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Design, synthesis, docking, ADMET and anticancer evaluations of *N*-alkyl substituted iodoquinazoline derivatives as dual VEGFR-2 and EGFR inhibitors†

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Fifteen new 1-alkyl-6-iodoquinazoline derivatives **5a–d** to **9a–e** were designed and synthesized and their anticancer activities were evaluated against HepG2, MCF-7, HCT116 and A549 cancer cell lines via dual targeting of EGFR and VEGFR-2. The newly synthesized compounds were designed based on the structure requirements of the target receptors and were confirmed using spectral data. Compound **9c** showed the highest anticancer activities with $EC_{50} = 5.00, 6.00, 5.17$ and $5.25 \mu\text{M}$ against HepG2, MCF-7, HCT116 and A549 cell lines correspondingly. Moreover, compounds **5d**, **8b**, **9a**, **9b**, **9d**, and **9e** exhibited very good anticancer effects against the tested cancer cell lines. The highly effective seven derivatives **5d**, **8b**, **9a–e** were examined against VERO normal cell lines to estimate their cytotoxic capabilities. Compounds **9c**, **9b**, **9d**, **9a**, **9e** and **5d** excellently inhibited VEGFR-2 activity with $IC_{50} = 0.85, 0.90, 0.90, 1.00, 1.20$ and $1.25 \mu\text{M}$ respectively. Moreover, compounds **9c**, **9d**, **9e**, **5d**, **8b** and **9b** excellently inhibited EGFR^{T790M} activity with $IC_{50} = 0.22, 0.26, 0.30, 0.40, 0.45$ and $0.50 \mu\text{M}$ respectively. Also, compounds **9c**, **9d** and **9e** excellently inhibited EGFR^{WT} activity with $IC_{50} = 0.15, 0.20$ and $0.25 \mu\text{M}$ respectively. As planned, compound **9c** showed excellent dual EGFR/VEGFR-2 inhibitory activities. Consonantly, ADMET study was calculated *in silico* for the supreme three worthwhile compounds **9b**, **9c** and **9e** in contrast to sorafenib and erlotinib as reference drugs. The obtained results concluded that, our compounds might be useful as prototype for design, optimization, adaptation and investigation to have more powerful and selective dual VEGFR-2/EGFR^{T790M} inhibitors with higher antitumor activity.

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1. Introduction

The prevention of tumor progression has involved blocking angiogenesis as a prevalent strategy. Vascular endothelial growth factor receptor (VEGFR) shows a fundamental role in VEGF-induced response regulation in endothelial cells and

increased microvascular permeability in solid tumors.¹ VEGFRs bind to VEGF and other related ligands, starting a cascade of signals leading to the proliferation and endothelial cell migration and new blood vessel development.² Also, in both normal and abnormal angiogenesis, VEGFR is a main component, making it a target for the development of new cancer

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treatments.³ VEGFR has three subtypes, which are VEGFR-1, VEGFR-2 and VEGFR-3. The most important for angiogenesis is the VEGFR-2.⁴ In various cancers such as breast, lung, liver and colorectal cancer, VEGFR-2 is overexpressed. So, blocking VEGFR-2 with suitable tyrosine kinase inhibitors is the most significant target in anti-angiogenic therapy.^{5,6}

Existing research is being conducted to improve new strategies targeting signaling pathways.^{7,8} Tyrosine kinases modulate the growth factors signaling so they are considered as a significant target in cancer therapy. One of the most popular targets in anticancer therapy is protein tyrosine kinases (PTKs).⁹ Targeted therapy showed fewer toxic side effects compared to chemotherapy as they aimed at cancer-specific molecules and signaling pathways.¹⁰

VEGFR-2 inhibitors can be divided into three classes: class I, which binds to the ATP binding region (hinge region). Class II binds to the ATP binding site and extends into the adjacent hydrophobic back pocket, showing higher affinity and selectivity compared to class I; and class III blocks the receptor through hydrophobic interactions in the allosteric hydrophobic back pocket. Due to their prolonged TK suppression, researchers are actively seeking to develop new type II and III VEGFR-2 inhibitors.^{11–13}

There are several quinazoline derivatives were reported as antitumor agents targeting VEGFR-2 kinase inhibitors.^{14,15} Alternatively, several potential anticancer agents were designed with numerous quinazoline scaffolds.^{16–18} Also, secondary resistance to approved VEGFR-2 inhibitors after the initial course of treatment highlights the need for novel therapeutic alternatives.^{19,20}

On the other hand, VEGF and epidermal growth factor (EGF) act through similar signaling pathways and can act independently of each other in cancer development and resistance to treatment.²¹ EGFR (epidermal growth factor receptor) is the most common important PTK in cellular proliferation, differentiation and signaling pathways.²² Multitarget agents are involved in signaling pathways preventing cancer progression by targeting different biological receptors resulting in reduced toxicity synergistic effects.²³ Thus, the simultaneous VEGFR and EGFR signaling pathways blockade proved to be an attractive approach to cancer treatment.¹⁴ Research suggests that tumors with EGFR mutations are more reliant on VEGF than those with the normal EGFR gene.²⁴ As a result, targeting both VEGF and EGFR simultaneously is considered a rational approach for treating EGFR-mutant non-small cell lung cancer (NSCLC) treatment.²⁵

EGFR mutation is a common mechanism of resistance to tyrosine kinase inhibitors (TKIs) involved in cancer progression.²⁶ In 90% of EGFR-mutant NSCLC cases, an exon 19 deletion or exon 21 L858R mutations occur, both of which render tumors responsive to EGFR TKIs.^{27,28} Also T790M, known as Thr790Met, is a gatekeeper mutation of EGFR. The mutation substitutes a threonine (T) with a methionine (M) at position 790 of exon 20. The EGFR^{T790M} mutation restores the affinity of the kinase for ATP to wild-type levels, and therefore at cellular ATP levels, the potency of the TKI is reduced.²⁹ In metastatic cases, EGFR TKIs are considered first-line therapy due to their excellent tolerability and progression-free survival (PFS).^{30,31}

Despite being initially effective, resistance to EGFR TKI therapy consequently occurs. The survival rate for individuals with metastatic lung cancer caused by a mutant EGFR gene is only 15% over five years, highlighting the need for new and improved therapeutic approaches. One possible addition to EGFR inhibition in non-small cell lung cancer is targeting the VEGF pathway.^{32,33} The suppression of the VEGF signaling pathway is a crucial step in cancer treatment.³⁴ Lately, there is considerable attention paid to designating dual TKIs that can prevent cancer development.^{35,36}

Numerous FDA approved quinazoline based small molecule inhibitors display clinical importance *via* targeting either VEGFR-2 or EGFR signaling pathways and inhibiting tumor progression.^{37,38}

Quinazoline is a good scaffold for drug discovery. In addition to anticancer activities targeting different biological targets, they also show good potential as antiviral³⁹ and antibacterial⁴⁰ drug discovery.

In this research we decide to synthesize new iodoquinazolinone derivatives as anticancer agents due to iodine's high atomic number, and ease of attachment to organic compounds, and it has also found favour as a non-toxic radiocontrast material. Because of the specificity of its uptake by the human body, radioactive isotopes of iodine can also be used to treat thyroid cancer. Moreover, there are many approved drugs containing iodo substitutions *e.g.* 2-iodo-4'-methoxychalcone,⁴¹ amiodarone and iodochlorhydroxyquinoline (Clioquinol) (Fig. 1). On the other hand, there are many quinazoline derivatives targeting either EGFR and/or VEGFR-2 signaling pathways such as Gefitinib, Erlotinib, Vandetanib, Lapatinib, WHI-P180 and Afatinib³⁸ (Fig. 1).

The hybridization strategy aimed at combining the EGFR, VEGFR inhibition and anticancer activity of quinazolines with other privileged pharmacophores possessing antitumor potency could enhance the activity of the compounds.^{2,42} In this work, we designed some new iodoquinazolinones as potential VEGFR-2 and EGFR inhibitors by keeping the acetamide linker attached to the N-3 of the iodoquinazolinone ring and modification of the distal side chains to be aliphatic, aromatic and/or heterocyclic rings.

The current investigation is an extension to our preceding work^{12,43–51} that reveals the efficacy of the hybridization strategy between iodoquinazolinone and different moieties as EGFR, VEGFR-2 inhibitors and anticancer agents. We designed and synthesized a series of new 6-iodoquinazolinone bearing different moieties with acetamide linker as dual EGFR-mutant/VEGFR-2 inhibitors for the treatment of various cancer types, as liver (HepG2), breast (MCF-7), colorectal (HCT-116) and NSCLC (A549). *In silico* molecular modeling study was performed for our compounds as dual VEGFR-2/EGFR inhibitors to explore their binding mode in the enzymes active sites or predict their ADMET properties.

1.1. Rationale and structure-based design

Our derivatives were designed based on the pharmacophoric features of both EGFR and VEGFR-2 inhibitors. We synthesized



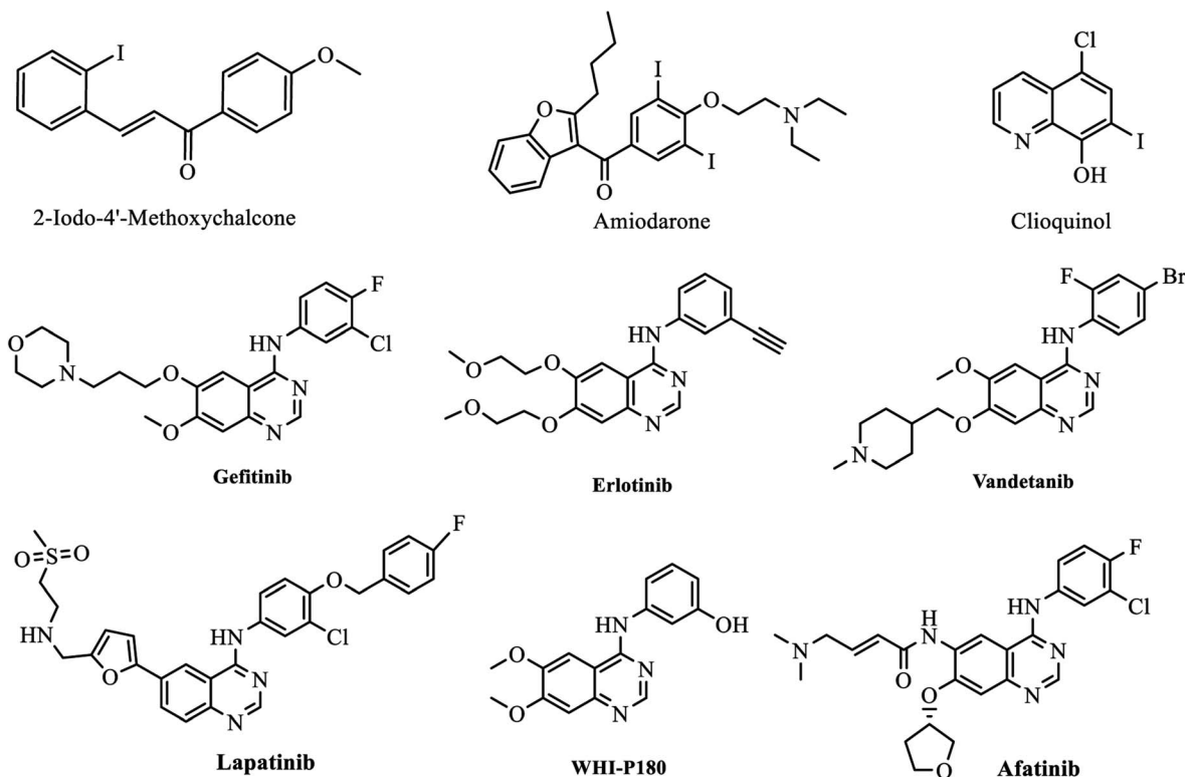


Fig. 1 Examples of drugs containing iodo substitutions and some quinazoline multitargeted TKIs.

a series of new iodoquinazoline derivatives inspired by the bioisosteric modifications of VEGFR-2 inhibitors (sorafenib and sunitinib) at four different positions: (1) a flat hetero-aromatic ring that occupies the ATP binding domain, (2) a central hydrophobic spacer, (3) a linker with a functional group that has both H-bond acceptor and donor properties to bind with key amino acids (Glu885 and Asp1046), and (4) a terminal hydrophobic moiety that fits into the allosteric hydrophobic pocket (Fig. 2).^{52,53}

Also, the iodoquinazolinone derivatives **5a–d** to **9a–e** possess the essential pharmacophoric features of EGFR-TKIs. These derivatives are based on structural modifications of erlotinib at four different positions: (1) a terminal hydrophobic head that fits into hydrophobic region I, (2) a flat hetero-aromatic system that occupies the adenine binding pocket, (3) HBD and/or HBA (NH as HBD) that forms important hydrogen bond interactions with amino acid residues in the linker region, and (4) a hydrophobic tail that fits into hydrophobic region II (Fig. 3).⁵⁴

2. Results and discussion

2.1. Chemistry

The synthetic strategy for preparation of the target compounds **5a–d** to **9a–e** is depicted in Scheme 1 and 2. Synthesis was initiated by reaction of chloroacetyl chloride with the appropriate amine **1a–c** to afford the corresponding chloroacetamide **2a–c**.^{55,56} The chloroacetamide **2a** underwent reaction with the appropriate methyl and/or ethylamine according to the mixed anhydride procedure⁵⁶ to afford the corresponding derivatives **3a,b**

respectively. On the other hand anthranilic acid (**1**) reacted with iodine to afford 5-iodoanthranilic acid (**2**) which underwent alkylation with ethyl iodide and/or propyl iodide to give the corresponding *N*-alkyl derivatives (**3a,b**).⁵⁷ Fusion of 5-iodo-*N*-alkylanthranilic acid (**3a,b**) with urea afforded the corresponding 6-iodo-1-alkylquinazoline-2,4(1*H*,3*H*)-dione (**4a,b**).⁵⁷ *In situ* reaction of 6-iodo-1-alkylquinazoline-2,4(1*H*,3*H*)-dione (**4a,b**) with the appropriate chloroacetamide **IIb,c** afforded the corresponding ester derivatives **5a–d**. Also reaction of 6-iodo-1-alkylquinazoline-2,4(1*H*,3*H*)-dione (**4a,b**) with the appropriate chloroacetamide **IIIa,b** to get the corresponding amide derivatives **6a,b** (Scheme 1). Reaction of the appropriate ethyl ester **5b,d** with hydrazine hydrate resulted in the corresponding hydrazide derivative **7a,b** which underwent cyclization with carbon disulfide giving the corresponding 5-sulfanyloxadiazole **8a,b** respectively. On the other hand, condensation of the appropriate acid hydrazide **7a,b** with the appropriate benzaldehyde afforded the corresponding Schiff's bases **9a–e** respectively (Scheme 2).

