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Novel pyridine-heterocycle hybrids: synthesis *via* Hantzsch and Biginelli reactions, docking simulations, and anticancer activity

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Cancer is a serious global health issue and remains one of the top causes of death worldwide. To overcome the problems of the existing anticancer drugs in terms of specificity and resistance, a new class of hybrid bis-heterocyclic compounds with a pyridine bridge has been designed and synthesized *via* the Hantzsch reaction. The results of elemental analysis and spectral data were used for the confirmation of the synthesized compounds. Among the tested analogs, the highest cytotoxic activity was shown by compound 7 against the HepG2, A549, and MCF7 cancer cell lines, with IC₅₀ values of 18.07, 14.45, and 30.89 μg mL⁻¹, respectively, while the cytotoxicity against normal fibroblasts was negligible with an IC₅₀ greater than 100 μg mL⁻¹. The structure–activity relationship results emphasized the key role of the molecular planarity and sulfur atom substitution. The results from molecular docking, molecular dynamics simulations, and MM/PBSA and MM/GBSA binding free energy calculations indicated a strong binding of compound 7 with EGFR. The results of the EGFR inhibition assay further strengthened the binding of the EGFR with the mentioned compound. The results from the ADME and toxicological analyses indicated a good pharmacokinetic and safety profile for the compound.

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1 Introduction

Cancer, a disease characterized by the unregulated growth of cells that can invade other parts of the body, is identified as one of the most perilous threats to the world's health. According to the GLOBOCAN 2022 database, there were 19.9 million incident cases of cancer as well as 9.6 million deaths due to cancer worldwide. Cancer has proven itself to be one of the principal causes of morbidity as well as mortality rates in the global scenario. Future years will experience a tremendous rise in the incidence of cancer in the global scenario.¹ Consequently, it is essential to seek out innovative medications that possess high efficacy, minimal toxicity, and low mutagenicity, along with targeted anticancer effects that can effectively combat the resistance commonly developed against existing treatments. Surgery, radiation therapy, and chemotherapy can be utilized separately or in combination with cancer treatments.^{2,3}

Heterocyclic anticancer drugs represent a significant category of medications used in cancer treatments. They are extensively researched in medicinal chemistry because of their various biological effects, particularly their crucial involvement in the development of anticancer drugs. These substances are essential in cancer therapy because of their capacity to target different biological pathways, such as DNA replication, enzyme inhibition, and the induction of apoptosis. The kind or dimension of the ring of heterocycles, together with the substituent groups and structural variations, allows for interactions with distinctly different structural configurations of enzyme binding sites, which significantly affect the physico-chemical traits and, in turn, the pharmacological properties of the molecules.^{4,5} Among various heterocycles, pyridine derivatives have attracted considerable interest as possible anticancer agents because of their capacity to engage with diverse biological targets, such as enzymes, receptors, and DNA.^{6–10}

Fused heterocyclic compounds containing a pyridine element also exhibit a variety of biological activities across different areas. Generally, substances with pyridine and dihydropyridine are predominantly used for diverse applications, including combating microbes, viruses, cancers, hypertension, diabetes, malaria, and inflammation, while also serving as antioxidants and psychopharmacological antagonists, and addressing amoebic infections.^{11–15} These structures are

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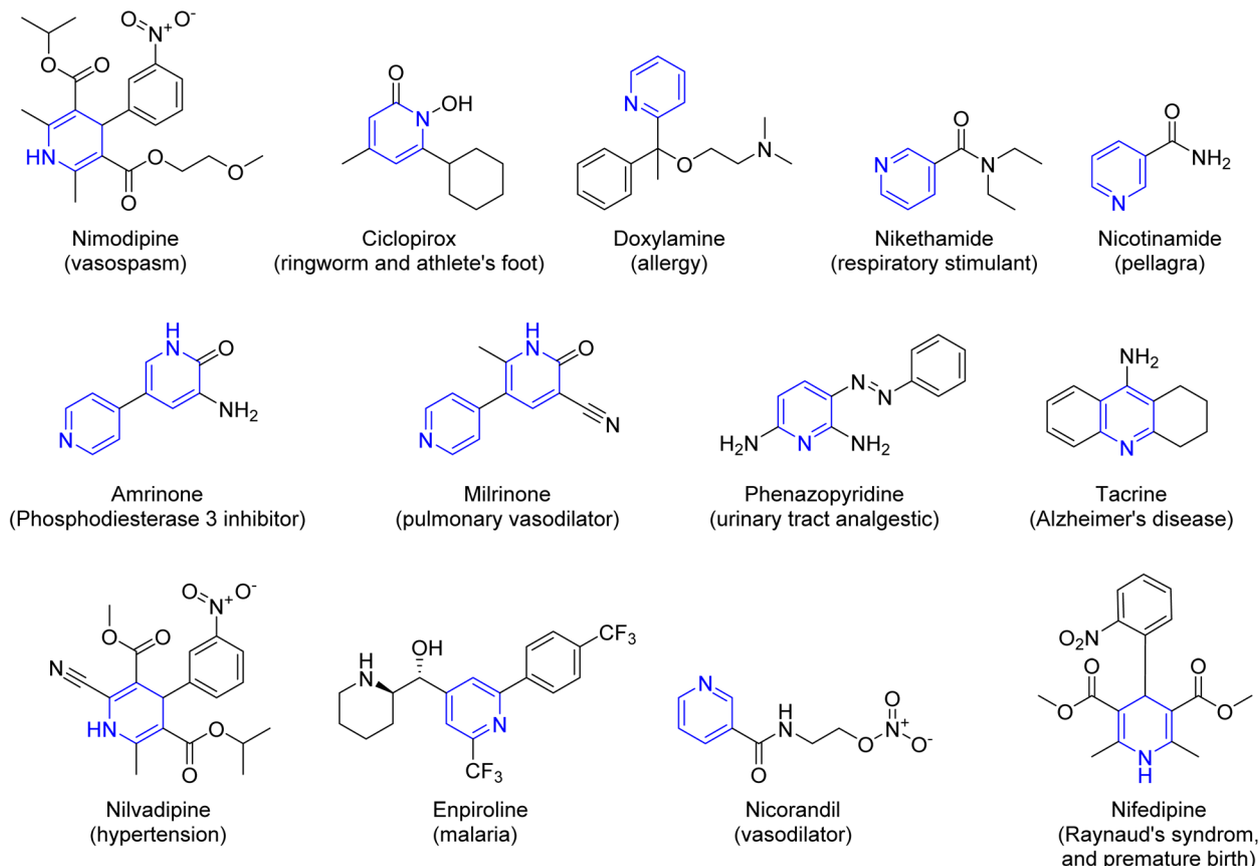


Fig. 1 Structures of some approved pyridine-containing medications on the market.

essential components in the chemical frameworks of many approved drugs available today (Fig. 1).

Molecular hybridization is a method in drug design and development that entails merging pharmacophoric components from different bioactive substances to produce a new hybrid compound with improved affinity and efficacy relative to the original drugs. In recent years, significant advancements have occurred in this domain, with several hybrid compounds featuring strong heterocycles frequently found in the drug discovery field within the pharmaceutical and agrochemical industries.^{16,17} This study details the creation of new hybrid compounds, including pyridine-derived polyhydroquinoline, bis(hexahydroacridine), bis(hexahydropyrimido[4,5-*b*]quinoline), and bis(tetrahydropyrazolo[3,4-*b*:4',3'-*e*]pyridine), using Hantzsch reactions. The cytotoxic effects and *in vitro* anticancer activities of the newly created compounds were evaluated against chosen human cancer cell lines. This study builds upon our earlier investigations into generating bioactive heterocycles and bis(heterocycles).^{18–46}

2 Experimental

2.1 Chemistry

2.1.1 General. All reagents were purchased at the highest available purity from Sigma-Aldrich and were used without further purification. Melting points were measured on

a Gallenkamp melting point apparatus. Elemental analyses were carried out at the Microanalytical Center of Cairo University, Giza, Egypt. IR spectra were obtained on Shimadzu FT-IR 8101 PC infrared spectrophotometers (Shimadzu, Tokyo, Japan) using KBr disks. NMR spectra were recorded on a JEOL JNM-LA500 spectrometer, operating at 500 MHz for ¹H NMR, and 125.65 MHz for ¹³C NMR using DMSO-*d*₆ as solvent. Chemical shifts are given in parts per million and are related to that of the solvent. Mass spectra were recorded on a Bruker Daltonics spectrometer.

2.1.1.1 Diethyl 4,4'-(((pyridine-2,6-diylbis(methylene))bis(oxy))bis(4,1-phenylene))bis(3,7,7-trimethyl-5-oxo-1,4,5,6,7,8-hexahydroquinoline-2-carboxylate) (7). A mixture of bis-aldehyde **4** (0.35 g, 1 mmol), dimedone (**5**) (0.28 g, 2 mmol), ethyl acetoacetate (**6**) (0.26 g, 2 mmol) was heated in absolute ethanol (30 mL) containing ammonium acetate (0.15 g, 2 mmol) for 6 h. The excess solvent was evaporated under reduced pressure, and the collected residue was recrystallized from EtOH.

