridin-3(2H)-ylidene]hydrazino}-N-pyrimidin-2ylbenzenesulfonamide (13).

Orange solid, yield (80%), m.p. 298–300 °C. IR v: 3457, 3302, 3205 (NH + NH₂), 3075, 3027 (C-H aromatic), 2935, 2867, 2812, 2734 (C-H aliphatic), 2215 (C \equiv N), 1574 (C \equiv N); ¹H δ 2.57 (s, 3H, $C\underline{H}_3$), 7.08 (s, 1H, $C\underline{H}_{arom}$), 7.03–7.19 (m, 6H, $4C\underline{H}_{arom} + N\underline{H}_2$ D_2O exchangeable), 8.52–8.53 (m, 3H, 2CH_{arom}+ N<u>H</u>, D_2O exchangeable), 9.19 (br, 1H, NH), 11.91 (br, 1H, NH, D₂O exchangeable), 13.37 (s, 1H, NH, D₂O exchangeable); 13 C NMR δ : 12.87 (CH₃), 79.92 (C-CN), 114.59, 116.16 (2C), 122.51, 129.08, 139.92, 153.79, 154.48, 155.11, 157.32, 158.88, 168.04; anal. calcd for C₁₇H₁₅N₉O₂S₂ (441.49): C, 46.25; H, 3.42; N, 28.55. Found: C, 46.38; H, 3.54; N, 28.41.

4.3.10. N-(4-Amino-5-cyano-6-(methylthio)pyridin-2-yl)-3oxo-2-{[4-(pyrimidin-2-ylsulfamoyl) phenyl]hydrazono}butanamide (14).

Yellow solid, yield (83%), m.p. > 300 °C, IR v: 3415, 3386, 3341 (NH + NH₂), 3039 (C-H aromatic), 2930, 2871, 2816, 2734 (C-H aliphatic), 2208 ($C \equiv N$), 1668 (br, C = O), 1582 (C = N); ¹H NMR δ : 2.52 (s, 3H, C<u>H</u>₃), 7.06–7.09 (t, 1H, J = 4 Hz, C<u>H</u>_{arom}), 7.19 (br, 2H, N \underline{H}_2), 7.38 (s, 1H, C $\underline{H}_{pyridyl}$), 7.65–7.67 (d, 2H, J=8 Hz, $C\underline{H}_{arom}$), 8.01–8.03 (d, 2H, J = 8 Hz, $C\underline{H}_{arom}$), 8.52–8.53 (d, 2H, J $= 4 \text{ Hz}, C_{\text{Harom}}$, 11.38 (s, 1H, NH), 11.82 (br, 1H, NH), 13.37 (s, 1H, NH); anal. calcd for C₂₁H₁₉N₉O₄S₂ (525.56): C, 47.99; H, 3.64; N, 23.99. Found: C, 48.11; H, 3.51; N, 23.83.

4.4. Biological evaluation

4.4.1. In vitro anti-proliferative activities. All procedures were provided in the ESI.†

4.4.2. EGFR^{WT} and EGFR^{T790M} kinase inhibitory assay. All procedures were provided in the ESI.†

4.5. Docking studies and ADMET estimation

All procedures were provided in the ESI.†57-62

Conclusion

A new series of sulfadiazine (5-14) was effectively synthesized via diazotization of sulfadiazine with some active methylene compounds. The cytotoxic activity of all products was investigated against A431, A549, and H1975 cancer cell panels via the MTT assay technique, utilizing erlotinib, gefitinib, and osimertinib as positive controls. Specifically, compounds 8, 12, and 14 were revealed to be the most potent analogs on A431, A549, and H1975 cancer cell panels with the IC_{50} values of (4.12, 4.78, and 2.31 μ M), (6.38, 5.15, and 6.12 μ M), and (6.15, 7.56, and 4.59 μ M) compared to **erlotinib** (IC₅₀ = 11.09, 17.50, and 12.86 μ M), gefitinib (IC₅₀ = 9.60, 15.75, and 10.15 μ M), and osimertinib (IC₅₀ = 5.96, NT, and 0.93 μ M), respectively. Moreover, compound 12 was the most potent tested one versus both $\mathrm{EGFR}^{\mathrm{WT}}$ and $\mathrm{EGFR}^{\mathrm{T790M}}$ protein kinases with IC_{50} values of 14.5 and 35.4 nM, respectively, compared to gefitinib and osimertinib as reference drugs with IC50 values of 18.2 and 368.2 nM and 57.8 and 8.5 nM, respectively. The moleculardocking results of scaffolds 8, 12, and 14 inside both the EGFRWT and EGFRT790M ATP active sites elucidated their ability to afford the critical fitting interactions made within the kinase domain by the references erlotinib, gefitinib, and osimertinib, respectively. Furthermore, investigating the physicochemical characteristics of the most outstanding EGFR inhibitors, compounds 8, 12, and 14, through the Swiss ADME web server illustrated good GIT absorption, lipophilicity, and low BBB penetration characteristics. The valuable outcomes from this simulation study are that scaffolds 8 and 12 are assumed to be the most promising analogs regarding their dual EGFRWT and EGFR^{T790M} inhibitory activity, as well as promising antiproliferative lead compounds with good pharmacokinetic properties. These findings suggest that these scaffolds should be further investigated and optimized for more potent and selective antiproliferative activity in the future.

Data availability

All data generated or analyzed during this study are included in this published article (and its ESI†).

Conflicts of interest

The authors declare no conflict of interest.

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