2.2. Docking studies

For molecular docking investigations, Molsoft software was used. The PDB IDs for VEGFR-2 (PDB ID 4ASD)^{48,58} and EGFR^{T790M} (PDB ID 3W2O)^{55,56} were used in each experiment, respectively.

2.2.1. Docking studies using VEGFR-2 inhibitors. The results obtained demonstrated that all the investigated congeners have the same place and location inside the identified binding site of VEGFR-2, which exposes a sizable gap limited by a membrane-binding domain that acts as an entry conduit for



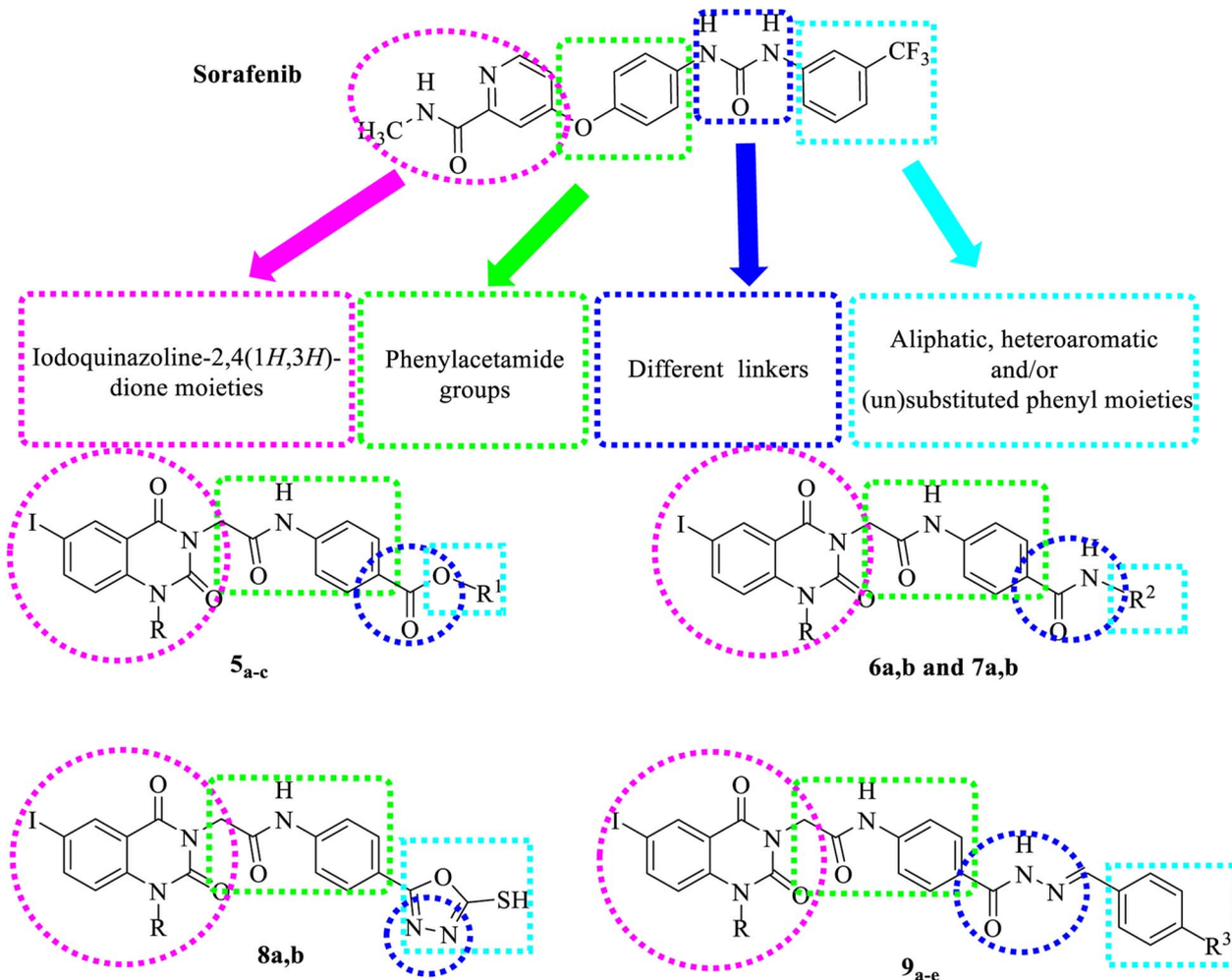


Fig. 2 Basic structural features of VEGFR-2 inhibitors.

substrate to the active site (Fig. 4). The majority of these drugs showed strong receptor binding affinity, as indicated by the binding free energy (ΔG) as shown in (Table 1).

The **Sorafenib**-suggested binding mechanism produced 5 H-bonds and an energy value of $-99.50 \text{ kcal mol}^{-1}$. It generated 2 hydrogen bonds with Cys919 (2.51 Å and 2.10 Å), two with Glu885 (1.77 Å and 2.75 Å), and one with Asp1046 (1.50 Å). The pocket created by Cys919, Phe918, Leu1035, Lys920, Glu917, Val848 and Leu840 occupied by the *N*-methylpicolinamide group. Additionally, the hydrophobic groove formed by Cys1045, Leu1035, Thr916, Lys868, and Val848 occupied by the central phenyl linker. Additionally, Cys1045, Asp1046, Hie1026, Ile888, Ile892 and Glu885 formed a hydrophobic channel that occupied by the terminal 3-trifluoromethyl-4-chlorophenyl group (Fig. 5). The urea spacer had a crucial part in the binding of the VEGFR-2 enzyme, but it also played a crucial role in the high binding affinity of **Sorafenib**. These results encourage us to experiment with various linkers in order to obtain efficient VEGFR-2 inhibitors.

Compound **9c** predicted binding mode is nearly identical to that of **Sorafenib**, which showed an affinity value of $-110.10 \text{ kcal mol}^{-1}$ and 6 H-bonds. 1-Ethyl-6-iodoquinazoline ring formed one H-bond

with Cys919 (2.33 Å). The acetamide linker also formed another H-bond with Cys919 (2.96 Å). The acid hydrazide linker formed three H-bonds with Asp1046 (2.50 Å) and Glu885 (2.15 Å and 2.21 Å), while the 4-methoxy group at the side chain formed one H-bond with Ile1025 (2.98 Å). The pocket formed by Leu1035, Lys920, Cys919, Phe918, Glu917, Val848 and Leu840 occupied by the Iodoquinazoline scaffold. Additionally, the hydrophobic channel created by Hie1026, Ile1025, Ile892, Ile888 and Glu885 occupied by the distal 4-methoxyphenyl moiety (Fig. 6).

Compound **9d** predicted binding mode is nearly identical to that of **Sorafenib** and **9c**, which showed an affinity value of $-108.45 \text{ kcal mol}^{-1}$ and 5 H-bonds. One H-bond was created with the 6-iodo-1-propylquinazoline ring and Cys919 (2.70 Å). The acetamide linker also formed another H-bond with Cys919 (2.97 Å). The carbonyl of hydrazide linker made two H-bonds with Lys868 (2.02 Å and 2.63 Å), and its NH group designed a third hydrogen bond with Asp1046 (2.43 Å) (Fig. 7). Moreover, compound **9b** suggested binding mode has five H-bonds and an affinity value of $-107.14 \text{ kcal mol}^{-1}$, which is almost identical to that of compound **9c**. It formed five H-bonds with Cys919 (2.74 Å and 2.98 Å), Lys868 (2.00 Å and 2.80 Å) and Asp1046 (2.28 Å) (Fig. 8).



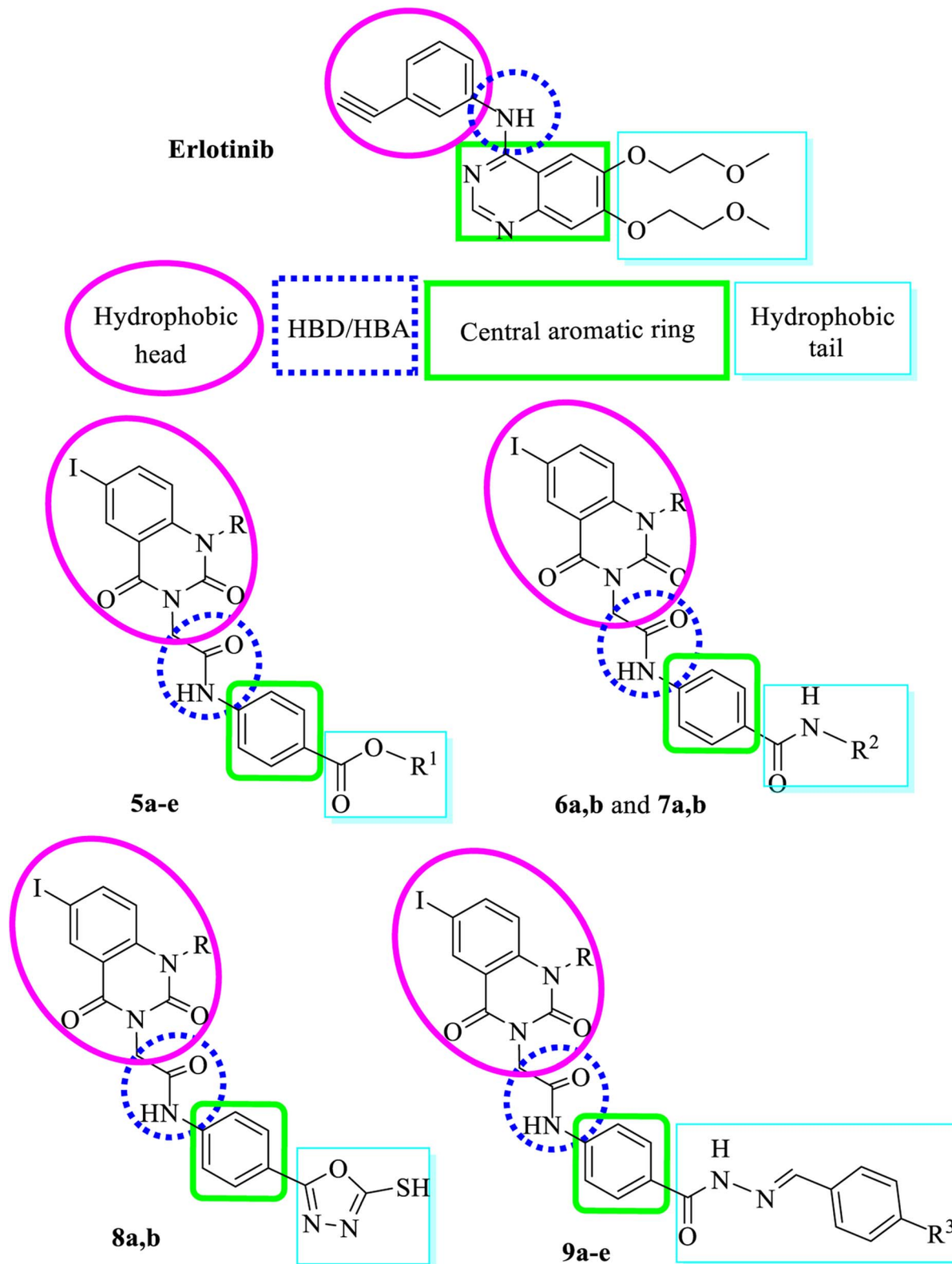
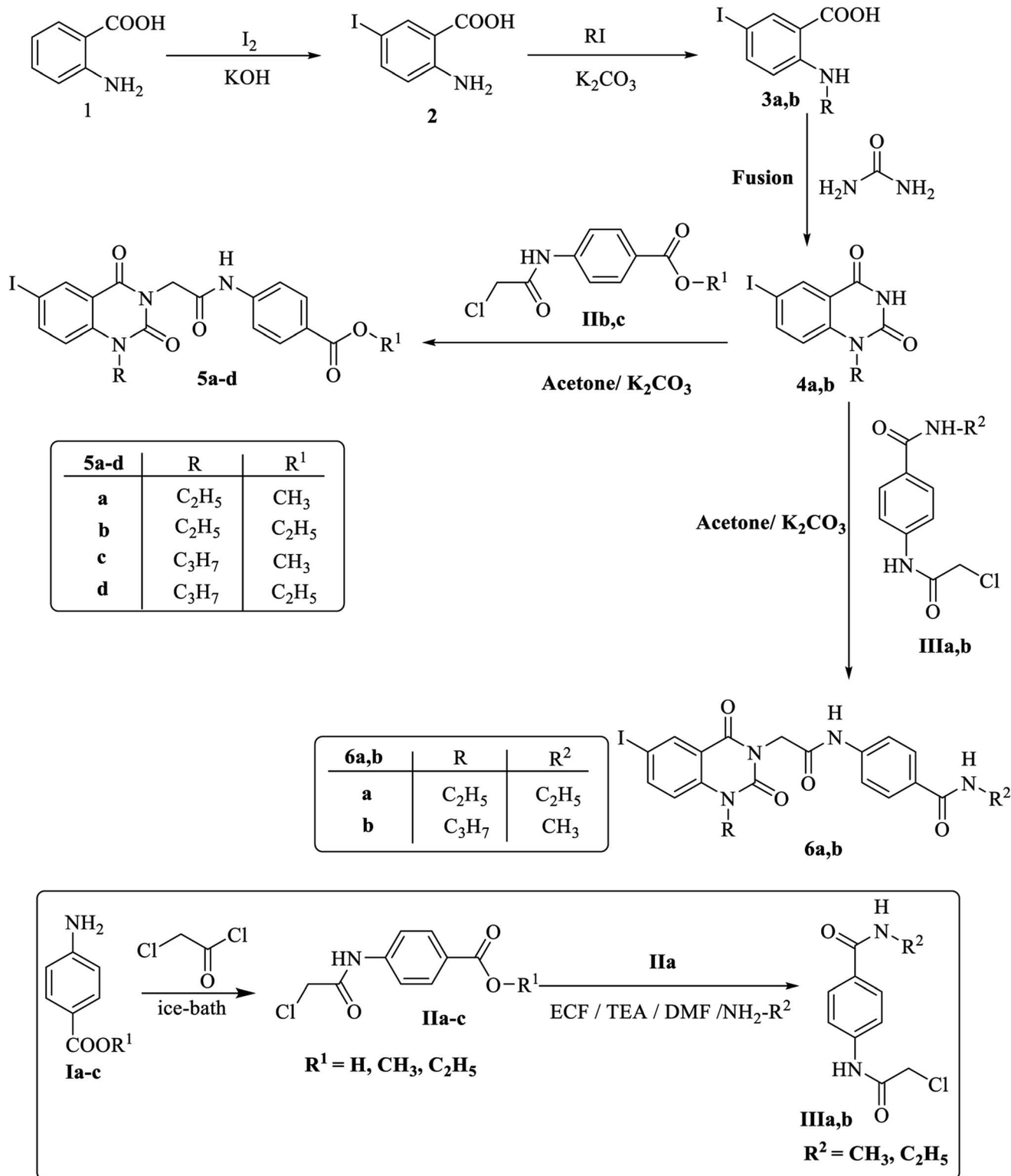