Yellow powder (0.67 g, 83%); Mp 272–274 °C; IR (KBr): ν 3279 (NH), 1697 (C=O ester), 1643 (C=O ketone) cm⁻¹; ¹H NMR (500 MHz, DMSO-*d*₆) δ 0.82 (s, 6H, 2CH₃), 0.97 (s, 6H, 2CH₃), 1.09 (t, *J* = 7.1 Hz, 6H, 2CH₃CH₂COO), 1.93 (d, 2H, H8, *J* = 15 Hz) 2.11 (d, 2H, H8, *J* = 15 Hz), 2.25 (s, 6H, 2CH₃), 2.28–2.39 (m, 4H, 2CH₂), 3.94 (q, *J* = 6.9 Hz, 4H, 2CH₃CH₂COO), 4.77 (s, 2H, 2CH), 5.05 (s, 4H, 2OCH₂), 6.82 (d, *J* = 8.4 Hz, 4H, Ar-H), 7.04 (d, *J* = 8.2 Hz, 4H, Ar-H), 7.39 (d, *J* = 7.8 Hz, 2H, pyridine-H), 7.80 (m, *J* =



7.9 Hz, 1H, pyrdine-H), 9.03 (s, 2H, 2NH) ppm; ^{13}C NMR (126 MHz, DMSO- d_6) δ 14.7, 18.8, 27.1, 29.6, 32.7, 35.6, 40.9, 50.9, 59.6, 70.7, 104.4, 110.7, 114.5, 121.1, 129.1, 138.3, 141.1, 145.3, 149.9, 156.8, 157.1, 167.5, 194.9 ppm; MS (EI, 70 eV): m/z (%) 813 [M^+]; anal. calcd for $\text{C}_{49}\text{H}_{55}\text{N}_3\text{O}_8$: C, 72.30; H, 6.81; N, 5.16. Found: C, 72.26; H, 6.78; N, 5.08%.

2.1.1.2 9,9'-(((Pyridine-2,6-diylbis(methylene))bis(oxy))bis(4,1-phenylene))bis(3,3,6,6-tetramethyl-3,4,6,7,9,10-hexahydroacridine-1,8(2H,5H)-dione) (**8**). A mixture of bis-aldehyde **4** (0.35 g, 1 mmol) and dimedone (**5**) (0.56 g, 4 mmol) was heated in glacial acetic acid (20 mL) containing ammonium acetate (0.15 g, 2 mmol) for 6 h. The excess solvent evaporated under reduced pressure, and the crude product was purified by washing with hot EtOH.

Creamy powder (0.73 g, 88%); Mp 285–287 °C; IR (KBr): $\bar{\nu}$ 3289 (NH), 1636 (C=O) cm^{-1} ; ^1H NMR (500 MHz, DMSO- d_6) δ 0.84 (s, 12H, 4CH₃), 0.97 (s, 12H, 4CH₃), 1.94–2.15 (m, 8H, 4CH₂), 2.26–2.44 (m, 8H, 4CH₂), 4.73 (s, 2H, 2CH), 5.04 (s, 4H, 2OCH₂), 6.79 (d, J = 8.2 Hz, 4H, Ar-H), 7.04 (d, J = 8.2 Hz, 4H, Ar-H), 7.39 (d, J = 7.8 Hz, 2H, pyrdine-H), 7.80 (t, J = 7.8 Hz, 1H, pyrdine-H), 9.23 (s, 2H, 2NH).ppm; ^{13}C NMR (126 MHz, DMSO- d_6) δ 27.1, 29.6, 32.5, 32.7, 40.7, 50.9, 70.7, 112.2, 114.4, 121.1, 129.2, 138.3, 140.6, 149.7, 156.6, 157.1, 195.0 ppm; MS (EI, 70 eV): m/z (%) 833 [M^+]; anal. calcd for $\text{C}_{53}\text{H}_{59}\text{N}_3\text{O}_6$: C, 76.32; H, 7.13; N, 5.04. Found: C, 76.24; H, 7.09; N, 5.01%.

2.1.1.3 9,9'-(((Pyridine-2,6-diylbis(methylene))bis(oxy))bis(4,1-phenylene))bis(3,3,6,6-tetramethyl-10-(*p*-tolyl)-3,4,6,7,9,10-hexahydroacridine-1,8(2H,5H)-dione) (**11**). *Method A*. A mixture of bis-aldehyde **4** (0.35 g, 1 mmol), dimedone (**5**) (0.28 g, 2 mmol), and enaminone **10** (0.46 g, 2 mmol) was heated in pyridine (10 mL) for 6 h. The precipitated solid was collected and washed thoroughly with hot EtOH.

Method B. A mixture of bis-aldehyde **4** (0.35 g, 1 mmol), dimedone (**5**) (0.56 g, 4 mmol), *p*-toluidine (**12**) (0.21 g, 2 mmol) was heated in pyridine (10 mL) for 6 h. The precipitated solid was collected and washed thoroughly with hot EtOH.

Yellow crystals (0.75 g, 74%, method A; 0.72 g, 71%, method B); Mp 300 < °C; IR (KBr): $\bar{\nu}$ 1636 (C=O) cm^{-1} ; ^1H NMR (500 MHz, DMSO- d_6) δ 0.68 (s, 12H, 4CH₃), 0.84 (s, 12H, 4CH₃), 1.72–1.98 (m, 8H, 4CH₂), 2.12–2.17 (m, 8H, 4CH₂), 2.38 (s, 6H, 2CH₃), 4.96 (s, 2H, 2CH), 5.08 (s, 4H, 2OCH₂), 6.88 (d, J = 8.4 Hz, 4H, Ar-H), 7.19 (d, J = 8.2 Hz, 8H, Ar-H), 7.37 (d, J = 8.0 Hz, 4H, Ar-H), 7.41 (d, J = 7.7 Hz, 2H, pyrdine-H), 7.82 (t, J = 7.8 Hz, 1H, pyrdine-H).ppm; MS (EI, 70 eV): m/z (%) 1013 [M^+]; anal. calcd for $\text{C}_{67}\text{H}_{71}\text{N}_3\text{O}_6$: C, 79.34; H, 7.06; N, 4.14. Found: C, 79.28; H, 7.03; N, 4.18%.

2.1.2 General procedure for synthesis of compounds 15 and 16. A mixture of bis-aldehyde **4** (0.35 g, 1 mmol), dimedone (**5**) (0.28 g, 2 mmol), and 6-aminouracil (0.25 g, 2 mmol) or 6-aminothiouracil (0.29 g, 2 mmol) was heated in glacial acetic acid (20 mL) for 6 h. The precipitated solid was collected and washed thoroughly with hot EtOH.

2.1.2.1 5,5'-(((Pyridine-2,6-diylbis(methylene))bis(oxy))bis(4,1-phenylene))bis(8,8-dimethyl-5,8,9,10-tetrahydropyrimido[4,5-*b*]quinoline-2,4,6(1H,3H,7H)-trione) (**15**). Creamy powder (0.62 g, 77%); Mp 300 < °C; IR (KBr): $\bar{\nu}$ 3408, 3269, 3192 (3NH), 1717 (C=O ketone), 1665, 1655 (2C=O amide) cm^{-1} ; ^1H NMR (500 MHz,

DMSO- d_6) δ 0.86 (s, 6H, 2CH₃), 0.98 (s, 6H, 2CH₃), 1.97–2.17 (m, 4H, 2CH₂), 2.40–2.45 (m, 4H, 2CH₂), 4.67 (s, 2H, 2CH), 5.06 (s, 4H, 2OCH₂), 6.82 (d, J = 8.2 Hz, 4H, Ar-H), 7.07 (d, J = 8.2 Hz, 4H, Ar-H), 7.39 (d, J = 7.8 Hz, 2H, pyrdine-H), 7.81 (t, J = 7.8 Hz, 1H, pyrdine-H), 8.81 (s, 2H, 2NH), 10.26 (s, 2H, 2NH), 10.65 (s, 2H, 2NH).ppm; ^{13}C NMR (126 MHz, DMSO- d_6) δ 27.1, 29.5, 32.7, 32.7, 39.5, 39.7, 39.9, 40.0, 40.2, 40.4, 40.5, 40.8, 50.7, 70.7, 90.4, 100.0, 112.0, 114.5, 121.0, 129.1, 138.4, 139.9, 149.5, 156.9, 157.1, 163.3, 163.3, 194.9 ppm; MS (EI, 70 eV): m/z (%) 809 [M^+]; anal. calcd for $\text{C}_{45}\text{H}_{43}\text{N}_7\text{O}_8$: C, 66.74; H, 5.35; N, 12.11. Found: C, 66.71; H, 5.27; N, 12.05%.

2.1.2.2 5,5'-(((Pyridine-2,6-diylbis(methylene))bis(oxy))bis(4,1-phenylene))bis(8,8-dimethyl-2-thioxo-2,3,5,8,9,10-hexahydropyrimido[4,5-*b*]quinoline-4,6(1H,7H)-dione) (**16**). Pale yellow powder (0.72 g, 86%); Mp 300 < °C; IR (KBr): $\bar{\nu}$ 3412, 3263, 3200 (3NH), 1670 (C=O ketone), 1653 (C=O amide) cm^{-1} ; ^1H NMR (500 MHz, DMSO- d_6) δ 0.87 (s, 6H, 2CH₃), 0.99 (s, 6H, 2CH₃), 1.98–2.20 (m, 4H, 2CH₂), 2.34–2.47 (m, 4H, 2CH₂), 4.68 (s, 2H, 2CH), 5.07 (s, 4H, 2OCH₂), 6.84 (d, J = 8.4 Hz, 4H, Ar-H), 7.09 (d, J = 8.1 Hz, 4H, Ar-H), 7.39 (d, J = 7.6 Hz, 2H, pyrdine-H), 7.81 (t, J = 7.8 Hz, 1H, pyrdine-H), 8.43 (s, 2H, 2NH), 11.64 (s, 2H, 2NH), 12.14 (s, 2H, 2NH).ppm; ^{13}C NMR (126 MHz, DMSO- d_6) δ 27.1, 29.4, 32.4, 32.7, 40.7, 50.7, 70.7, 94.9, 100.1, 111.8, 114.6, 121.0, 129.2, 138.4, 139.1, 143.9, 149.0, 157.1, 160.7, 173.9, 194.9 ppm; MS (EI, 70 eV): m/z (%) 841 [M^+]; anal. calcd for $\text{C}_{45}\text{H}_{43}\text{N}_7\text{O}_6\text{S}_2$: C, 64.19; H, 5.15; N, 11.64. Found: C, 64.12 H, 5.10; N, 11.60%.

2.1.2.3 2,6-Bis((4-(3,5-dimethyl-1,4,7,8-tetrahydropyrazolo[3,4-*b*:4',3'-*e*]pyridin-4-yl)phenoxy)methyl)pyridine) (**18**). A mixture of bis-aldehyde **4** (0.35 g, 1 mmol), dimedone (**5**) (0.28 g, 2 mmol), 3-methyl-1H-pyrazol-5(4H)-one (**17**) (0.20 g, 2 mmol) was heated in absolute ethanol (30 mL) containing ammonium acetate (0.15 g, 2 mmol) for 6 h. The collected precipitate was isolated and washed with hot EtOH.

Orange crystals (0.61 g, 92%); Mp 300 < °C; IR (KBr): $\bar{\nu}$ 3173, 3129 (NH) cm^{-1} ; ^1H NMR (500 MHz, DMSO- d_6) δ 2.05 (s, 12H, 4CH₃), 4.74 (s, 2H, 2CH), 5.18 (s, 4H, 2OCH₂), 6.85 (d, J = 8.8 Hz, 4H, Ar-H), 7.01 (d, J = 8.5 Hz, 4H, Ar-H), 7.40 (d, J = 7.7 Hz, 2H, pyrdine-H), 7.81 (t, J = 7.8 Hz, 1H, pyrdine-H), 10.84 (br, 6H, 6NH).ppm; MS (EI, 70 eV): m/z (%) 665 [M^+]; anal. calcd for $\text{C}_{37}\text{H}_{35}\text{N}_{11}\text{O}_2$: C, 66.75; H, 5.30; N, 23.14. Found: C, 66.70; H, 5.21; N, 23.12%.

2.2 In vitro cytotoxicity assay

In the MTT assay^{47,48} synthetic compounds (**16**, **15**, **8**, **7**, **11**, and **18**) were tested for their toxicity on HepG2 (liver cancer), A549 (lung cancer), MCF7 (breast cancer), and HFB4 (normal skin) cell lines. The cell lines used in this study were purchased from the Production Company of Vaccines, Sera, and Drugs (VAC-SERA), Egypt. Cell adhesion was accomplished by cultivating cells at 5×10^3 per well in 96-well plates for 24 hours. All test substances were administered to cells at varying concentrations (0–100 $\mu\text{g mL}^{-1}$) for 48 hours. Add 100 μL of MTT solution (0.5 mg mL^{-1}) to each well and incubate for 4 hours at 37 °C. After dissolving the formazan crystals in 100 μL of DMSO, measure the absorbance at 570 nm using a microplate reader.



Three experiments on dose–response curves were used to establish the IC₅₀ values, which indicate the concentration required to reduce cell viability by 50%. The selectivity index (SI) was established by calculating the ratio of IC₅₀ values between normal cells (HFB4) and cancer cells (HepG2, A549, or MCF7).

The study on the structure–activity relationship (SAR) was conducted to understand how changes in structure within the synthesized chemical series affect their anticancer properties. The examination considered molecular structures, heteroatom modifications, scaffold flexibility and rigidity, as well as the electronic characteristics of the substances. Crucial elements comprised the existence of fused polycyclic structures, hydrogen-bond donor/acceptor ability, characteristics of heteroatoms (such as nitrogen and sulfur), and the equilibrium between hydrophilic and lipophilic areas.^{49,50} These characteristics were methodically linked to cellular absorption, target-binding capacity, and selectivity. Significant focus was directed toward how scaffold rigidity, the placement of functional groups, and steric effects influence bioactivity and pharmacokinetic properties. This method offered insights into how changes in structure influenced variations in cytotoxic effectiveness and selectivity, providing useful direction for the logical design of enhanced anticancer compounds in upcoming research.

2.3 Molecular docking

In this research, bis-polyhydroquinoline 7 was docked^{23,51–53} to analyze its binding modes and affinities for five cancer-related protein targets: FGFR4, Aurora A kinase, CDK1, EGFR, and VEGFR2. They were selected due to their essential roles in cancer progression, particularly in hepatocellular carcinoma (HepG2), lung carcinoma (A549), and breast cancer (MCF7) cells, where compound 7 exhibited significant cytotoxicity. Hepatocellular carcinoma is associated with FGFR4 and Aurora A, while the proliferation of lung cancer cells and angiogenesis rely on EGFR and VEGFR2. CDK1 was incorporated due to its regulation of the cell cycle in various tumor types.

The Protein Data Bank (PDB) provided the three-dimensional X-ray structures of the selected target. We used the Molecular Operating Environment (MOE 2015.01) program to make the protein.^{54,55} For protein preparation, each protein structure was examined meticulously to identify any missing residues. Next, any portions that weren't needed were removed, like water molecules, ions, and heteroatoms. Hydrogen atoms were introduced, and the protonation states were adjusted according to the body's pH. For ligand preparation, MOE's Builder module was utilized to construct the 2D structure of compound 7, and then it was changed into a 3D conformer. The MMFF94x force field was employed to minimize the structure's energy and optimize its conformation to the lowest-energy state, ensuring the proper shape and protonation before docking.

The MOE Dock module was utilized to execute docking simulations. The coordinates of the co-crystallized ligand were used to determine the location of the binding site. MOE's site finder (dummy atoms) was employed to find the active pocket. The search region (grid sphere) was altered so that it now

includes all the relevant residues that are within roughly 10 Å of the co-crystallized ligand or dummy atom centroid. This made sure that the pocket was fully closed.

The triangle matcher placement method was used for the docking process, with an initial generation of 100 poses ranked by the London dG scoring function. The London dG scoring technique was used to rank these positions, and then the top 50 poses from each simulation were selected in rigid receptor mode for further refinement. After that, the GBVI/WSA dG scoring tool was used to assign a new score to the new poses. This was to find out the free energy of binding.

The best binding pose for each target was picked based on three things: (1) the lowest GBVI/WSA dG score, which shows the best binding affinity; (2) the presence of important non-covalent interactions (hydrogen bonds, hydrophobic contacts, π – π stacking, and salt bridges) with critical active-site residues; and (3) pharmacophoric alignment with the co-crystallized inhibitor or a known active ligand of the target protein. Finally, a visual check was performed to ensure that there were no steric conflicts and that compound 7 was correctly positioned in the active site and was chemically stable.

2.4 Molecular dynamics simulations

The highest consensus docking scores from compound 7 with the protein receptor Epidermal Growth Factor Receptor tyrosine kinase domain (EGFR TKD)⁵⁶ were subjected to all-atom molecular dynamics (MD) simulations to investigate the relative stabilities of the protein–ligand interactions and to screen the compound for subsequent binding energy calculations. GROMACS V2023 package^{57–59} and the CHARMM36m^{60,61} force field were utilized for all simulations. The parameters and topological files for the compound we chose were generated using the latest CHARMM/CGenFF force field using CHARMM-GUI.^{62–65} The protein–ligand complex was placed in the middle of a box of solvated water molecules with a TIP3P explicit solvation model. To mimic the physiological salt concentrations, 0.154 M ions (98 Na⁺ and 99 Cl[−]) were added to provide charge neutralization and electrostatic screening, which extended the edge distance 20 Å from the protein. The rectangular cubic system for CHARMM and the periodic boundary conditions were set to 104.0 Å in the *x*, *y*, and *z* axis.

The MD procedures included minimization, equilibration (in two steps), and production. The 200 ns MD production simulations did not limit any atoms. The preliminary energy of the solvated protein–ligand complex was minimized through 1 000 000 steps at a constant temperature of 300 K. This was followed by equilibration in two steps: the first was a canonical (NVT) ensemble with 4 000 000 steps, and the second was an isothermal–isobaric (NPT) ensemble with 4 000 000 steps and a 1 fs time integration step. Finally, the production MD simulation step was performed for the isothermal–isobaric (NPT) ensemble with 100 000 000 runs and a 2-fs time integration step for 200 ns. Using NVT and NPT ensembles, the pressure was set to 1 atm. Using the Langevin thermostat, the temperature was regulated to 300.0 K.⁶⁶ Force-field settings that didn't include scaling of 1.2 nm were used to minimize



and balance the complexes in the water box. All atoms, even those of hydrogen, were shown clearly. Trajectory data every 0.05 ns has been obtained. Tools from the GROMACS and VMD packages⁶⁷ were used to look at the full system residues, utilizing the produced trajectories we got during the manufacturing stage. A distance cut-off of 1.0 nm was used for short-range nonbonded interactions with a pair list distance of 1.2 nm, while Lennard-Jones interactions were smoothly cut off at 1.2 nm (rvdm, rcoulomb = 1.2 nm). The particle-mesh Ewald (PME) approach^{68,69} was used to handle long-range electrostatic interactions. For the sake of consistency, the same method for all the MD simulations was utilized.