Fig. 3 Basic structural features of EGFR inhibitors.

The obtained binding moods in docking results (Table 1) led us to the conclusion that the used linkers shared the same groove as the urea linker of **Sorafenib** and performed the same function,

both of which are necessary for greater affinity towards the VEGFR-2 enzyme. The establishment of an H-bond with cysteine 919 and the emergence of hydrophobic interactions may have





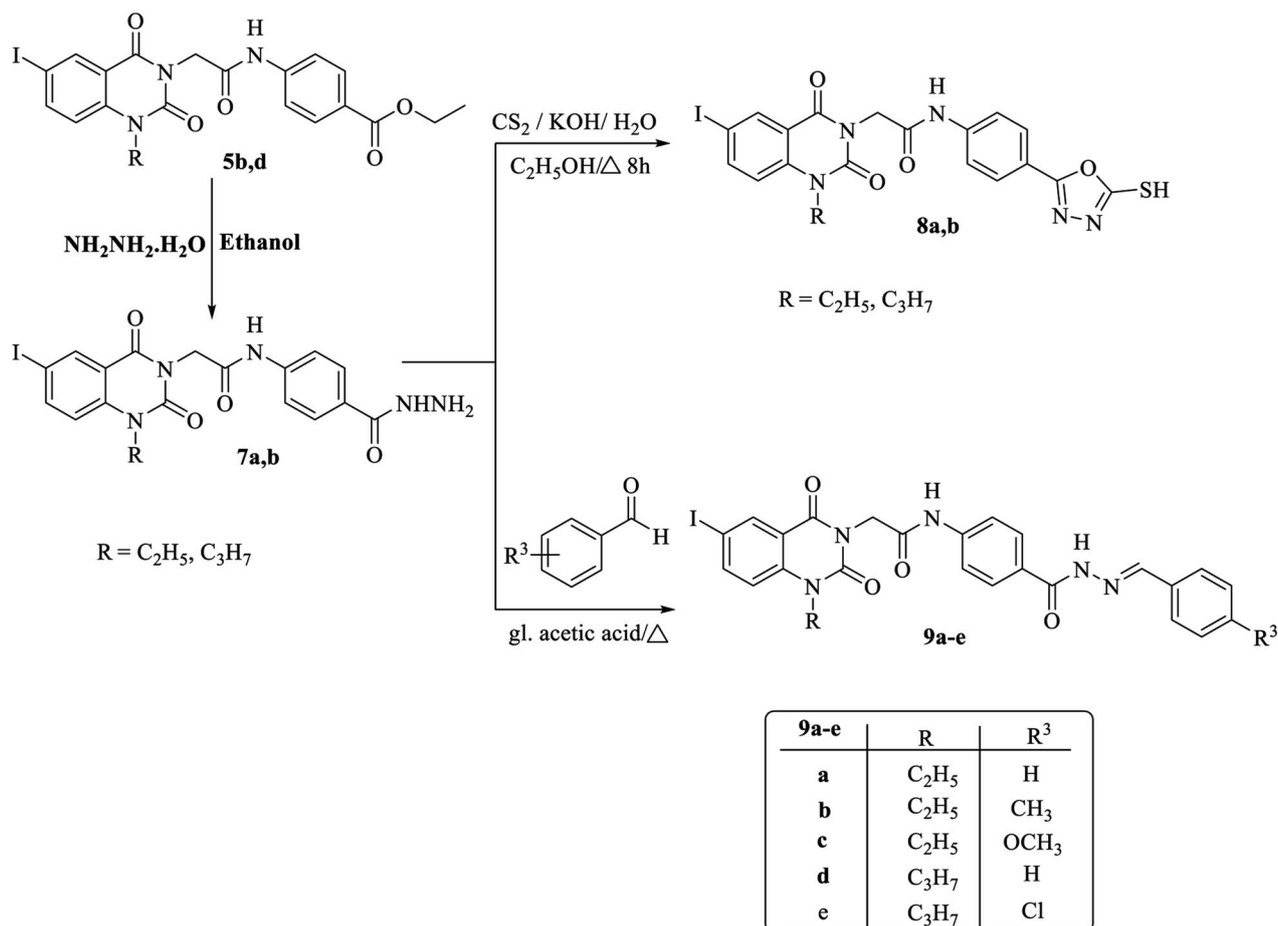
Scheme 1 Synthetic route for preparation of the target compounds 5a–d and 6a,b.

contributed to the heterocyclic *N*-alkyl-6-iodoquinazoline scaffold's higher affinity towards the VEGFR-2 enzyme.

2.2.2. Docking results for EGFR^{T790M} inhibitors studies. The obtained results demonstrated that each derivative in this study has a comparable position and orientation within the

identified binding site of EGFR (Fig. 9). The computed values mirrored the general trend and the picked up findings of the free energy of binding (ΔG) explained that the majority of these compounds had strong binding affinity for the receptor (Table 2).





Scheme 2 Synthetic route for preparation of the target compounds 7a,b to 9a-e.

Erlotinib's hypothesized binding mode revealed an binding energy value of $-82.77 \text{ kcal mol}^{-1}$ and four H-bonds, one of the two 2-methoxyethoxy groups established one H-bond with the

residing amino acid Cys797 (2.05 Å), The creation of two H bonds with Val726 (2.97 Å) and Met793 (1.82 Å) and stabilized the quinazoline moiety. The NH spacer and Thr854 (2.99 Å) established

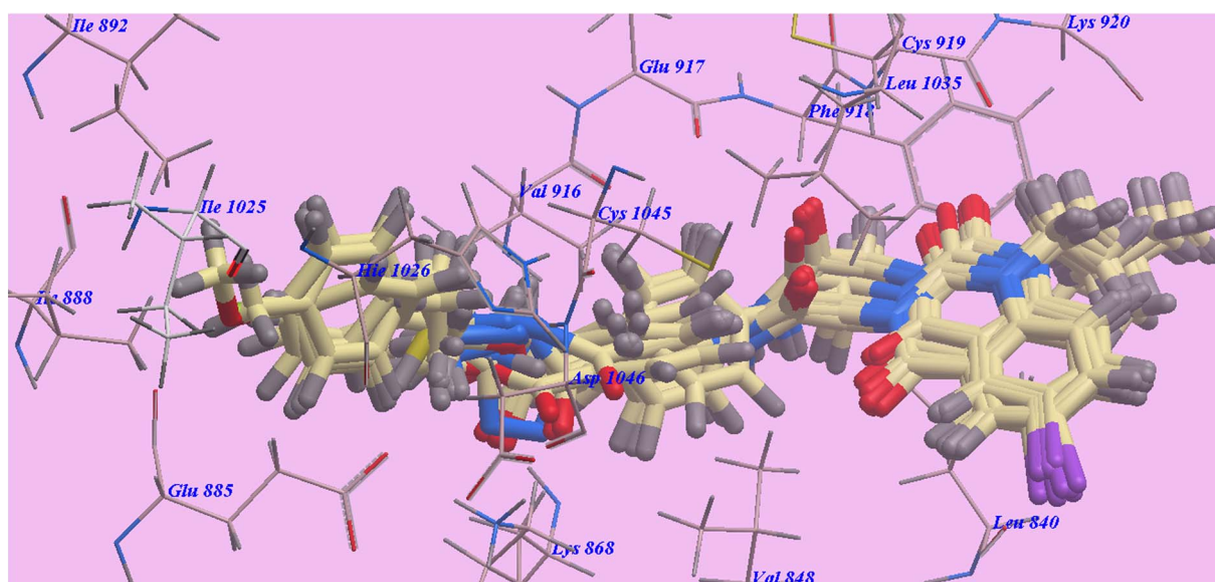


Fig. 4 Compounds superimposed inside the active site of 4ASD.



Table 1 Calculated free energy (ΔG in kcal mol⁻¹) of ligands binding with VEGFR-2

Compound	ΔG [kcal mol ⁻¹]	Compound	ΔG [kcal mol ⁻¹]
5a	-86.20	8a	-88.77
5b	-90.99	8b	-93.50
5c	-87.37	9a	-102.82
5d	-95.01	9b	-107.14
6a	-91.18	9c	-110.10
6b	-88.47	9d	-108.45
7a	-79.59	9e	-102.12
7b	-83.13	Sorafenib	-99.50

one H-bond. The hydrophobic region I, which formed by Val726, Asp855, Thr854, Glu791, Met790, Leu777, Glu762, Ile759, and Phe723, was occupied by the 3-ethynylphenyl head. Additionally, the 2-methoxyethoxy tail occupied the hydrophobic area II created by Val845, Leu844, Pro794, Met793, and Leu718 (Fig. 10).

Compound **9c** predicted binding mode is nearly identical to that of **erlotinib**, which displayed an affinity value of -90.95 kcal mol⁻¹ and 6 H-bonds. Iodoquinazoline scaffold made two hydrogen bonds with Thr854 (2.22 Å) and Met790 (2.96 Å), while acetamide linker formed one hydrogen bond with Met793 (2.27 Å). Moreover, the hydrazide linker formed three H-bonds with Met793 (2.96 Å), Pro794 (2.77 Å) and Phe795 (2.95 Å). Moreover, the 4-methoxy group formed H-bond with Tyr801 (2.98 Å). The hydrophobic region II, which created by Val845, Leu844, Cys797, Phe795, Pro794, Met793, Leu718 and Tyr801, was occupied by the distal 4-methoxyphenyl side chain and the acid hydrazide linker. The hydrophobic region I, which was created by Asp855, Thr854, Met790, Leu788 and Gly724 was occupied by iodoquinazoline scaffold (Fig. 11).

Compounds **9d** and **9e** have suggested binding mechanisms that are nearly identical to those of **erlotinib** and **9c**, with affinities of -89.30 kcal mol⁻¹ (formed five H-bonds) (Fig. 12) and -88.10 kcal mol⁻¹ (formed 5 H-bonds) (Fig. 13), respectively.

2.3. Biological evaluation

2.3.1. Cytotoxicity evaluation. The MTT assay, created by Mosmann,⁵⁹⁻⁶¹ was used to assess the novel iodoquinazoline derivatives **5a-c** to **9a-e**. The drugs were tested on four types of human tumor cell lines (HepG2, MCF-7, HCT-116 and A549). In Table 3, half maximal effective concentration (EC₅₀) values for the study's reference cytotoxic drugs, sorafenib and erlotinib, are shown. The findings showed that most of the compounds had very good to moderate EC₅₀ efficacy against the cancer cell types under study. Particularly, compound **9c** showed the highest anticancer activities with EC₅₀ = 5.00, 6.00, 5.17 and 5.25 μM against HepG2, MCF-7, HCT116 and A549 cell lines correspondingly. Despite it showed lower activities than sorafenib (EC₅₀ = 4.00, 5.05, 5.58 and 4.04 μM) versus HepG2, MCF-7 and A549 contrarily subagent performance against HCT116, it demonstrated higher actions than erlotinib (EC₅₀ = 7.73, 13.91, 8.20 and 5.49 μM), against the four tested cell lines.

Regarding HepG2, compounds **5b-d**, **6a-b**, **8a-b** and **9a,b,d,e** exhibited very good anticancer effects with EC₅₀ ranging from 5.65 to 8.50 μM. Derivative **5a** with EC₅₀ = 12.53 μM, showed good cytotoxicity. On the other hand, derivatives **7a** and **7b** exhibited moderate anticancer effects with EC₅₀ = 16.00 and 15.50 μM respectively.

Concerning MCF-7, derivatives **5b-d**, **6a-b**, **8a-b** and **9a,b,d,e** exhibited very good anticancer effects with EC₅₀ ranging from

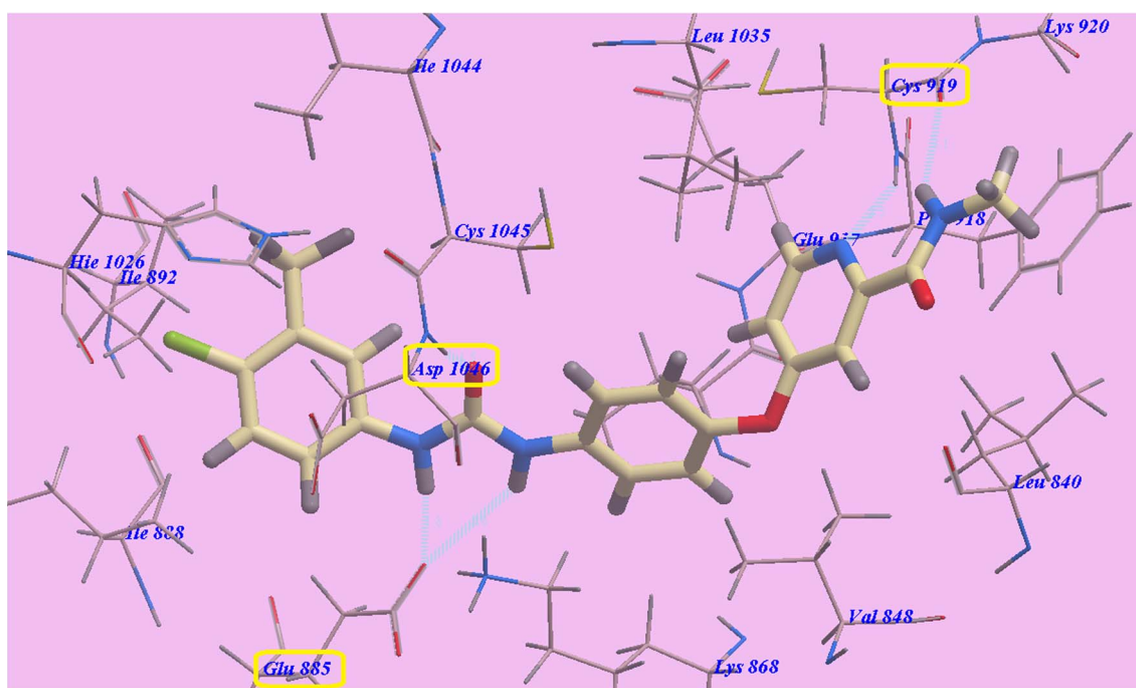


Fig. 5 Sorafenib predicted mechanism of binding with 4ASD. Dotted lines represent atoms that are H-bonded.



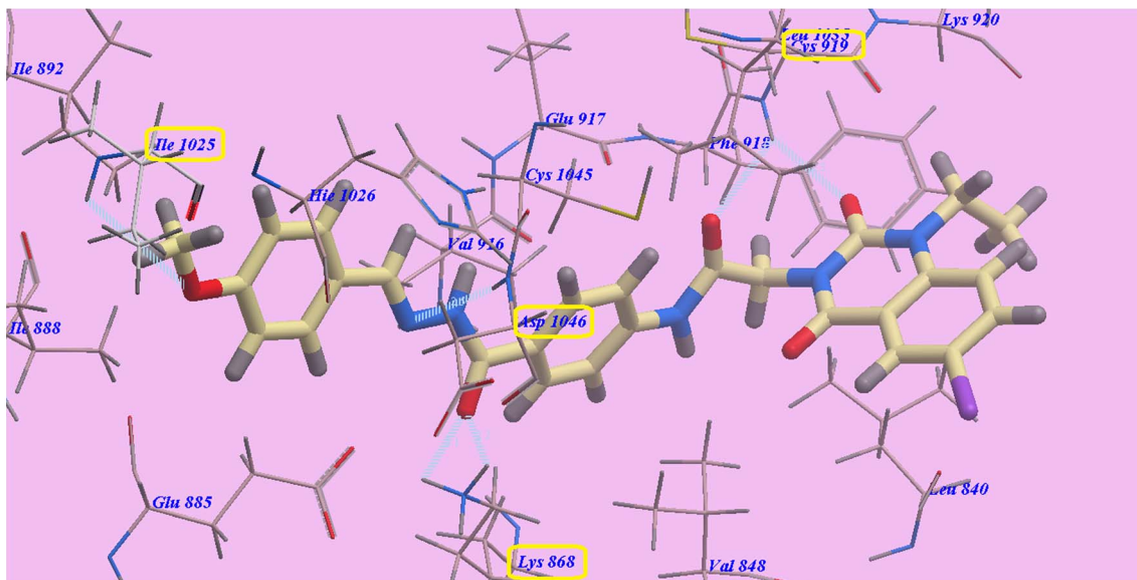


Fig. 6 9c predicted binding mode with 4ASD.

6.45 to 8.50 μM . Derivatives **5a** and **7b** with EC_{50} = 11.60 and 13.90 μM respectively, showed good cytotoxicity. On the other hand, derivative **7a** exhibited moderate anticancer effects with EC_{50} = 15.50 μM .

Derivatives **5b–d**, **6a–b**, **8a–b** and **9a,b,d,e** exhibited very good anticancer effects with EC_{50} ranging from 5.93 to 8.35 μM against HCT-116. Derivatives **5a** and **7a** with EC_{50} = 11.44 and 13.25 μM respectively, showed good cytotoxicity. On the other hand, derivative **7b** exhibited moderate anticancer effects with EC_{50} = 15.90 μM .

Derivative **5a–d**, **6a–b**, **8b** and **9a,b,d,e** exhibited very good anticancer effects with EC_{50} ranging from 5.48 to 8.45 μM

against A549. Derivatives **7a** and **8a** with EC_{50} = 13.66 and 10.10 μM , showed good cytotoxicity. On the other hand, derivative **7b** presented moderate anticancer effect with EC_{50} = 15.10 μM .

2.4. Selectivity index (SI)

Finally, the highly effective six derivatives **5d**, **8b** and **9a–e** were examined against VERO normal cell lines to estimate their cytotoxic capabilities. All conclusions revealed that each discovered compound possessed low toxicity regarding VERO normal cells with 50% cytotoxic concentration (CC_{50}) extending from 45.50 to 55.00 μM . One criterion for the anticancer drug to be good it should not affect the non-cancer cells. A molecule

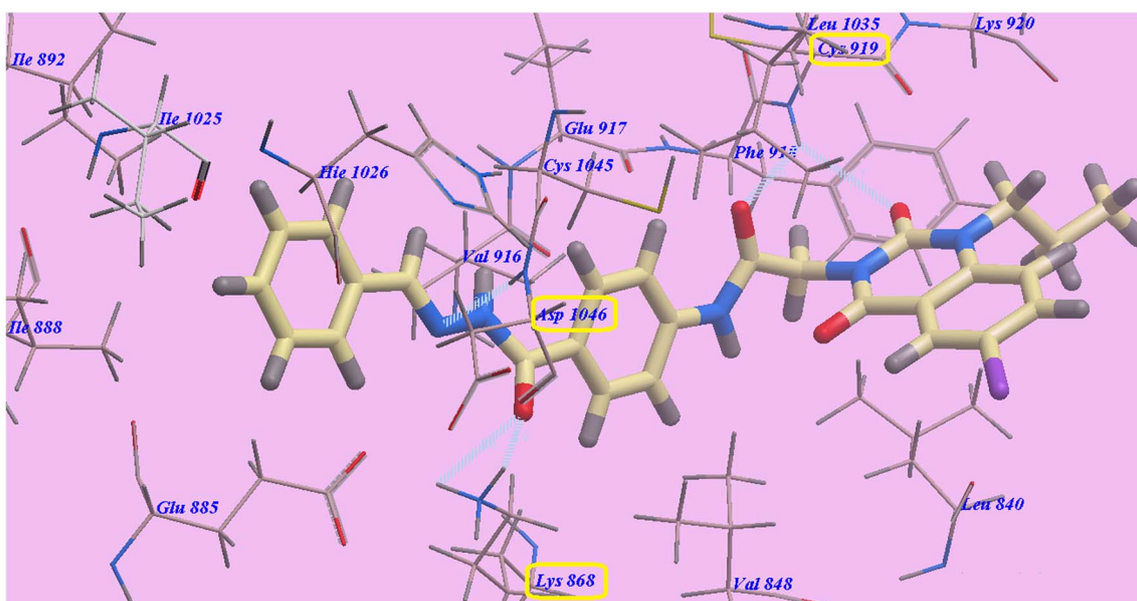


Fig. 7 Docking results for 9d with 4ASD.



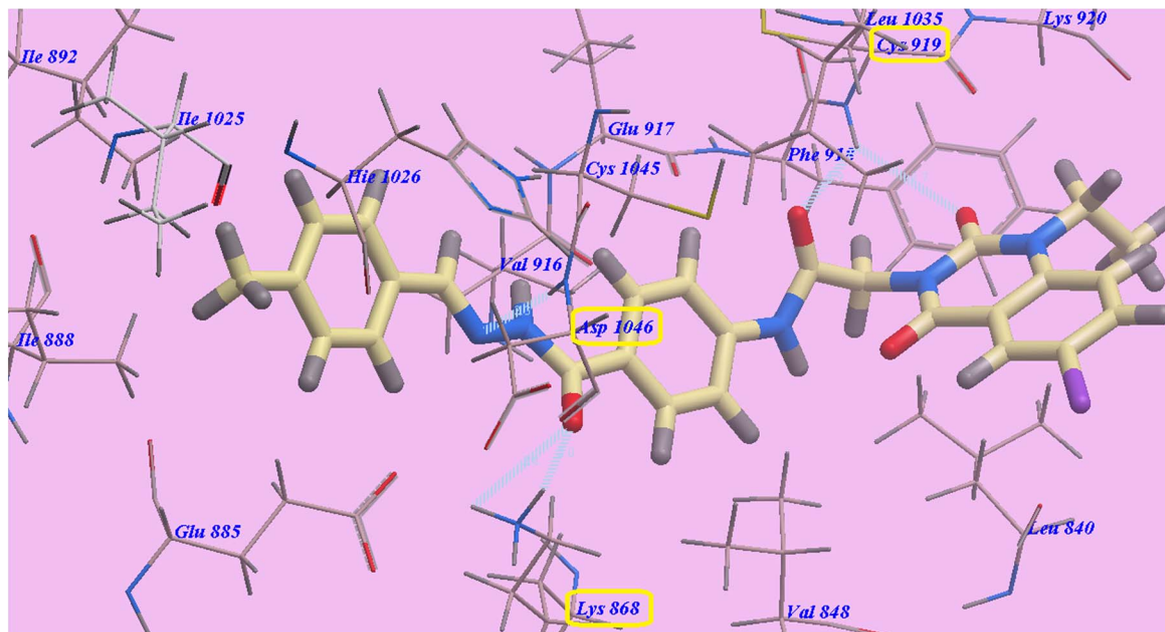


Fig. 8 Docking results for 9b with 4ASD.

could be considered as highly selective if it presented SI value ≥ 5 . A molecule with moderate selectivity presents SI value > 2 while low selectivity is considered if the SI is lower than 2.⁶² In this study, derivatives 5d, 8b, 9a, 9b, 9c, 9d and 9e are correspondingly 6.21, 6.77, 6.69, 7.94, 11.00, 8.92 and 6.25 folds more toxic respecting HepG2 than normal VERO cells. Uniformly, derivatives 5d, 8b, 9a, 9b, 9c, 9d and 9e are consequently 6.32, 6.81, 6.07, 7.35, 9.17, 7.81 and 6.30 folds toxic in MCF-7 than in normal VERO cells. Furthermore, structures 5d, 8b, 9a, 9b, 9c,

9d and 9e are respectively 7.15, 7.52, 7.23, 8.31, 10.64, 7.75 and 6.87 folds grander lethality in HCT-116 than in ordinary VERO cells. Furthermore, products 5d, 8b, 9a, 9b, 9c, 9d and 9e are respectively 6.98, 7.57, 6.74, 7.30, 10.48, 9.20 and 7.30 folds toxicity in A549 than in ordinary VERO cells. All derivatives exhibited high selectivity against the tested cancer cell lines where SI values > 5 .

2.4.1. *In vitro* VEGFR-2 kinase inhibitory assay. Additionally, all derivatives were evaluated for their ability to inhibit

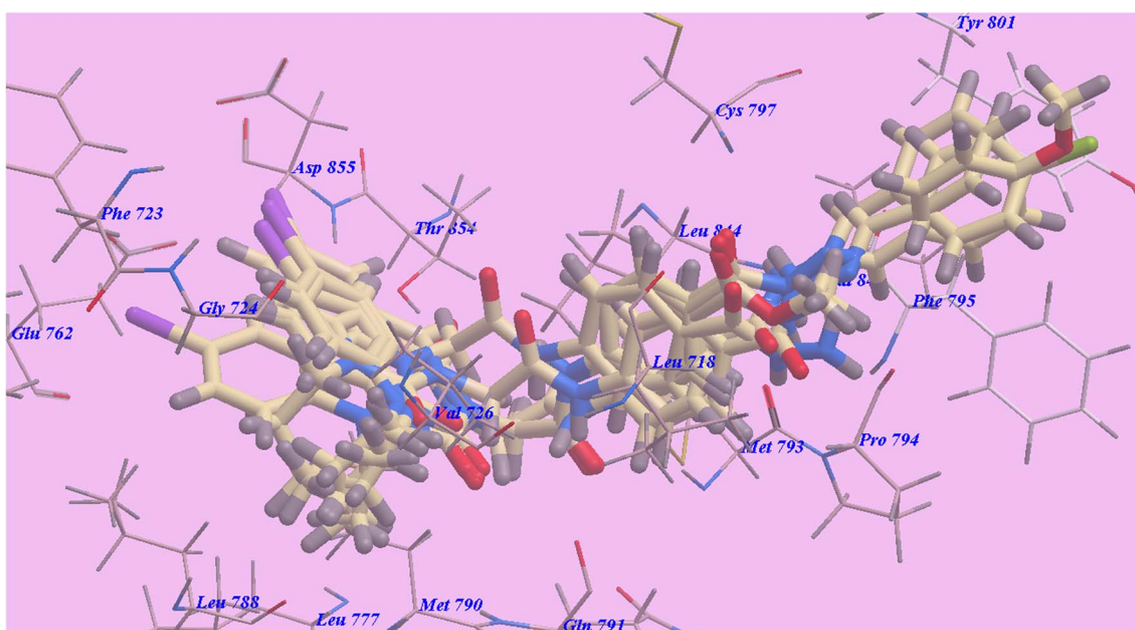


Fig. 9 Certain docked compounds superimposed inside the binding pocket of 3W2O.