The binding free energy of the simulated complex was calculated using MD simulations by GROMACS, utilizing snapshots from the system's trajectories over 200 ns, encompassing 4000 trajectories. This was followed by the gmx_MMPBSA tool, based on the AMBER tools MMPBSA.py module with GROMACS files,⁷⁰ where the ligand (L) binds to the protein receptor (R) to form the complex (RL). We are solely interested in computations of relative binding energy, which is provided by Gibbs relative binding energy:

$$\Delta G_{\text{bind}} = \Delta G_{\text{bind,vaccum}} + \Delta G_{\text{RL,solvation}} - (\Delta G_{\text{R}} - \Delta G_{\text{L}})$$

2.5 EGFR inhibition assay

Following the manufacturer's instructions, the EGFR kinase inhibitor screening assay kit (Cat. no. 40321, BPS Bioscience, USA) was used to test how well compound 7 worked against EGFR. This colorimetric test measures the phosphorylation of a poly(Glu-Tyr) substrate by EGFR. An HRP-conjugated anti-phosphotyrosine antibody and TMB substrate were used to find it. The concentrations of compound 7 ranged from 7.81 $\mu\text{g mL}^{-1}$ to 1000 $\mu\text{g mL}^{-1}$, and sorafenib was used as a positive control. After 30 minutes of incubation at 30 °C with EGFR and ATP, the detection reagents were added sequentially. A microplate reader was utilized to assess absorbance at 450 nm. Percent inhibition was figured out, and IC_{50} values were evaluated for compound 7 and sorafenib.

2.6 ADMET prediction

The pkCSM pharmacokinetics web interface⁷¹ evaluates absorption, distribution, metabolism, excretion, and toxicity factors by utilizing graph-based signatures to forecast compound 7's ADMET characteristics. The outcomes were automatically calculated following the input of compound 7's representation into the pkCSM web interface. Furthermore, we utilized *in silico* permeability data from the ADMETlab 3.0 platform,⁷² including PAMPA projections, to get a better idea of how well compound 7 may be absorbed.⁷³ To aid research on compound 7's anticancer capabilities, we evaluated its drug-likeness and forecasted its pharmacokinetic and safety profiles *in silico* to enhance experimental findings and bolster its prospects as an anticancer candidate.

2.7 Statistical analysis

Each experiment was conducted in triplicate ($n = 3$), and results were displayed as mean \pm standard deviation (SD). We used GraphPad Prism 9 software for statistical analysis and fitted dose-response curves to get IC_{50} .

3 Results and discussion

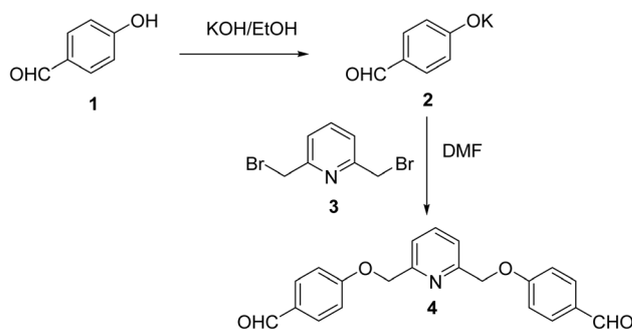
3.1 Chemistry

To reach our objective, 2,6-bis(bromomethyl)pyridine 3 was chosen as an essential intermediate and can serve as a primary element for the creation of 4,4'-((pyridine-2,6-diylbis(methylene))bis(oxy))dibenzaldehyde 4. The interaction of compound 3 with the potassium salt 2 (obtained from the reaction of 4-hydroxybenzaldehyde 1 with ethanolic KOH) in boiling DMF led to the formation of compound 4 in 88% yield (Scheme 1).^{74,75}

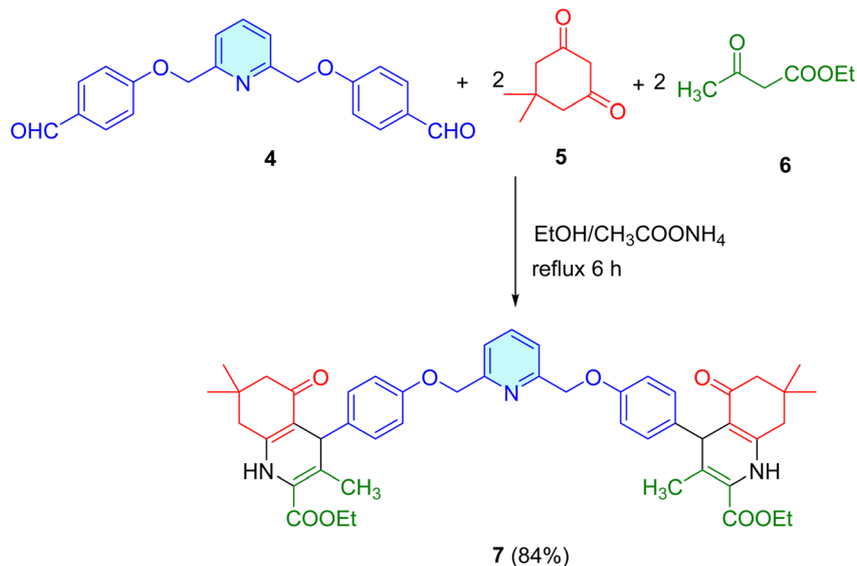
Our primary emphasis was on synthesizing bis-polyhydroquinoline 7 through Hantzsch's four-component condensation reactions that include bis-aldehyde 4, dimedone 5, ethyl acetoacetate 6, and ammonium acetate (Scheme 2). The product was obtained with a yield of 84% after being heated in ethanol at reflux for 6 hours.

Spectral information was employed to establish the structure of compound 7. Consequently, the NH group was recognized at $\bar{\nu}$ 3279 cm^{-1} according to its IR spectra. Additionally, it showed the existence of the carbonyl group at $\bar{\nu}$ 1697 and 1643 cm^{-1} . The ^1H NMR spectrum of compound 7 exhibited characteristic singlets at δ 4.77 for H-4, δ 5.05 for OCH_2 protons, and δ 9.03 for 2NH. It also showed singlets at 0.82, 0.97, and 2.25 ppm, each integrated to 6H assigned to the different methyl groups, as well as two multiplets for two CH_2 groups at δ 1.92–2.16 and 2.27–2.41 ppm. The ethyl protons appeared as a triplet and a quartet at δ 1.09 and 3.94 ppm. Signals for aromatic protons appeared at the expected positions. Moreover, the ^{13}C NMR examination of compound 7 validated its proposed structure, showing characteristic peaks at δ 167.5 and 194.9 ppm for ester and ketonic $\text{C}=\text{O}$, respectively.

The reaction of bis(aldehyde) 4 with dimedone (5) and ammonium acetate in AcOH under reflux produced a 73% yield of bis(3,3,6,6-tetramethyl-3,4,6,7,9,10-hexahydroacridine-



Scheme 1 Synthesis of 4,4'-((pyridine-2,6-diylbis(methylene))bis(oxy))dibenzaldehyde 4.



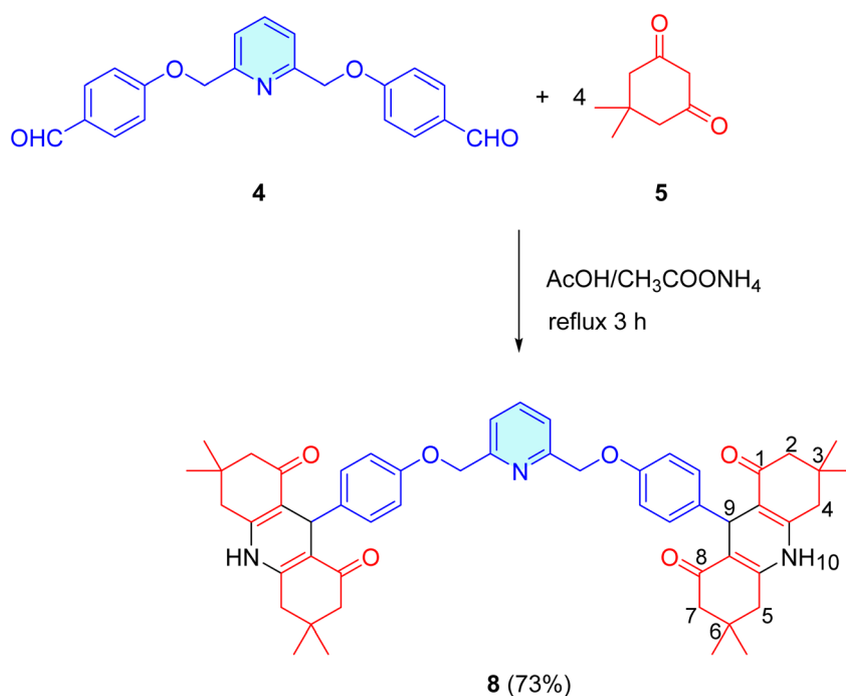
Scheme 2 Synthesis of bis-polyhydroquinoline 7.

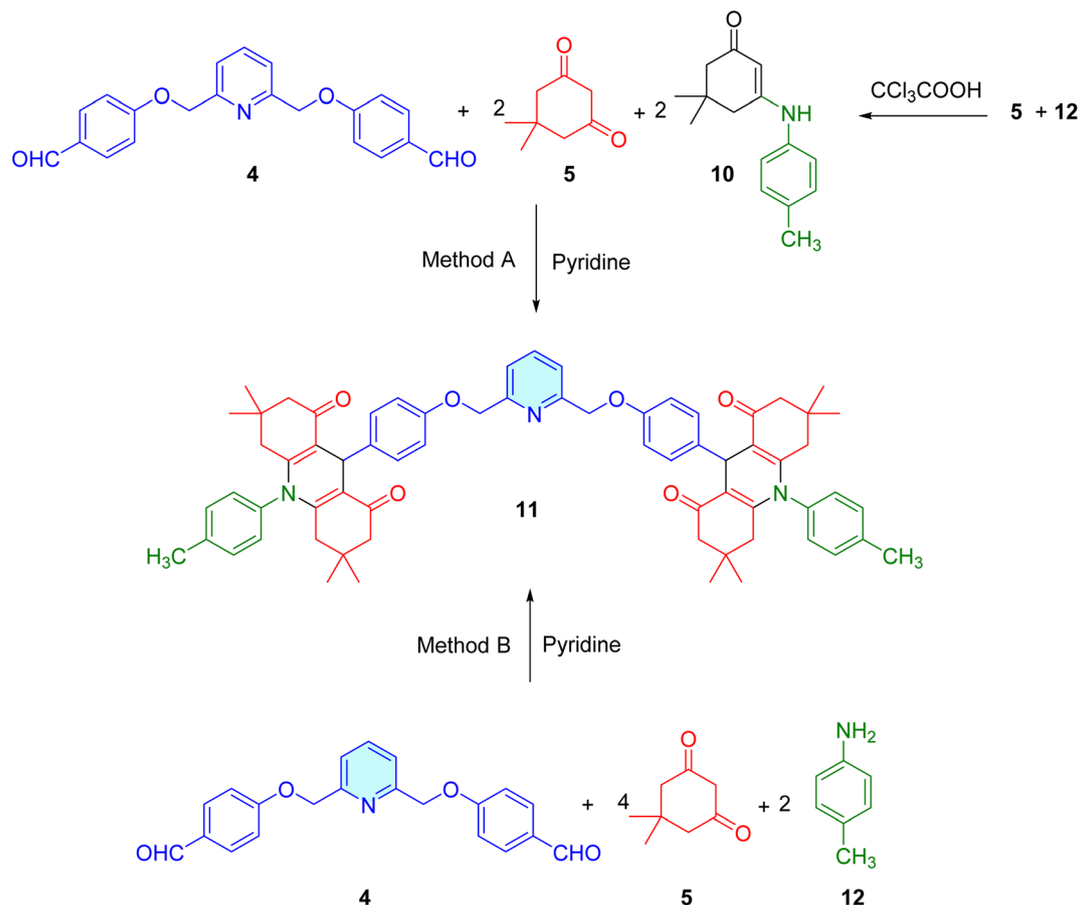
1,8(*2H,5H*)-dione **8** (Scheme 3). The structural examination of the target compound was conducted utilizing spectral data.

The composition of product **8** was established through multiple spectral methods. The IR spectra of compound **15** showed NH groups at $\bar{\nu}$ 3289 cm^{-1} , and the carbonyl groups were observed at $\bar{\nu}$ 1636 cm^{-1} . The ^1H NMR spectrum exhibited two singlets, totaling 24 protons at δ 0.84 and 0.97 ppm, which correspond to eight CH_3 groups. The 16 protons constituting the four methylene groups (H2, H4, H5, H7) were detected as a pair of multiplets at δ 1.94–2.15 and 2.26–2.44 ppm.

Furthermore, the singlet signals at δ 4.73 and 5.04 ppm correspond to the acridine-H9 and $-\text{OCH}_2$, respectively. The NH group appeared as a singlet at δ 9.23 ppm. All other signals were as expected. The proton at the H-5 position resonates at δ = 4.65 ppm, which is a characteristic signal of the chromene moiety present in this class of compounds.⁷⁶

Since acridines hold biological significance, we expanded our study to include the synthesis of bis(9-aryl-substituted acridine). Consequently, when bis-aldehyde **4** is reacted with two equivalents each of dimerone **5** and enaminone **10** under

Scheme 3 Synthesis of bis(hexahydroacridine-1,8-dione) **8**.



Scheme 4 Synthesis of bis(9-aryl-substituted acridine) 11.

reflux conditions in pyridine, it leads to the generation of bis(3,3,6,6-tetramethyl-10-(*p*-tolyl)-3,4,6,7,9,10-hexahydroacridine-1,8(2*H*,5*H*)-dione) **11** in 74% yield (method A) (Scheme 4). The product was obtained with a limited quantity of the target substance when aldehyde **4** reacted with four moles of dimesone **5** and two moles of *p*-toluidine **12** (method B), as illustrated in Scheme 4. It should be noted that the formation of enamine **10** involved the interaction of dimesone **5** with arylamines **12** in the presence of trichloroacetic acid (Scheme 4).⁷⁷

The spectral data helped to identify the structure of compound **11**. Consequently, the IR spectrum revealed a carbonyl group at $\bar{\nu}$ 1636 (C=O) cm^{-1} . The existence of 24 protons was validated in the ^1H NMR spectrum of compound **11**, displaying two singlets at δ 0.68 and 0.84 ppm that were attributed to eight CH_3 groups. Additionally, the singlet peak at δ 4.96 ppm suggests the presence of acridine-H9. All other signals were found exactly where they should have been.

A reaction involving three components, namely dibenzaldehyde **4**, two equivalents of dimesone **5**, and either 6,6-aminopyrimidine-2,4-dione **13** or 6-amino-2-thioxo-2,3-dihydropyrimidin-4(1*H*)-one **14**, produced bis(2,3,5,8,9,10-hexahydropyrimido[4,5-*b*]quinoline-4,6(1*H*,7*H*)diones) **15** and **16** with yields of 91% and 77%, respectively (Scheme 5).

The IR spectra of Compound **15** showed NH groups at $\bar{\nu}$ 3408, 3269, and 3192 cm^{-1} . Furthermore, the carbonyl groups were

observed at $\bar{\nu}$ 1717, 1665, and 1655 cm^{-1} . The ^1H NMR spectrum of compound **15** showed a singlet peak at 4.67 ppm, which was attributed to H5. The NH group displayed wide signals at δ 8.81, 10.26, and 10.65 ppm. Every other signal was in the proper location. The structure of compound **15** was further supported based on 2D-HMBC NMR spectroscopy that indicated 3J -cross peak correlation between H5 at δ = 4.65 and CO at δ = 195.82 (Fig. S18).

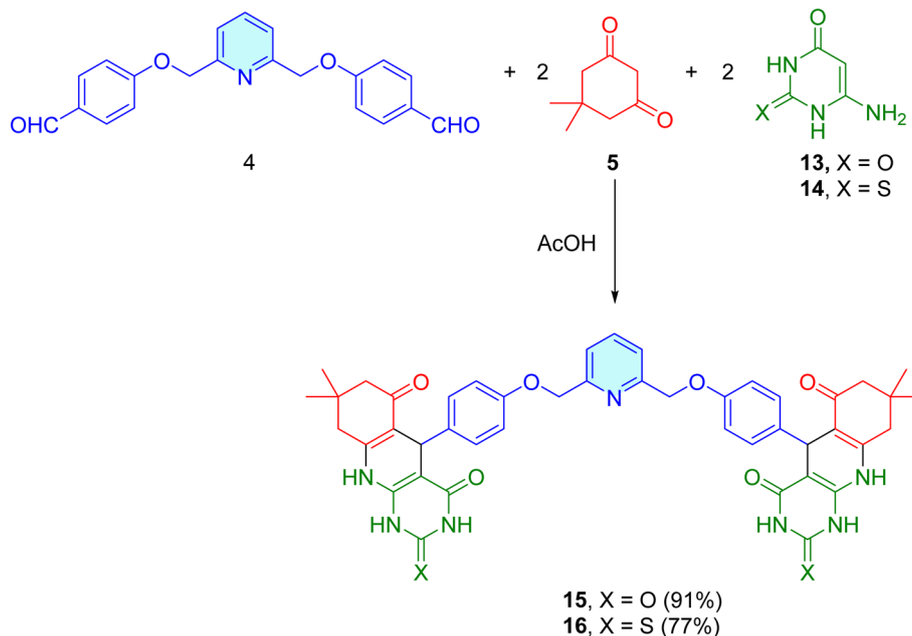
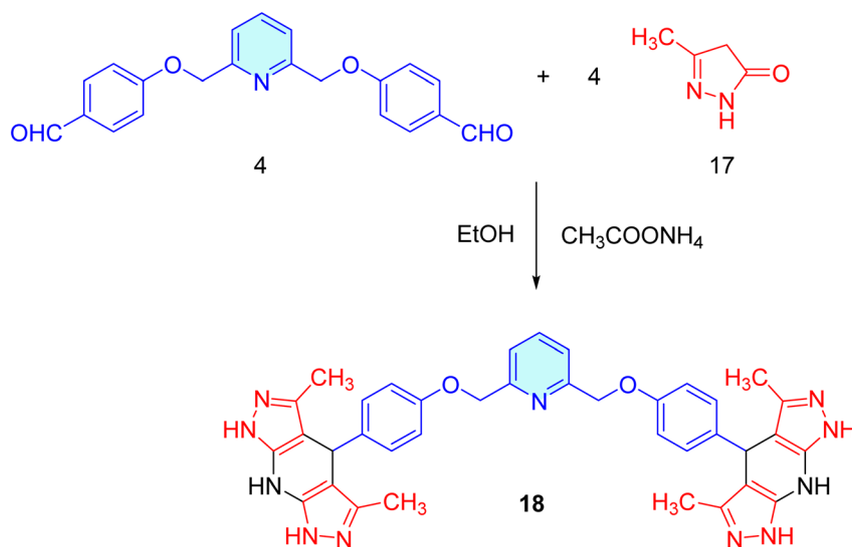
The study was expanded to include the synthesis of bis(3,5-dimethyl-1,4,7,8-tetrahydropyrazolo[3,4-*b*:4',3'-*e*]pyridin-4-yl) alkanes **18** *via* a multicomponent reaction that utilized 3-methyl-1*H*-pyrazol-5(4*H*)-one (**17**), bis(aldehyde) **4**, and ammonium acetate in heated ethanol (Scheme 6).