Table 2 Calculated free energy (ΔG in kcal mole⁻¹) of synthesized compounds binding with EGFR^{T790M}

Compound	ΔG [kcal mol ⁻¹]	Compound	ΔG [kcal mol ⁻¹]
5a	-82.03	8a	-80.89
5b	-83.02	8b	-85.85
5c	-80.71	9a	-83.59
5d	-86.58	9b	-84.20
6a	-81.91	9c	-90.95
6b	-83.76	9d	-89.30
7a	-73.37	9e	-88.10
7b	-69.77	Erlotinib	-82.77

VEGFR-2 as a result of applying an anti-phosphotyrosine antibody with the Alpha Screen system (PerkinElmer, USA).^{62,63} In Table 3, the findings were presented as the IC₅₀ (50% inhibitory concentration). Sorafenib was used as a positive standard in this evaluation. The evaluated candidates showed an excellent to low inhibitory activities with IC₅₀ range of 0.85–2.50 μ M.

Compounds **9c**, **9b**, **9d**, **9a**, **9e** and **5d** excellently inhibited VEGFR-2 activity with IC₅₀ = 0.85, 0.90, 0.90, 1.00, 1.20 and 1.25 μ M respectively. Candidates **5b**, **5c**, **6a**, **6b**, **8a** and **8b** significantly inhibited VEGFR-2 at IC₅₀ range of 1.38–1.65 μ M. Alternatively, compounds **5a**, **7a** and **7b** moderately inhibited VEGFR-2 at IC₅₀ = 2.00, 2.50 and 2.25 μ M, respectively.

2.4.2. In vitro EGFR^{T790M} kinase inhibitory assay. The inhibitory activities against mutant EGFR^{T790M} kinases for all compounds were also assessed. This test used a homogeneous time resolved fluorescence (HTRF) assay.^{63,64} Erlotinib was used as standard with IC₅₀ = 0.24 μ M. A comparison of the IC₅₀ values for the investigated compounds is shown in Table 3.

Compounds **9c**, **9d**, **9e**, **5d**, **8b** and **9b** excellently inhibited EGFR^{T790M} activity with IC₅₀ = 0.22, 0.26, 0.30, 0.40, 0.45 and 0.50 μ M respectively. Candidates **5a–c**, **6b**, **8a** and **9a** significantly inhibited EGFR^{T790M} at IC₅₀ range of 0.60–1.00 μ M. Alternatively, compound **9a** moderately inhibited EGFR^{T790M} at IC₅₀ = 1.50 μ M. The least inhibitory effectiveness against EGFR^{T790M} was displayed by compound **7b**. As planned, compound **9c** showed excellent dual EGFR^{T790M}/VEGFR-2 inhibitory activities.

2.4.3. In vitro EGFR^{WT} kinase inhibitory assay. The inhibitory activities of compounds **5d**, **8b**, **9b**, **9c**, **9d** and **9e** against EGFR^{WT} kinases were also assessed (Table 3).⁶⁵ Erlotinib was used as standard with IC₅₀ = 0.18 μ M. Compounds **9c**, **9d** and **9e** excellently inhibited EGFR^{WT} activity with IC₅₀ = 0.15, 0.20 and 0.25 μ M respectively. In addition compounds **5d**, **9b** and **8b** significantly inhibited EGFR^{WT} activity with IC₅₀ = 0.32, 0.35, and 0.38, μ M respectively.

2.4.4. Structure activity relationship (SAR). Generally, the 1-alkyl-6-iodoquinazolidione ring, the spacer and the linker, as well as the lipophilicity and electronic nature of the substituents, all played a significant influence in the anticancer action. The HCT-116 and A549 cell lines were the most sensitive to the effect of our compounds. Generally, compounds **9a–e** with Schiff's bases side chains displayed higher anticancer activities than that with aliphatic side chains (compounds **5a–d**, **6a,b** and **7a,b**) and that with heteroaromatic ones (compounds **8a,b**). It was shown that compound **9c** showed the highest anticancer activity compared to the other compounds against HepG2, HCT116, A549 and MCF-7 respectively. Also, it displayed the highest activities as dual EGFR^{T790M} and VEGFR-2 inhibitor.

Based on the structure of the synthesized derivatives and the data in Table 3, we can divide the tested compounds into three

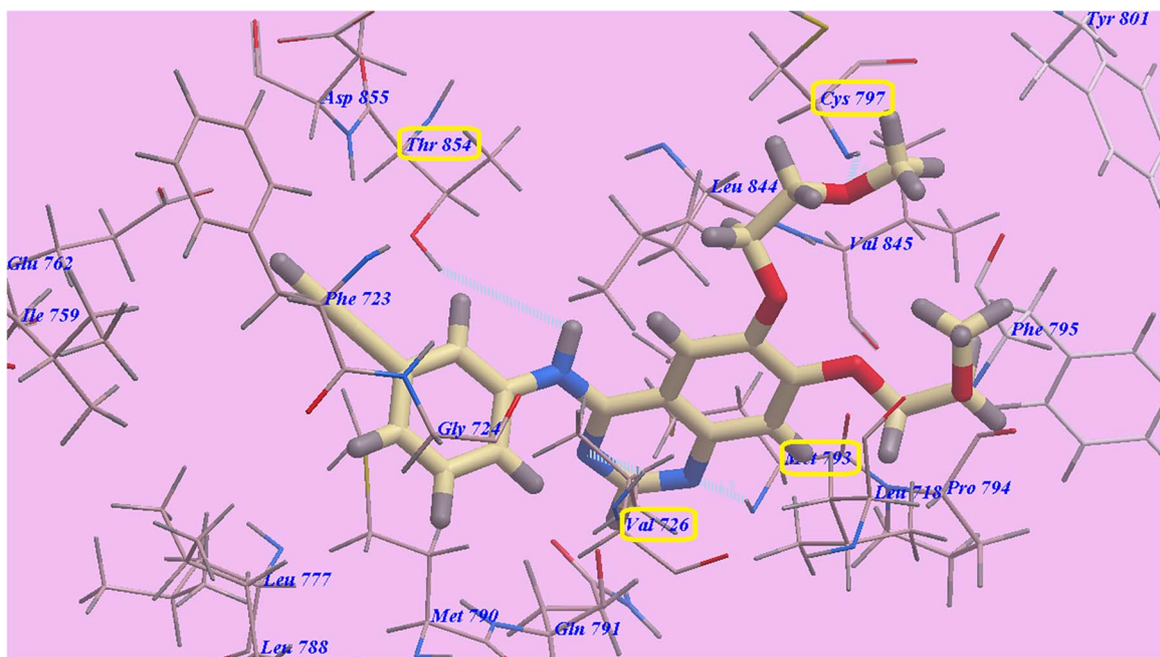


Fig. 10 Docking results for Erlotinib with 3W2O, H-bonded atoms are specified by dotted lines.



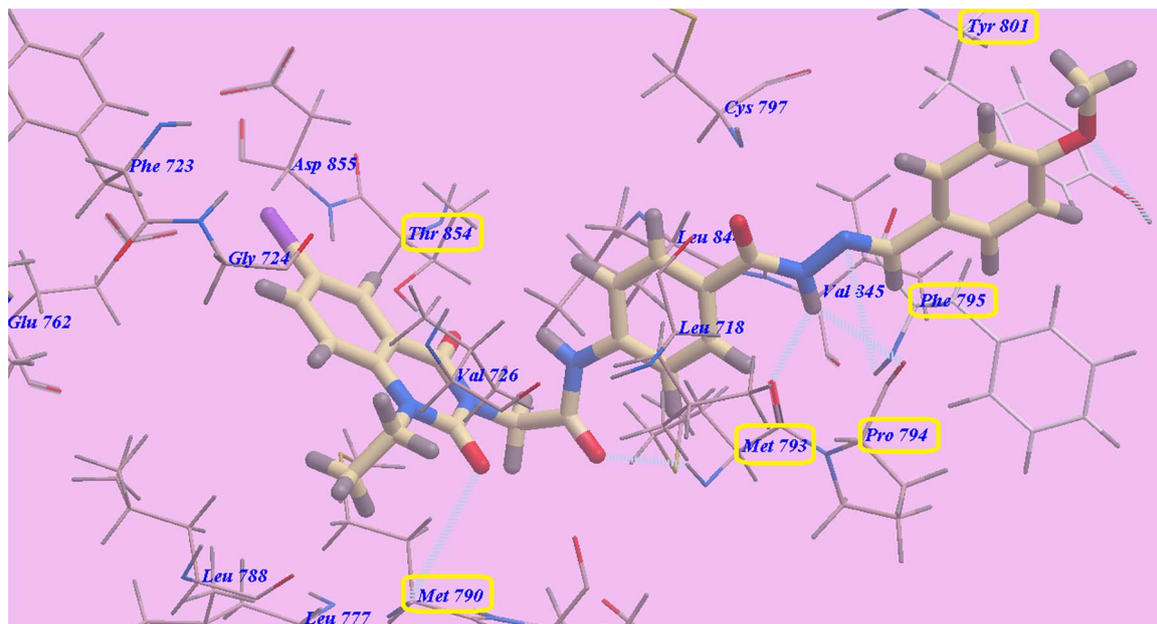


Fig. 11 Predicted binding mode for 9c with 3W2O.

groups. Compounds **5a–d**, **6a,b** and **7a,b** which have ester, amide and hydrazide moieties respectively as side chains, belong to the first category. Generally in this group esters exhibited higher anticancer activities than amides and hydrazides respectively.

Compound **5d** with the more hydrophobic 1-propyl and ethyl ester side chain exhibited higher anticancer activities than compound **5b** with 1-ethyl and ethyl ester side chain, compound **5c** with 1-propyl and methyl ester side chain and **5a**

with 1-ethyl and methyl ester side chain against the four tested cell lines except A549 where the order of activity is **5d** > **5c** > **5b** > **5a**. Moreover, the ester derivatives **5d–c** showed higher activities than the amides **6a** and **6b** against HepG2, MCF-7 and HCT-116, but in case of A549 the order of activity is **5d** > **6b** > **5c** > **5b** > **6a**. In addition, the amide **6a** with 1-ethyl and ethyl amide side chain displayed higher activities than the amide **6b** with 1-propyl and methyl amide side chain against HepG2 and HCT-116 but **6b** showed higher activities than **6a** against MCF-7

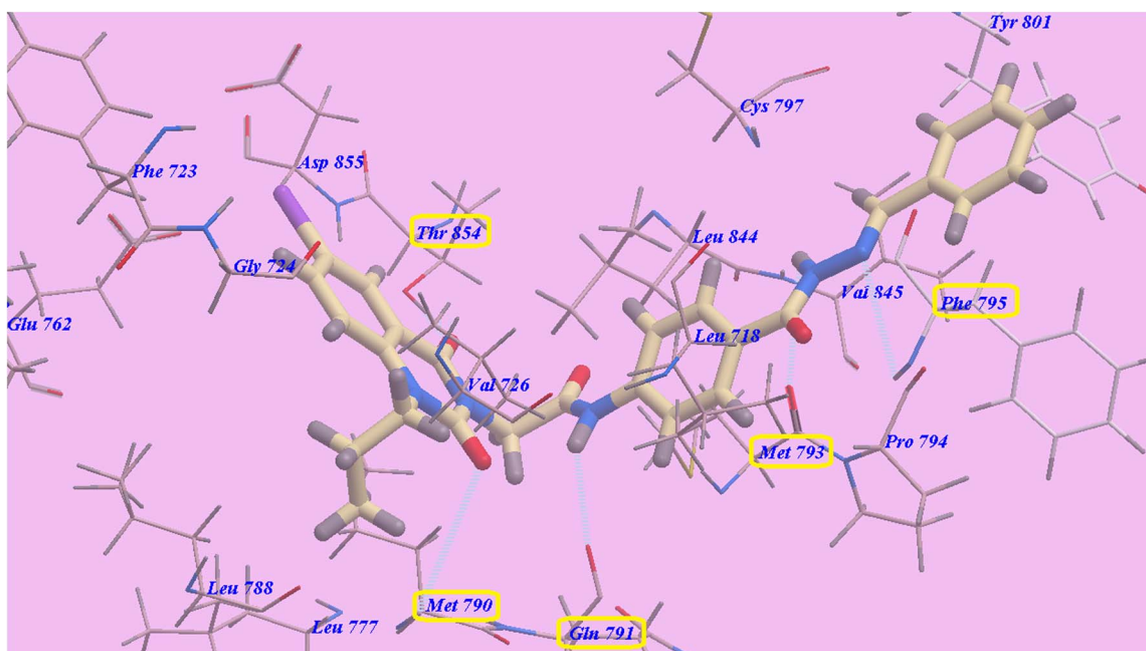


Fig. 12 Docking results for compound 9d with 3W2O.