The IR spectrum for compound **18** displayed the NH groups at $\bar{\nu}$ 3173 and 3129 cm^{-1} . The singlet peak in the ^1H NMR spectrum of compound **18** exhibited an integration of 12 protons at δ 2.05, which corresponds to four CH_3 groups. Moreover, the signals at δ 4.74 and 5.18 ppm correspond to OCH_2 and H4, respectively. The NH groups appeared as a broad singlet peak at δ 10.68. All the other signals were seen in their expected positions.

3.2 *In vitro* cytotoxic activity evaluation

HepG2 (liver cancer), A549 (pulmonary cancer), MCF7 (breast cancer) and normal human skin fibroblast (HFB4) cells were



Scheme 5 Synthesis of bis(hexahydropyrimido[4,5-*b*]quinolinediones) 15 and 16.Scheme 6 Synthesis of bis(tetrahydrodipyrzolo[3,4-*b*:4',3'-*e*]pyridin-4-yl)alkanes 18.

assessed for *in vitro* cytotoxic effects of synthetic compounds 16, 15, 8, 7, 11, and 18 through the MTT assay after 48 hours of exposure, as shown in Fig. 2. Tables 1 and S1 present pertinent IC₅₀ values and selectivity indices (SI). Certain cell lines (HepG2, A549, and MCF7) were selected for both strategic and scientific purposes. The commonly utilized model of hepatocellular carcinoma, HepG2, represents a substantial worldwide cancer load characterized by treatment resistance and rapid proliferation.⁷⁸ A549 cells, derived from human alveolar epithelial carcinoma, serve as a recognized model for non-small cell lung cancer (NSCLC), which is one of the leading causes of cancer-related deaths globally.⁷⁹ MCF7 cells serve as models for

hormone-sensitive breast cancer and are widely used to test anticancer compounds that affect the growth of tumor tissue in response to the hormone estrogen.⁸⁰ These three cell lines offer a solid foundation for assessing the anticancer effects of new compounds sourced from various tissue origins and types of cancer. The evaluated compounds showed a dose-dependent decrease in cancer cell viability according to MTT assay results. Drawing from the bis-polyhydroquinoline framework, 7 exhibited the most significant cytotoxic impact among them. Against HepG2 cells, it exhibited IC₅₀ values of 18.07 ± 1.17 μg mL⁻¹, while against A549 cells and MCF7 cells, it was 14.45 ± 3.73 μg mL⁻¹ and 30.89 ± 6.63, respectively. Notably,



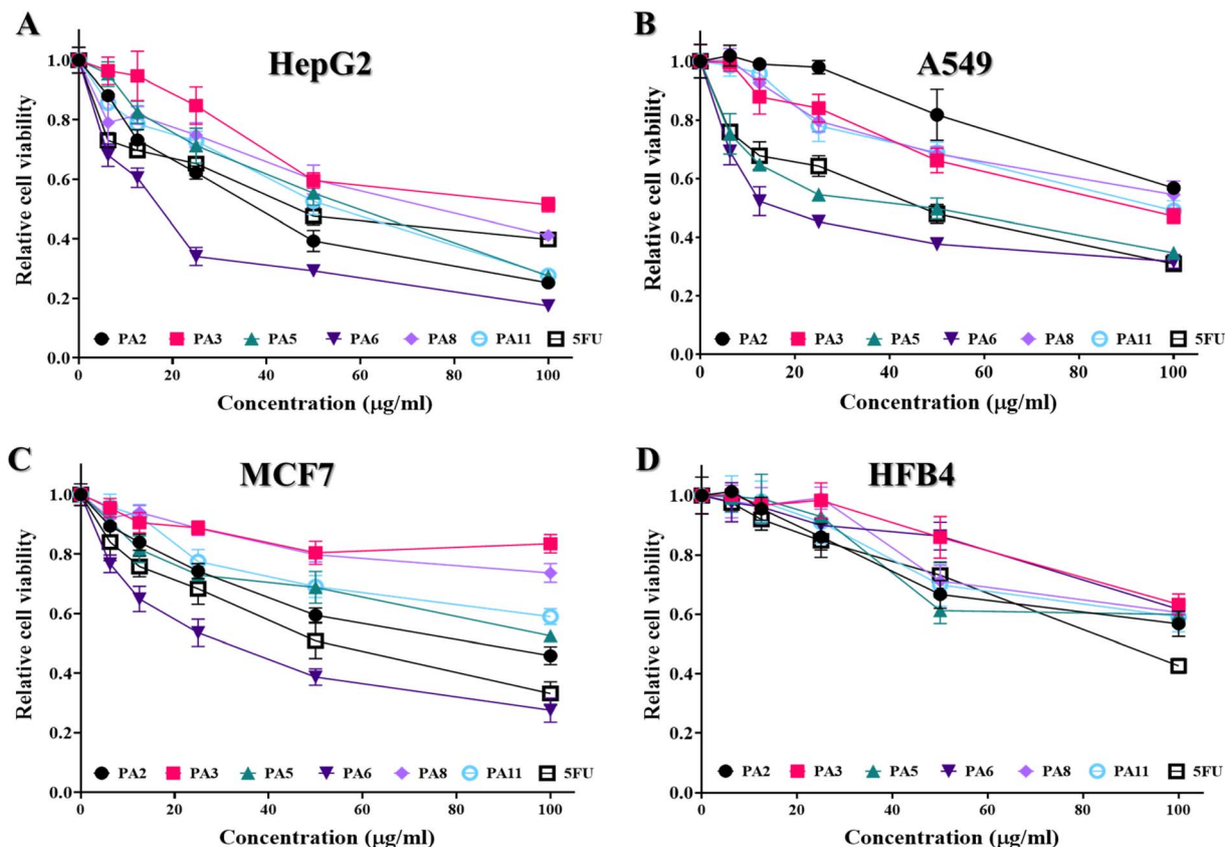


Fig. 2 *In vitro* cytotoxic activities of the novel derivatives (7, 8, 11, 15, 16, and 18) against HepG2 cells (A), A549 cells (B), MCF7 cells (C), and HFB4 cells (D) using MTT assay following 48 h of treatment. Compound PA6 (7); compound PA5 (8); compound PA8 (11); compound PA3 (15); compound PA2 (16); compound PA11 (18).

Table 1 IC_{50} values ($\mu\text{g mL}^{-1}$) of novel derivatives (7, 8, 11, 15, 16, and 18) toward different human cells, HepG2, A549, MCF7, and HFB4 cells, using MTT assay following 48 h of treatment

Compound	IC_{50}^a ($\mu\text{g mL}^{-1}$)				SI^b		
	HepG2	A549	MCF7	HFB4	HepG2	A549	MCF7
16 (PA2)	39.43 ± 3.25	>100	80.70 ± 10.50	>100	>2.53	>1	>1.24
15 (PA3)	65.78 ± 1.87	82.54 ± 6.80	>100	>100	>1.52	>1.21	>1
8 (PA5)	59.14 ± 8.53	44.38 ± 17.84	>100	68.24 ± 4.73	1.15	>1.53	0.68
7 (PA6)	18.07 ± 1.17	14.45 ± 3.73	30.89 ± 6.63	>100	>5.53	>6.92	>3.23
11 (PA8)	73.81 ± 14.90	>100	>100	>100	>1.35	>1	>1
18 (PA11)	55.82 ± 7.43	95.87 ± 3.61	>100	>100	>1.79	>1.04	>1
5-Fluorouracil	45.45 ± 5.97	48.00 ± 4.32	51.04 ± 11.48	88.33 ± 2.53	1.94	1.84	1.73