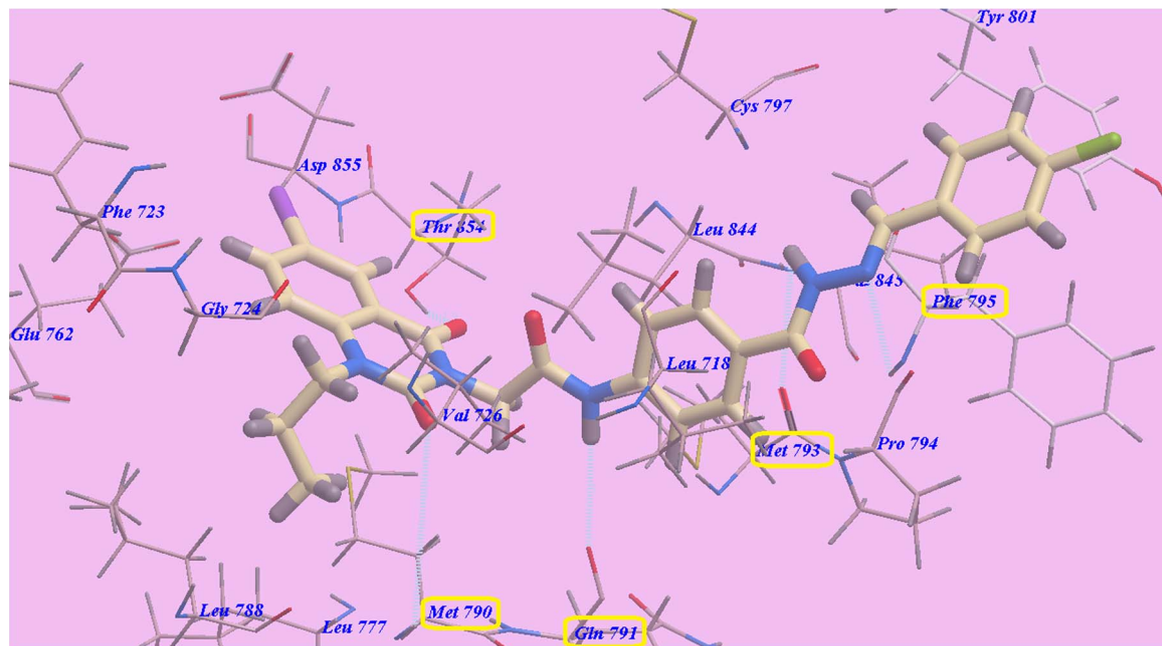


Fig. 13 Docking results for compound 9e with 3W2O.

and A549. The hydrazone derivatives **7a** and **7b** displayed the lowest anticancer activities along this series. The hydrazone **7b** with 1-propyl substitution exhibited higher activities than **7a** with 1-ethyl against HepG2 and MCF-7 cell lines but in case of HCT-116 and A549 cell lines **7a** showed higher activities than **7b**.

Compounds **8a** and **8b** in the second group have a distal hydrophobic heteroaromatic oxadiazole moiety. Compound **8b** with 1-propyl and oxadiazol-5-thiol side chain groups showed stronger effects than compound **8a** with 1-ethyl and oxadiazol-5-

thiol side chain groups against the four HepG2, MCF-7, HCT-116 and A549 cell lines. This finding indicated that the long chain alkyl group at position-1 increases the anticancer activities.

The third group consists of compounds **9a–e** with distal hydrophobic Schiff's bases side chains. Compound **9c** with 1-ethyl and highly electron donating methoxy group at position-4 of the phenyl side chain showed strongest activities against the four cell lines along this group. Derivative **9d** with 1-propyl and unsubstituted phenyl side chain showed higher activities than

Table 3 EGFR^{T790M} and VEGFR-2 kinase assays and *in vitro* cytotoxic effects against the HepG2, MCF-7, HCT-116, A549 and VERO cell lines

Comp.	EC ₅₀ (μM) ^a				CC ₅₀ (μM) ^b	IC ₅₀ (μM) ^c		
	HepG2	MCF-7	HCT116	A549	VERO	VEGFR-2	EGFR ^{T790M}	EGFR ^{WT}
5a	12.53 ± 0.7	11.60 ± 0.7	11.44 ± 0.7	8.45 ± 0.7	NT ^d	2.00 ± 0.25	0.86 ± 0.50	NT ^d
5b	7.59 ± 0.7	7.40 ± 0.7	6.55 ± 0.7	6.78 ± 0.7	NT ^d	1.50 ± 0.10	0.80 ± 0.50	NT ^d
5c	7.75 ± 0.7	7.96 ± 0.7	6.93 ± 0.7	6.73 ± 0.7	NT ^d	1.50 ± 0.50	0.92 ± 0.50	NT ^d
5d	7.50 ± 0.7	7.37 ± 0.7	6.52 ± 0.7	6.68 ± 0.7	46.60 ± 0.50	1.25 ± 0.10	0.40 ± 0.35	0.32 ± 0.03
6a	7.80 ± 0.8	8.50 ± 0.8	7.60 ± 0.8	7.95 ± 0.8	NT ^d	1.40 ± 0.20	0.90 ± 0.50	NT ^d
6b	8.50 ± 0.8	8.15 ± 0.8	8.35 ± 0.8	6.70 ± 0.8	NT ^d	1.65 ± 0.10	0.60 ± 0.35	NT ^d
7a	16.00 ± 1.5	15.50 ± 1.5	13.25 ± 1.5	13.66 ± 1.5	NT ^d	2.50 ± 0.25	1.50 ± 0.50	NT
7b	15.50 ± 1.5	13.90 ± 1.5	15.90 ± 1.5	15.10 ± 1.5	NT ^d	2.25 ± 0.25	2.05 ± 0.50	NT ^d
8a	7.55 ± 0.9	8.25 ± 0.9	7.25 ± 0.9	10.10 ± 0.9	NT ^d	1.50 ± 0.10	1.00 ± 0.50	NT ^d
8b	7.50 ± 0.7	7.45 ± 0.7	6.75 ± 0.7	6.70 ± 0.5	50.75 ± 0.50	1.38 ± 0.10	0.45 ± 0.35	0.38 ± 0.03
9a	6.80 ± 0.7	7.50 ± 0.7	6.29 ± 0.7	6.75 ± 0.7	45.50 ± 0.50	1.00 ± 0.10	0.75 ± 0.50	NT ^d
9b	6.20 ± 0.7	6.70 ± 0.6	5.93 ± 0.6	6.75 ± 0.6	49.25 ± 0.50	0.90 ± 0.10	0.50 ± 0.35	0.35 ± 0.03
9c	5.00 ± 0.5	6.00 ± 0.5	5.17 ± 0.5	5.25 ± 0.5	55.00 ± 0.50	0.85 ± 0.10	0.22 ± 0.30	0.15 ± 0.03
9d	5.65 ± 0.5	6.45 ± 0.5	6.50 ± 0.5	5.48 ± 0.5	50.39 ± 0.50	0.90 ± 0.10	0.26 ± 0.30	0.20 ± 0.03
9e	7.36 ± 0.7	7.30 ± 0.7	6.70 ± 0.7	6.30 ± 0.7	46.00 ± 0.50	1.20 ± 0.10	0.30 ± 0.30	0.25 ± 0.03
Sorafenib	4.00 ± 0.33	5.05 ± 0.50	5.58 ± 0.55	4.04 ± 0.33	NT ^d	0.84 ± 0.04	NT ^d	NT ^d
Erlotinib	7.73 ± 0.67	13.91 ± 1.3	8.20 ± 0.34	5.49 ± 0.45	NT ^d	NT ^d	0.24 ± 0.22	0.18 ± 0.02

^a EC₅₀ (50% effective concentration). ^b CC₅₀ (50% cytotoxic concentration). ^c IC₅₀ (50% inhibitory concentration). ^d NT: not tested compounds.



Table 4 The highest effective compounds, sorafenib and erlotinib; ADMET profile

Parameter	9b	9c	9e	Sorafenib	Erlotinib
Physicochemical properties					
Molecular weight	609.424	625.423	643.869	464.831	393.443
Log <i>P</i>	3.49872	3.1989	4.2338	5.5497	3.4051
Rotatable bonds	7	8	8	5	10
Acceptors	7	8	7	4	7
Donors	2	2	2	3	1
Surface area	225.514	230.628	235.817	185.111	169.532
Absorption					
Water solubility	−4.304	−4.504	−4.457	−4.822	−4.736
Caco2 permeability	0.384	0.399	0.421	0.689	1.431
Human intest. absorption	86.879	82.751	88.2	89.043	94.58
Skin permeability	−2.75	−2.748	−2.743	−2.767	−2.741
Substrate for P-glycoprotein	+	+	+	+	−
Inhibitor of P-glycoprotein I	+	+	+	+	+
Inhibitor of P-glycoprotein II	+	+	+	+	+
Distribution					
VDss (human)	−0.388	−0.353	−0.326	−0.29	0.199
Human unbound fraction	0.101	0.092	0.086	0.065	0.059
Permeability throughout BBB	−0.7	−0.912	−0.878	−1.684	−0.745
Permeability to CNS	−2.286	−2.535	−2.266	−2.007	−3.216
Metabolism					
CYP2D6 substrate	−	−	−	−	−
CYP3A4 substrate	+	+	+	+	+
Inhibition of CYP1A2	+	−	+	+	+
Inhibition of CYP2C19	+	+	+	+	+
Inhibition of CYP2C9	+	+	+	+	+
Inhibition of CYP2D6	−	−	−	−	−
Inhibition of CYP3A4	+	+	+	+	+
Excretion					
Clearance	−0.817	−0.838	−0.858	−0.219	0.702
Renal OCT2 substrate	−	−	−	−	−
Toxicity					
AMES toxicity	−	−	−	−	−
Human max. tolerated dose	0.421	0.421	0.533	0.549	0.839
Inhibitor of hERG I	−	−	−	−	−
hERG II inhibitor	+	+	+	+	+
Acute toxic activity (LD ₅₀)	2.439	2.468	2.346	2.538	2.393
Chronic toxic activity (LOAEL)	1.873	1.74	1.925	1.198	1.37
Hepatotoxic effect	+	+	+	+	+
Skin sensitization	−	−	−	−	−
T. Pyriformis toxicity	0.361	0.333	0.343	0.383	0.309
Minnow toxic activity	0.849	0.526	0.171	0.189	−0.1

compound **9b** 1-ethyl and 4-methylphenyl side chain and **9a** with 1-ethyl and unsubstituted phenyl side chain against the four cell lines except HCT-116 where **9b** > **9a** > **9d**. Moreover, derivatives **9b** demonstrated higher action than **9a** against HepG2, MCF-7, and HCT-116 but showed the same activity against A549. Finally, derivative **9d** with 1-propyl and unsubstituted phenyl side chain showed higher activities than compound **9e** with 1-propyl and the electron withdrawing 4-chlorophenyl side chain against the four tested cell lines. These findings enable us to conclude that substitution with electron donating groups at *para*-position showed higher activities than electron withdrawing ones.

2.5. ADMET; *in silico* studies profile

In silico report of the highly active derivatives **9b**, **9c** and **9e** was conducted for their physicochemical character evaluation and the proposed ADMET profile. It was predicted using pkCSM descriptor algorithm procedures⁶⁶ and matched to the rule of five described by Lipinski.⁶⁷ Good absorption properties were expected for the molecules that accomplish at least three rules: (i) no more than five hydrogen bond donors, (ii) no more than 10 hydrogen bond acceptors, (iii) molecular weight less than 500, (iv) not more than 5 for log *P*. In the current work, the standard anticancer agent sorafenib violate the log *P* rule while



our new compounds **9b**, **9c** and **9e** violate the molecular weight rule but erlotinib doesn't violate any rule.

As a result of obtaining data from Table 4, we can assume that compounds **9b**, **9c** and **9e** have very good GIT absorption in human (82.751 to 88.2) which indicates easier to cross different biological membranes.⁶⁸ So, they may show a significantly high bioavailability through GIT. Concerning CNS penetrability, our prepared compounds can reach CNS (CNS permeability values -2.266 to -2.535), comparable to that of sorafenib (-2.007) and erlotinib (-3.216).

It is well known that CYP3A4, the major drug-metabolizing enzyme, could be inhibited by sorafenib, erlotinib and also our derivatives **9b**, **9c** and **9e**. Elimination was expected depending on the total clearance which is a considerable factor in deciding dose intervals. The data showed that erlotinib confirmed higher clearance rates compared with sorafenib and our new compounds which demonstrated low clearance values. Thus, erlotinib could be eliminated faster, and as a result, supposed to have shorter dosing intervals. Unlike erlotinib, the prepared compounds exhibited slowly clearance rate, which signifies a long duration of action and extended dosing intervals. Toxicity is the final ADMET profile studied factor. Like presented in Table 4, sorafenib, erlotinib, and the novel compounds **9b**, **9c** and **9e** shared the drawback of unwanted hepatotoxic actions. Erlotinib, sorafenib, **9b**, **9c** and **9e** demonstrated high maximum tolerated dose. These involve the advantage of the broad therapeutic index of erlotinib, sorafenib, and our derivatives respectively. Lastly, the oral chronic toxic doses of the novel compounds **9b**, **9c** and **9e** are higher than that of sorafenib and erlotinib.