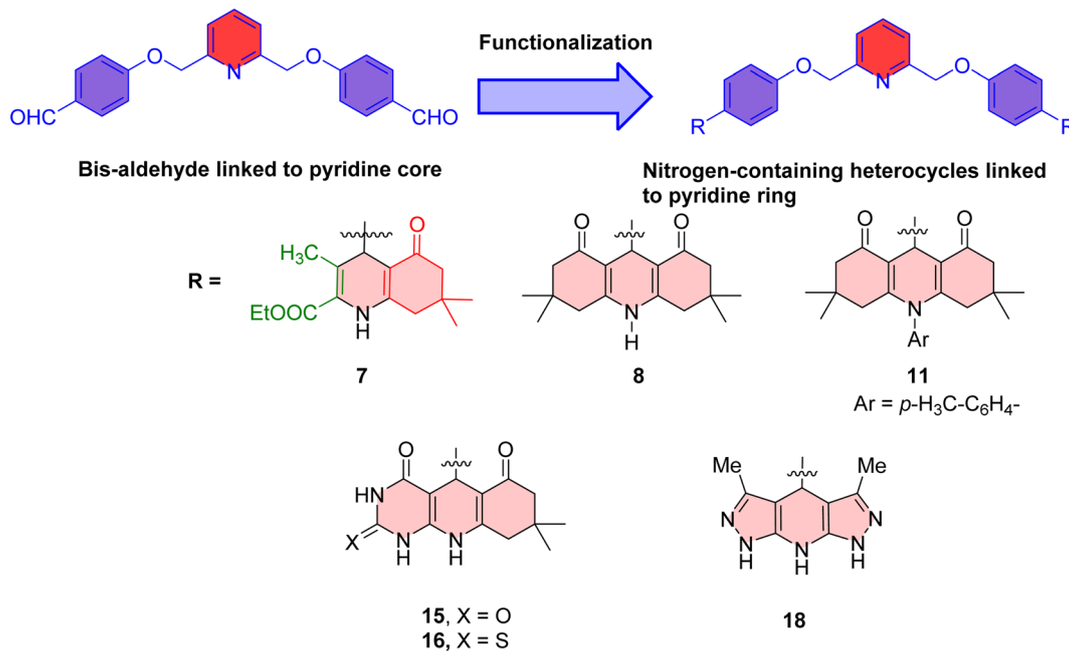
^a The data were expressed as the mean \pm SD of three independent experiments. ^b $SI = IC_{50}$ on HFB4/ IC_{50} on cancer cells.

compound 7 demonstrated pronounced activity against MCF7 cells comparable to its effects on HepG2 and A549 cells, with exceptional selectivity indices (>5.53 for HepG2, >6.92 for A549, and >3.23 for MCF7), and IC_{50} values exceeding $100 \mu\text{g mL}^{-1}$ ensuring minimal toxicity to normal fibroblast cells (HFB4). This notable selectivity implies that 7 specifically focuses on cancerous cells rather than healthy ones, a highly sought-after trait in anticancer drugs.

In comparison, 16, a sulfur-including bis(hexahydropyrimido[4,5-*b*]quinolinedione), showed minimal

effectiveness on A549 cells ($IC_{50} > 100 \mu\text{g mL}^{-1}$), but exhibited moderate cytotoxicity against HepG2 and MCF7 cells ($IC_{50} = 39.43 \pm 3.25 \mu\text{g mL}^{-1}$ and $80.70 \pm 10.50 \mu\text{g mL}^{-1}$, respectively). The selectivity index of 16 against HepG2 and MCF7 (>2.53 and 1.24, respectively) highlighted its safer profile in contrast to nonselective cytotoxic drugs. Compound 15, the nitrogen counterpart of 16, demonstrated a similarly diminished effectiveness relative to 16, presenting IC_{50} values of $65.78 \pm 1.87 \mu\text{g mL}^{-1}$ for HepG2 and $82.54 \pm 6.80 \mu\text{g mL}^{-1}$ for A549, and lacked cytotoxic activity against MCF7 cells ($IC_{50} > 100 \mu\text{g mL}^{-1}$).





Scheme 7 Design concept of nitrogen-containing heterocycles linked to a pyridine ring.

Although **15**'s selectivity indices (>1.52 and >1.21) are satisfactory, replacing sulfur with nitrogen reduced cytotoxic potency, suggesting that sulfur substitution enhances biological activity.⁸¹ The compound **8**—a bis(hexahydroacridine-1,8(2*H*,5*H*)-dione)—demonstrated a slight cytotoxic effect, with IC₅₀ values of 59.14 ± 8.53 μg mL⁻¹ for HepG2 cells and 44.38 ± 17.84 μg mL⁻¹ for A549 cells, and displayed no effective cytotoxicity against MCF7 cells (IC₅₀ > 100 μg mL⁻¹).⁸² However, the slightly lower selectivity indices (1.15, >1.53, and 0.68, respectively) suggested less cancer specificity when compared to **7**. With an IC₅₀ of 73.81 ± 14.90 μg mL⁻¹ for HepG2 and an IC₅₀ > 100 μg mL⁻¹ for A549 and MCF7, **11**, featuring bis(9-aryl-substituted acridine) components, exhibited reduced overall activity, suggesting that increased arylation and greater planarity may adversely influence cytotoxic efficacy, possibly due to reduced cellular absorption or impaired binding affinity to the targeted biomolecules.⁸³

The results show that compound **7** has considerable potential to serve as a treatment, considering that it was very detrimental to A549, HepG2, and MCF7 cells, with IC₅₀ values much lower than those of other similar compounds. It also had a high selectivity index against normal fibroblasts, meaning it was only harmful to cancer cells.

3.3 Structure–activity relationship (SAR) analysis

Key structural components influencing anticancer efficacy were identified through a detailed analysis of the structure–activity relationship (SAR) of the studied compounds. Upon examining the structures of the synthesized compounds, it is possible to conclude that there is a pyridine core linked to two fused-heterocyclic units *via* phenoxy-methyl linkers. The heterocyclic systems include hexahydroquinoline, hexahydroacridine,

tetrahydropyrimido[4,5-*b*]quinoline, and tetrahydrodipyrazolo[3,4-*b*:4',3'-*e*]pyridine. The design approach is illustrated in Scheme 7. The bis-polyhydroquinoline scaffold emerged as the most appropriate framework, as suggested by **7**. Target binding seems to be enhanced through a fused, semi-rigid tetracyclic design that boosts molecular planarity while maintaining a perfect equilibrium between flexibility and rigidity.⁸⁴ Moreover, the numerous hydrogen bond donors and acceptors present in the polyhydroquinoline framework are likely to facilitate positive interactions with cellular proteins involved in managing oxidative stress, such as kinases or enzymes linked to cancer development.^{85,86} Incorporating sulfur atoms, as seen in **16**, appears to enhance cytotoxic efficacy compared to **15**, a nitrogen analogue. Sulfur atom regulation of electrical properties, enhancement of lipophilicity, and facilitation of membrane permeability, all of which may aid in boosting intracellular accumulation and biological effectiveness.^{81,86} Conversely, the limited activity of **16** in comparison to **7** suggests a synergistic advantage from the polyhydroquinoline core that goes beyond mere heteroatom influences.⁸⁷ According to **8** and **11**, the derivatives of bis(hexahydroacridine) and bis(aryl-acridine) showed reduced efficacy and selectivity. Either excessive planarity or large aryl groups could lead to suboptimal interactions with biological targets, which may account for the steric hindrance caused by both.⁸⁸ Additionally, these structures may possess changed pharmacokinetic characteristics and diminished solubility, features considered mainly significant for cytotoxic efficacy. Moreover, **18** demonstrated moderate activity with a pyrazolo-pyridine structure, indicating that nitrogen-dense fused heterocycles hold potential but might need further refinement to achieve the same potency and selectivity as **7**.^{87,89} The inadequate performance of **18** further



emphasizes the importance of balancing hydrophilic and hydrophobic elements within the molecule for maximal anticancer effect. The connection with 5-FU was especially beneficial. While 5-FU is frequently employed in therapy, it showed lower selectivity compared to 7, highlighting the need for the creation of next-generation cancer treatments that merge strong cytotoxic effects with minimized off-target effects. In addition, the superior selectivity of 7 compared to 5-FU suggests a distinct or more targeted mechanism of action, necessitating further in-depth mechanistic research. Overall, the SAR analysis reveals that the type and placement of substituents on the bis-heterocyclic structure significantly affects the cytotoxic capability of the compounds. The existence of a central pyridine linker, along with electron-donating groups, seems to boost cellular absorption and enhance interactions with intracellular targets, probably due to better molecular planarity and π - π stacking ability. Moreover, the bis-polyhydroquinoline structure might enhance lipophilicity, promoting membrane penetration and accumulation within cells. These structural characteristics work together to explain the measured cytotoxic effects. Among the synthesized derivatives, compound 7 exhibited strong activity and significant selectivity, underscoring its potential as a key framework for the strategic development of more effective anticancer therapies.