3. Conclusion

Fifteen new 1-alkyl-6-iodoquinazoline derivatives bearing different aliphatic, aromatic and/or heteroaromatic moieties were designed and synthesized as dual EGFR^{T790M}/VEGFR-2 inhibitors. The cytotoxic activities of the new compounds were screened *in vitro* against HepG2, MCF-7, HCT116 and A549 using an MTT assay. Their ability to bind with both EGFR and VEGFR-2 receptors was examined by molecular modeling. Compound **9c** showed the highest anticancer activities with EC₅₀ = 5.00, 6.00, 5.17 and 5.25 μM against HepG2, MCF-7, HCT116 and A549 cell lines correspondingly. Despite it showed lower activities than sorafenib (EC₅₀ = 4.00, 5.05, 5.58 and 4.04 μM) *versus* HepG2, MCF-7 and A549 contrarily subjacent performance against HCT116, it demonstrated higher actions than erlotinib (EC₅₀ = 7.73, 13.91, 8.20 and 5.49 μM), against the four tested cell lines. Moreover, compounds **5d**, **8b**, **9a**, **9b**, **9d**, and **9e** exhibited very good anticancer effects against the tested cancer cell lines. The highly effective seven derivatives **5d**, **8b**, **9a**, **9b**, **9c**, **9d** and **9e** were examined against VERO normal cell lines to estimate their cytotoxic capabilities. All conclusions revealed that each discovered compound possessed low toxicity regarding VERO normal cells with CC₅₀ extending from 45.50 to 55.00 μM . Additionally, all derivatives were evaluated for their ability as dual inhibitors of VEGFR-2 and EGFR^{T790M} in comparing to sorafenib and erlotinib as

standards with IC₅₀ = 0.84 and 0.24 respectively. Moreover, compounds **9c**, **9d**, **9e**, **5d**, **8b** and **9b** excellently inhibited EGFR^{T790M} activity with IC₅₀ = 0.22, 0.26, 0.30, 0.40, 0.45 and 0.50 μM respectively. Also, compounds **9c**, **9d** and **9e** excellently inhibited EGFR^{WT} activity with IC₅₀ = 0.15, 0.20 and 0.25 μM respectively in comparing to erlotinib (IC₅₀ = 0.18 μM). As planned, compound **9c** showed excellent dual EGFR/VEGFR-2 inhibitory activities. Consonantly, ADMET study was calculated *in silico* for the supreme three worthwhile compounds **9b**, **9c** and **9e** in contrasting to sorafenib and erlotinib as reference drugs.

4. Experimental

4.1. Chemistry

All melting points were carried out by open capillary method on a Gallen kamp Melting point apparatus at faculty of pharmacy Al-Azhar University and were uncorrected. The infrared spectra were recorded on pye Unicam SP 1000 IR spectrophotometer at Microanalytical Unit, Faculty of Pharmacy, Cairo University using potassium bromide disc technique. Proton magnetic resonance ¹H NMR spectra were recorded on a Bruker 400 MHz-NMR spectrophotometer at Microanalytical Center, Ain Shams University, and/or a Mercury 300 MHz-NMR spectrophotometer Faculty of Science, Cairo University. ¹³C NMR spectra were recorded on a Bruker 100 MHz-NMR spectrophotometer at Microanalytical Unit, Faculty of pharmacy, Cairo University. TMS was used as internal standard and chemical shifts were measured in δ scale (ppm). The mass spectra were carried out on Direct Probe Controller Inlet part to Single Quadropole mass analyzer in Thermo Scientific GCMS model ISQ LT using Thermo X-Calibur software at the Regional Center for Mycology and Biotechnology, Al-Azhar University. Elemental analyses (C, H, N) were performed on a CHN analyzer at Regional Center for Mycology and Biotechnology, Al-Azhar University. All compounds were within ± 0.4 of the theoretical values. The reactions were monitored by thin-layer chromatography (TLC) using TLC sheets precoated with UV fluorescent silica gel Merck 60 F254 plates and were visualized using UV lamp and different solvents as mobile phases.

Alkyl 4-(2-chloroacetamido)benzoate (**IIb,c**), 4-(2-chloroacetamido)-*N*-alkylbenzamide (**IIIa,b**), 5-iodoanthranilic acid (**2**), 5-iodo-*N*-alkylantranilic acid (**3a,b**), and 6-iodo-1-alkylquinazoline-2,4(1*H*,3*H*)-dione (**4a,b**), were synthesized according to the reported procedures.^{55–57}

4.1.1. Alkyl 4-{2-[1-alkyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2*H*)-yl]acetamido}benzoates (**5a–e**)

4.1.1.1. General method. A mixture of equimolar quantities of 6-iodo-1-ethylquinazoline-2,4(1*H*,3*H*)-dione (**4a**) and/or 6-iodo-1-propylquinazoline-2,4(1*H*,3*H*)-dione (**4b**) (0.01 mole) and the appropriate alkyl 4-(2-chloroacetamido)benzoate **IIb,c** (0.01 mole) in dry acetone (50 ml) was heated under reflux for 10 h in the presence of K₂CO₃ (1.38 g, 0.01 mole). The reaction mixture was filtered off while hot. The reaction mixture was allowed to attain room temperature and the precipitated solids were filtered, dried and crystallized from ethanol to give the corresponding target compounds, **5a–d** respectively.



4.1.1.2. Methyl 4-[2-[1-ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]acetamido]benzoate (**5a**). Yield, 74%; m.p. 250–2 °C; IR_{vmax} (cm⁻¹): 3484 (NH), 3053 (C–H aromatic), 2917 (C–H aliphatic), 1752, 1718, 1701, 1685 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.23 (t, 3H, NCH₂CH₃), 3.82 (s, 3H, COOCH₃), 4.14–4.21 (q, 2H, NCH₂CH₃), 4.80 (s, 2H, CH₂CO), 7.31–8.10 (m, 7H, 7Ar–H), 10.63 (s, 1H, NH, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 10.1, 36.5, 46.4, 50.4, 86.9, 115.8, 118.2, 121.4 (2), 127.2, 127.9 (2), 136.8, 140.0, 145.4, 150.3, 155.3, 158.3, 163.4, 168.4; MS (*m/z*): 507 (M⁺, 17.09%), 420 (80.04%), 294 (100% base), 268 (69.10%), 144 (70.06%); Anal. Calcd. For C₂₀H₁₈IN₃O₅ (m.w. 507.28): C, 47.35; H, 3.58; N, 8.28. Found: C, 47.30; H, 3.66; N, 8.33.

4.1.1.3. Ethyl 4-[2-[1-ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]acetamido]benzoate (**5b**). Yield, 75%; m.p. 257–9 °C; IR_{vmax} (cm⁻¹): 3481 (NH), 3046 (C–H aromatic), 2969 (C–H aliphatic), 1725, 1672, 1653 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.21–1.39 (2t, 6H, 2 CH₂CH₃), 4.16–4.19 (q, 2H, NCH₂CH₃), 4.27–4.29 (q, 2H, OCH₂CH₃), 4.80 (s, 2H, –CH₂CO), 7.32–8.10 (m, 7H, 7Ar–H), 10.62 (s, 1H, NH, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 14.3, 21.5, 33.9, 42.6, 62.2, 89.3, 119.6, 130.3 (2), 130.5, 130.8 (2), 134.7, 141.9, 147.7, 150.7, 154.8, 158.3, 165.4, 167.2, 167.4; Anal. Calcd. For C₂₁H₂₀IN₃O₅ (m.w. 521.31): C, 48.38; H, 3.87; N, 8.06. Found: C, 48.45; H, 3.75; N, 7.98.

4.1.1.4. Methyl 4-[2-[6-iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl]acetamido]benzoate (**5c**). Yield, 74%; m.p. 250–2 °C; IR_{vmax} (cm⁻¹): 3310 (NH), 3070 (C–H aromatic), 2922 (C–H aliphatic), 1722, 1697, 1666, 1650 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.29–1.34 (t, 3H, CH₂CH₂CH₃), 3.57 (s, 3H, COOCH₃), 3.79–3.83 (m, 2H, CH₂CH₂CH₃), 4.26–4.31 (t, 2H, CH₂CH₂CH₃), 4.82 (s, 2H, CH₂CO), 7.34–8.09 (m, 7H, 7Ar–H), 10.62 (s, 1H, NH, D₂O exchangeable); MS (*m/z*): 521 (M⁺, 35.30%), 498 (48.96%), 421 (100% base), 338 (79.89%), 235 (78.10%), 221 (85.87%); Anal. Calcd. For C₂₁H₂₀IN₃O₅ (m.w. 521.31): C, 48.38; H, 3.87; N, 8.06. Found: C, 48.50; H, 3.80; N, 7.94.

4.1.1.5. Ethyl 4-[2-[6-iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl]acetamido]benzoate (**5d**). Yield, 75%; m.p. 260–2 °C; IR_{vmax} (cm⁻¹): 3318 (NH), 3058 (C–H aromatic), 2925 (C–H aliphatic), 1733, 1686, 1671 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 0.92–0.97 (t, 3H, CH₂CH₂CH₃), 1.29–1.33 (t, 3H, OCH₂CH₃), 1.63–1.70 (m, 2H, CH₂CH₂CH₃), 4.07–4.12 (t, 2H, CH₂CH₂CH₃), 4.25–4.31 (q, 2H, OCH₂CH₃), 4.80 (s, 2H, CH₂CO), 7.31–8.04 (m, 7H, 7Ar–H), 10.62 (s, 1H, NH, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 9.4, 14.2, 23.1, 46.3, 56.3, 89.9, 112.5, 113.5, 114.4, 120.9 (2), 130.4 (2), 137.7, 141.4, 145.0, 150.4, 153.4, 154.8, 156.4, 164.3, 168.1; Anal. Calcd. For C₂₂H₂₂IN₃O₅ (m.w. 535.34): C, 49.36; H, 4.14; N, 7.85. Found: C, 49.53; H, 4.06; N, 8.02.

4.1.2. 4-[2-[1-Alkyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]acetamido]-*N*-substitutedbenzamide (6a,b)

4.1.2.1. General method. A mixture of equimolar quantities of 6-iodo-1-alkylquinazolin-2,4(1*H*,3*H*)-dione (**4a,b**) (0.01 mole) and the appropriate 4-(2-chloroacetamido)-*N*-substitutedbenzamide (**IIIa,b**) (0.01 mole) in dry acetone (50 ml)

was heated under reflux for 12 h in the presence of K₂CO₃ (1.38 g, 0.01 mole). The reaction mixture was filtered off while hot. The reaction mixture was concentrated and the afforded precipitated solids were filtered, dried and crystallized from ethanol to give the corresponding target compounds **6a,b**.

4.1.2.2. 4-[2-[1-Ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]acetamido]-*N*-ethylbenzamide (6a). Yield, 75%; m.p. 273–5 °C; IR_{vmax} (cm⁻¹): 3315, 3283 (2NH), 3064 (C–H aromatic), 2922 (C–H aliphatic), 1703, 1659, 1632 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.21–1.25 (t, 3H, NHCH₂CH₃), 1.30–1.33 (t, 3H, NCH₂CH₃), 4.16–4.24 (q, 2H, NHCH₂CH₃), 4.27–4.31 (q, 2H, NCH₂CH₃), 4.81 (s, 2H, CH₂CO), 7.32–8.10 (m, 7H, 7Ar–H), 9.73 (s, 1H, NHCH₂CH₃, D₂O exchangeable), 10.63 (s, 1H, CONH, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 9.1, 13.7, 32.8, 43.7, 55.4, 91.9, 113.9, 119.53, 120.1 (2), 121.1, 126.8, 127.1 (2), 129.9, 132.9, 134.5, 138.9, 141.5, 166.2, 168.0; Anal. Calcd. For C₂₁H₂₁IN₄O₄ (m.w. 520.33): C, 48.48; H, 4.07; N, 10.77. Found: C, 48.40; H, 3.95; N, 10.80.

4.1.2.3. 4-[2-[6-Iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl]acetamido]-*N*-methylbenzamide (6b). Yield, 71%; m.p. 276–8 °C; IR_{vmax} (cm⁻¹): 3285 (2NH), 3062 (C–H aromatic), 2907 (C–H aliphatic), 1734, 1686, 1667 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 0.91–0.96 (t, 3H, CH₂CH₂CH₃), 1.59–1.66 (m, 2H, CH₂CH₂CH₃), 3.83 (s, 3H, COOCH₃), 3.96–4.02 (t, 2H, CH₂CH₂CH₃), 4.30 (s, 2H, CH₂CO), 7.23–8.02 (m, 7H, 7Ar–H), 10.60 (s, 1H, NHCH₃, D₂O exchangeable), 11.50 (s, 1H, CONH, D₂O exchangeable); MS (*m/z*): 522 (M⁺+2, 1.87%), 521 (M⁺+1, 13.78%), 520 (M⁺, 26.11%), 480 (53.91%), 352 (100% base), 293 (52.06%), 157 (70.02%); Anal. Calcd. For C₂₁H₂₁IN₄O₄ (m.w. 520.33): C, 48.48; H, 4.07; N, 10.77. Found: C, 48.58; H, 4.15; N, 10.59.

4.1.3. 2-(1-Alkyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl)-*N*-(4-(hydrazinecarbonyl)phenyl)-acetamide (7a,b)

4.1.3.1. General method. Hydrazine hydrate 100% (0.5 ml, 0.01 mol) was added dropwise to a stirred solution of the appropriate ethyl ester **5b** and/or **5d** (0.002 mol) in absolute ethanol (20 ml). The mixture was stirred well and heated at 70 °C for 6 h. The reaction mixture was concentrated, cooled and the crude product was filtered and recrystallized from ethanol to afford the corresponding hydrazide derivatives **7a,b** respectively.

4.1.3.2. 2-(1-Ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl)-*N*-(4-(hydrazinecarbonyl)phenyl)-acetamide (7a). Yield, 75%; m.p. 300–2 °C; IR_{vmax} (cm⁻¹): 3250, 3190 (2NH & NH₂), 3036 (C–H aromatic), 2927 (C–H aliphatic), 1710, 1685 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.23 (t, 3H, NCH₂CH₃), 4.16–4.19 (q, 2H, NCH₂CH₃), 4.42 (s, 2H, NH₂), 4.80 (s, 2H, CH₂CO), 7.31–8.10 (m, 7H, 7Ar–H), 9.63 (s, 1H, NHNH₂, D₂O exchangeable), 10.48 (s, 1H, NHPh, D₂O exchangeable); Anal. Calcd. For C₁₉H₁₈IN₅O₄ (m.w. 507.29): C, 44.99; H, 3.58; N, 13.81. Found: C, 45.12; H, 3.65; N, 13.70.

4.1.3.3. *N*-(4-(hydrazinecarbonyl)phenyl)-2-(6-iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl)acetamide (7b). Yield, 70%; m.p. 306–8 °C; IR_{vmax} (cm⁻¹): 3436, 3234, 3179 (2NH & NH₂), 3038 (C–H aromatic), 2915 (C–H aliphatic), 1685, 1670 (4C=O); ¹³CNMR (100 MHz, DMSO-*d*₆): 10.7, 22.9, 32.8, 46.6, 55.4, 90.7, 118.1, 126.7 (2), 127.4 (2), 135.5, 140.0, 143.1, 147.4,



152.3, 156.6, 161.7, 166.7, 171.6; Anal. Calcd. For $C_{20}H_{20}IN_5O_4$ (m.w. 521.32): C, 46.08; H, 3.87; N, 13.43. Found: C, 45.90; H, 3.96; N, 13.37.

4.1.4. 2-[1-Alkyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]-N-[4-(5-mercapto-1,3,4-oxadiazol-2-yl)phenyl]acetamides (8a,b). The appropriate hydrazide derivative **7a** and/or **7b** (0.01 mol) was dissolved in a solution of potassium hydroxide (0.56 g, 0.01 mole) in ethanol (40 ml) and water (2 ml). Carbon disulphide (3 ml, 4.61 g, 0.06 mol) was then added while stirring and the reaction mixture was refluxed for 24 h. The reaction mixture was concentrated, cooled to room temperature and acidified with dil.HCl. The obtained precipitated solids were filtered, washed with water and crystallized from ethanol to give the targeted compounds **8a,b** respectively.

4.1.4.1. 2-[1-Ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]-N-[4-(5-mercapto-1,3,4-oxadiazol-2-yl)phenyl]acetamide (8a). Yield, 74%; m.p. 283–5 °C; IR_{vmax} (cm⁻¹): 3301, 3192 (2NH), 3036 (C–H aromatic), 2919 (C–H aliphatic), 1689, 1665, 1643 (3C=O), 1263 (C=S); ¹HNMR (400 MHz, DMSO-*d*₆): 1.21–1.26 (t, 3H, CH₂CH₃), 4.17–4.19 (q, 2H, CH₂CH₃), 4.82 (s, 2H, COCH₂), 7.32–8.11 (m, 7H, 7Ar–H), 10.66 (s, 1H, NH, D₂O exchangeable), 14.59 (s, 1H, SH, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 12.3, 34.9, 52.1, 91.6, 114.8, 116.9, 122.8, 123.6, 124.6 (2), 125.4, 128.9 (2), 129.4, 139.5, 148.5, 153.1, 157.4, 163.7, 170.1; Anal. Calcd. for $C_{20}H_{16}IN_5O_4S$ (m.w. 549.34): C, 43.73; H, 2.94; N, 12.75; Found: C, 43.90; H, 3.06; N, 12.65.

4.1.4.2. 2-[6-Iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl]-N-[4-(5-mercapto-1,3,4-oxadiazol-2-yl)phenyl]acetamide (8b). Yield, 68%; m.p. 287–9 °C; IR_{vmax} (cm⁻¹): 3447, 3189 (2NH), 3050 (C–H aromatic), 2967 (C–H aliphatic), 1692, 1653 (3C=O), 1262 (C=S); ¹HNMR (400 MHz, DMSO-*d*₆): 0.92–0.97 (t, 3H, CH₂CH₂CH₃), 1.63–1.70 (m, 2H, CH₂CH₂CH₃), 4.07–4.12 (t, 2H, CH₂CH₂CH₃), 4.81 (s, 2H, COCH₂), 7.31–8.09 (m, 7H, 7Ar–H), 10.66 (s, 1H, NH, D₂O exchangeable), 14.59 (s, 1H, SH, D₂O exchangeable); MS (*m/z*): 565 (M⁺+2, 3.96%), 564 (M⁺+1, 9.96%), 520 (M⁺, 60.00%), 443 (85.24%), 334 (80.01%), 328 (100% base), 233 (79.97%); Anal. Calcd. for $C_{21}H_{18}IN_5O_4S$ (m.w. 563.37): C, 44.77; H, 3.22; N, 12.43. Found: C, 44.84; H, 3.11; N, 12.56.

4.1.5. 2-[1-Alkyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]-N-[4-(2-(un)substitutedbenzylidene-hydrazine-1-carbonyl)phenyl]acetamides (9a–e)

4.1.5.1. General method. Equimolar amounts of appropriate hydrazide **7a** and/or **7b** (0.002 mol) and the appropriate aromatic aldehyde namely benzaldehyde, 2-chlorobenzaldehyde, 4-chlorobenzaldehyde, 4-methylbenzaldehyde and 4-methoxybenzaldehyde (0.002 mol) were refluxed in gl. acetic acid (25 ml) for 4 hours and the reaction was followed up by TLC. After completion of the reaction, the mixture was cooled, poured into water, filtered and crystallized from ethanol to afford the corresponding target compounds **9a–e** respectively.

4.1.5.2. N-[4-(2-benzylidenehydrazine-1-carbonyl)phenyl]-2-[1-ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]acetamide (9a). Yield, 72%; m.p. 290–2 °C; IR_{vmax} (cm⁻¹): 3262 (2NH), 3050 (C–H aromatic), 2950 (C–H aliphatic), 1722, 1697, 1666, 1652 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.24–1.31 (t, 3H,

CH₂CH₃), 4.18–4.20 (q, 2H, CH₂CH₃), 4.82 (s, 2H, COCH₂), 7.32–8.12 (m, 12H, 12Ar–H), 8.45 (s, 1H, –N=CH), 10.68 (s, 1H, NHCO, D₂O exchangeable), 11.75 (s, 1H, N–NHCO, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 13.4, 34.9, 56.6, 91.7, 114.7, 116.8, 121.8 (2), 122.4, 123.6, 124.3, 124.5 (2), 125.6 (2), 128.9, 129.4, 129.5 (2), 140.0, 148.5, 152.3, 156.4, 160.0, 164.6, 168.7; Anal. Calcd. for $C_{26}H_{22}IN_5O_4$ (m.w. 595.40): C, 52.45; H, 3.72; N, 11.76. Found: C, 52.35; H, 3.78; N, 11.70.

4.1.5.3. 2-[1-Ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]-N-[4-[2-(4-methylbenzylidene)hydrazine-1-carbonyl]phenyl]acetamide (9b). Yield, 72%; m.p. 297–9 °C; IR_{vmax} (cm⁻¹): 3313, 3255 (2NH), 3033 (C–H aromatic), 2922 (C–H aliphatic), 1700, 1675 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.22–1.26 (t, 3H, CH₂CH₃), 2.34 (s, 3H, CH₃), 4.17–4.20 (q, 2H, CH₂CH₃), 4.82 (s, 2H, COCH₂), 7.25–8.11 (m, 11H, 11Ar–H), 8.42 (s, 1H, –N=CH), 10.57 (s, 1H, NHCO, D₂O exchangeable), 11.67 (s, 1H, N–NHCO, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 13.4, 21.8, 56.6, 87.4, 114.6 (2), 116.6 (2), 118.8, 121.8 (2), 123.9, 124.3, 125.6, 128.9, 129.3 (2), 129.5, 140.0, 143.3, 145.7, 147.7, 153.7, 156.6, 159.4, 160.9, 170.1; Anal. Calcd. for $C_{27}H_{24}IN_5O_4$ (m.w. 609.42): C, 53.21; H, 3.97; N, 11.49. Found: C, 53.14; H, 4.15; N, 11.65.

4.1.5.4. 2-[1-Ethyl-6-iodo-2,4-dioxo-1,4-dihydroquinazolin-3(2H)-yl]-N-[4-[2-(4-methoxybenzylidene)hydrazine-1-carbonyl]phenyl]acetamide (9c). Yield, 70%; m.p. 302–4 °C; IR_{vmax} (cm⁻¹): 3262, 3184 (2NH), 3036 (C–H aromatic), 2932 (C–H aliphatic), 1733, 1665, 1641 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 1.22–1.27 (t, 3H, CH₂CH₃), 3.81 (s, 3H, OCH₃), 4.18–4.20 (q, 2H, CH₂CH₃), 4.82 (s, 2H, COCH₂), 7.27–8.11 (m, 11H, 11Ar–H), 8.42 (s, 1H, –N=CH), 10.58 (s, 1H, NHCO, D₂O exchangeable), 11.75 (s, 1H, N–NHCO, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 14.7, 33.8, 50.6, 62.0, 87.4, 99.3, 113.6 (2), 115.9, 117.1, 123.7 (2), 131.2 (2), 134.9 (2), 136.5, 137.7, 138.8, 150.0, 151.0, 154.8, 158.4, 162.9, 165.6, 169.9, 178.5; MS (*m/z*): 626 (M⁺+1, 41.76%), 625 (M⁺, 65.91%), 558 (74.18%), 353 (100% base), 334 (81.89%), 289 (74.26%); Anal. Calcd. for $C_{27}H_{24}IN_5O_5$ (m.w. 625.42): C, 51.85; H, 3.87; N, 11.20. Found: C, 52.00; H, 3.58; N, 11.40.

4.1.5.5. N-[4-(2-benzylidenehydrazine-1-carbonyl)phenyl]-2-[6-iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl]acetamide (9d). Yield, 65%; m.p. 307–9 °C; IR_{vmax} (cm⁻¹): 3447, 3283 (2NH), 3063 (C–H aromatic), 2979 (C–H aliphatic), 1692, 1658, 1653 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 0.93–1.02 (t, 3H, CH₂CH₂CH₃), 1.64–1.71 (m, 2H, CH₂CH₂CH₃), 4.08–4.13 (t, 2H, CH₂CH₂CH₃), 4.81 (s, 2H, COCH₂), 7.32–8.11 (m, 12H, 12Ar–H), 8.45 (s, 1H, –N=CH), 10.57 (s, 1H, NHCOCH₂, D₂O exchangeable), 11.74 (s, 1H, N–NHCO, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 11.8, 21.7, 47.3, 56.8, 89.9, 112.8 (2), 116.3, 124.5 (2), 127.8, 129.1 (2), 130.5 (2), 131.3, 138.3, 138.9, 146.6, 150.6, 153.7, 157.4, 157.9, 161.5, 165.3, 168.7, 171.6; Anal. Calcd. for $C_{27}H_{24}IN_5O_4$ (m.w. 609.42): C, 53.21; H, 3.97; N, 11.49. Found: C, 53.30; H, 4.05; N, 11.42.

4.1.5.6. N-[4-[2-(4-chlorobenzylidene)hydrazine-1-carbonyl]phenyl]-2-[6-iodo-2,4-dioxo-1-propyl-1,4-dihydroquinazolin-3(2H)-yl]acetamide (9e). Yield, 70%; m.p. >310 °C; IR_{vmax} (cm⁻¹): 3316, 3280 (2NH), 3073 (C–H aromatic), 2950 (C–H aliphatic), 1701, 1659 (4C=O); ¹HNMR (400 MHz, DMSO-*d*₆): 0.93–0.98 (t, 3H,



CH₂CH₂CH₃), 1.64–1.69 (m, 2H, CH₂CH₂CH₃), 4.08–4.13 (t, 2H, CH₂CH₂CH₃), 4.82 (s, 2H, COCH₂), 7.32–8.11 (m, 11H, 11Ar-H), 8.86 (s, 1H, -N=CH), 10.59 (s, 1H, NHCOCH₂, D₂O exchangeable), 11.98 (s, 1H, N-NHCO, D₂O exchangeable); ¹³CNMR (100 MHz, DMSO-*d*₆): 10.9, 21.3, 46.8, 54.9, 96.8, 113.9 (2), 118.9, 122.8 (2), 123.6 (2), 124.4, 125.6, 129.7, 129.9 (2), 131.8, 142.7, 143.5, 148.8, 152.5, 153.2, 160.0, 169.4; MS (*m/z*): 645 (M⁺+2, 14.69%), 643 (M⁺, 44.72%), 580 (58.22%), 492 (60.98%), 471 (100% base), 367 (46.13%), 87 (82.00%); Anal. Calcd. for C₂₅-H₁₉ClIN₅O₄ (m.w. 643.87): C, 50.37; H, 3.60; N, 10.88. Found: C, 50.22; H, 3.74; N, 11.08.

4.2. Docking studies

Both VEGFR-2 (PDB ID 4ASD)^{48,58} and EGFR^{T790M} (PDB ID 3W2O)^{55,56} were used by Molsoft program to carry out docking studies.

4.3. Biological testing

4.3.1. *In vitro* anti-cancer activity. Our derivatives were tested against four cell lines, HCT-116, HepG2, MCF-7 and A549 using MTT colorimetric assay.^{59–61}

4.3.2. *In vitro* VEGFR-2 assay. All compounds were assessed for their inhibitory activities against VEGFR-2.^{62,63}

4.3.3. EGFR^{T790M} assay. Using the HTRF assay, all drugs were assessed for their ability to inhibit mutant EGFR^{T790M}.^{63,64}

4.3.4. EGFR^{WT} assay. The *in vitro* inhibitory activities of some synthesized compounds against EGFR^{WT} were carried out using EGFR Kinase Assay Kit (BPS biosciences).⁶⁵

Author contributions

K. El-Adl, Samir A. Salama, Abeer Mohamed, Ahmed Aljohani, Ahmed El-morsy, Doaa Keshek and Marwa Alsulaimany were responsible for the conception and rational design of the work. K. El-Adl, Abeer Mohamed, Ahmed Aljohani, Omar Alatawi, Hussam Alharbi, Majed Aljohani and Marwa Alsulaimany were responsible for the data collection and synthesis of the new compounds. K. El-Adl performed the molecular docking study. Hussam Alharbi, Majed Aljohani, Ahmed El-morsy, Ahmed Aljohani, Marwa Alsulaimany and K. El-Adl were responsible for spectral data analysis. Doaa Keshek, Sara Almadani, Abdulrahman Alsimaree, Samir A. Salama and K. El-Adl conducted the cytotoxicity assay. Sara Almadani, Abdulrahman Alsimaree, Samir A. Salama and K. El-Adl conducted the *in silico* pharmacokinetic study. All authors discussed the results and contributed to the writing and revision of the original manuscript.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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