3.4 Molecular docking of 7

The results of the MTT assay indicated that 7 was safe for normal HFB4 cells but exhibited significant cytotoxic effects on HepG2 and A549 cancer cells. These findings indicate that 7 could exhibit selective anticancer effects through specific molecular interactions with key oncogenic targets. Through molecular docking⁹⁰ of bis-polyhydroquinoline 7 with cancer-related targets, a detailed understanding of its binding affinities, interaction characteristics, and potential mechanisms explaining its cytotoxic effects on HepG2, A549, and MCF7

cancer cells was achieved, thereby investigating the molecular processes behind these experimental findings. The choice of the type of protein receptors was determined by how important they were in key signaling pathways linked to hepatocellular carcinoma (HepG2), non-small cell lung cancer (A549), and breast cancer (MCF7). FGFR4⁹¹ and VEGFR2⁹² were chosen because they are essential for the growth of hepatocellular carcinoma and the formation of new blood vessels, respectively. EGFR⁹³ and Aurora A kinase⁹⁴ are often present in high amounts in A549 and MCF7 cells, and they are thought to be important for tumor development and cell cycle progression. CDK1⁹⁵ was added because it controls mitosis in all cells, and cancer cells that divide quickly have higher levels, including breast, lung, and liver malignancies. This reasoning guarantees that the docking targets are pathologically and therapeutically relevant to the cancer models used *in vitro*.

The initial validation of docking data was achieved by self-docking of the reference co-crystallized ligand (4-anilinoquinazoline inhibitor, erlotinib) with its respective receptor (EGFR; PDB ID: 1M17). The RMSD value was determined to be under 2 Å, and an impressive superimposition of the re-docked (cyan) and co-crystallized ligand (green) inside the active site was shown, as shown in Fig. 3.

3.4.1 Docking with FGFR4 (PDB ID: 4XCU). Docking studies indicated that 7 binds effectively to FGFR4 (Fig. S1 and Table 2), with a docking score of -8.18 kcal mol⁻¹ and multiple stabilizing contacts in the kinase domains. Particularly, a hydrogen bond donor interaction was observed between the carboxylate group of ASP630, located 3.15 Å away, a crucial residue in the ATP-binding site, and the nitrogen atom (N44) of 7. Additionally, reinforcing the intricate structure within the active site, there are more hydrogen bond acceptor interactions between the oxygen atoms (O71 and O107) of 7 and the side chains of ARG483 and LYS503, respectively (2.88 Å and 3.21 Å). Additionally, a π -cation interaction between the six-membered aromatic ring of 7 and the positively charged NH₂ group of

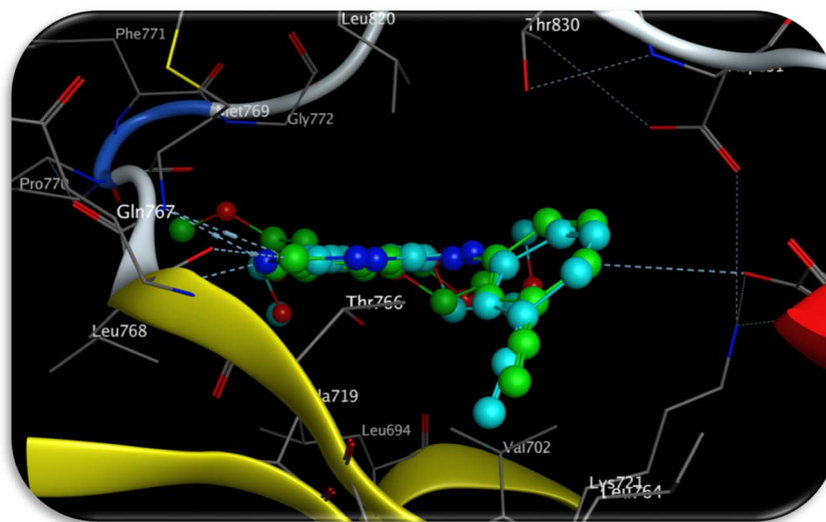


Fig. 3 Superimposition of the re-docked (carbon atoms with cyan color) and the co-crystallized ligand (carbon atoms with green color) within the binding pocket of its protein receptor (EGFR, PDB ID: 1M17).



ARG483 (3.86 Å) strengthened the binding, a type of non-covalent interaction frequently crucial for high-affinity kinase inhibitors. Additional stabilizing interactions were provided by a π -hydrogen bond with ASN557 (4.71 Å).

FGFR4 is known to promote cell proliferation, survival, and metastases in hepatocellular carcinoma (HepG2) by activating the MAPK and PI3K/AKT pathways.⁹⁶ A significant link between 7's strong interaction characteristics within the active area of FGFR4 and the noted reduction in HepG2 cell survival during cytotoxicity assessments indicates that the drug could effectively inhibit FGFR4-mediated signaling. This aligns with reports showing that FGFR4-selective inhibitors reduced HepG2 proliferation in both *in vitro* and *in vivo* circumstances.⁹⁷ The comparable docking score and interaction profile of 7 imply that it may work in the same way as a selective FGFR4 antagonist.

3.4.2 Docking with Aurora A kinase (PDB ID: 3H0Z). 7 exhibited a docking score of -8.04 kcal mol⁻¹ with Aurora A kinase, indicating a strong binding affinity. The ligand formed two key hydrogen bonds (Fig. S2 and Table 2); one was between its oxygen atom (O70) and the backbone nitrogen of THR292 (2.93 Å), and the other was between its oxygen atom (O107) and the NZ atom of LYS143 (2.90 Å). Moreover, a π -hydrogen interaction was observed between the aromatic ring of 7 and the side chain of THR292 (4.16 Å).

The formation of the mitotic spindle, alignment of chromosomes, and maturation of centrosomes are all largely reliant on Aurora A kinase. The overexpression of Aurora A results in genomic instability and cancer.^{98,99} The documented interaction of 7 with key residues in the Aurora active site suggests that 7 could lessen the activity of the kinase, consequently promoting mitotic arrest and cell death, outcomes frequently observed in cancer cells during treatment. This clarifies a mechanism through which 7 induces cytotoxicity and modifies cell cycle progression, particularly during the G2/M phase. Similar binding characteristics and inhibitory capabilities have been shown for established Aurora kinase inhibitors, including MLN8237 (alisertib), that exhibited analogous hydrogen bonding with THR residues in preclinical studies of lung and liver malignancies.^{100,101} This confirms that 7's binding mode is properly configured.

3.4.3 Docking with CDK1 (PDB ID: 6GU6). With a docking score of -7.51 kcal mol⁻¹, 7 demonstrates a strong binding affinity to CDK1 (PDB ID: 6GU6). The ligand formed three hydrogen bonds (Fig. S3 and Table 2): one between an oxygen atom (O70) and the nitrogen backbone of GLU12 (3.23 Å), another between its carbon atom (C34) and ASP86 (3.10 Å), and a third involving another oxygen atom (O97) with the backbone nitrogen of LEU83 (3.11 Å). Additionally, a π -hydrogen interaction was identified between the aromatic ring of 7 and PHE80 (4.27 Å).

CDK1, often elevated in different tumors, plays a crucial role in regulating the G2/M transition during the cell cycle.¹⁰² The identified binding mechanism indicates that 7 attaches directly within the ATP-binding pocket of CDK1, potentially disrupting its kinase function. Even though the docking score was slightly less than that of FGFR4 and EGFR, the nature and abundance of

the interactions suggest that 7 could effectively lower CDK1 activity, thereby aiding in the observed inhibition of proliferation and triggering of cell cycle arrest in both HepG2 and A549 cells. Interestingly, the interaction pattern is quite like that described for roscovitine (Seliciclib), which is a well-known CDK1 inhibitor. This suggests that 7 may have a similar inhibitory profile.¹⁰³

3.4.4 Docking with EGFR (PDB ID: 1M17). Compound 7 exhibited the highest binding affinity among the examined targets with a docking score of -8.98 kcal mol⁻¹, making it the most advantageous protein target assessed. Deep inside the ATP-binding pocket (Fig. 4 and Table 2), a singular yet significant hydrogen bond formed between the oxygen atom (O97) of 7 and the amine group of LYS721's side chain (2.84 Å).

Particularly in non-small cell lung cancer (NSCLC), EGFR plays a vital role in cancer progression by enhancing cell survival, proliferation, and angiogenesis through the activation of downstream pathways such as RAS/RAF/MEK/ERK and PI3K/AKT.^{104,105} Considering 7's strong binding affinity to EGFR, it appears that it could serve as an effective EGFR inhibitor, thus inhibiting these proliferative signals. For A549 cells, which primarily depend on EGFR signaling, this holds significant importance. Consequently, the strong molecular rationale for the significant antiproliferative effect of 7, shown experimentally against A549 lung cancer cells, is derived from the docking data. The docking pattern resembles that of erlotinib, which is an EGFR inhibitor that has been authorized for use in humans and likewise engages with LYS721 and fills the ATP-binding cleft in a comparable manner.¹⁰⁶ This enhances the molecular understanding of 7's antiproliferative activity *via* EGFR suppression.

3.4.5 Docking with VEGFR2 (PDB ID: 3VHE). With a docking score of -7.60 kcal mol⁻¹, 7 demonstrated outstanding binding to VEGFR2 (PDB ID: 3VHE). In addition to a hydrogen bond between the oxygen atom (O70) of 7 and ASN923 (3.20 Å), several stabilizing interactions were noted, including hydrogen bonds involving CYS1045 (4.28 Å) and ASP1046 (3.26 Å). Additionally, π -hydrogen interactions with GLU886 and VAL899 further contributed to stabilizing the ligand at the active site (Fig. S4 and Table 2).

Tumor angiogenesis, which involves the formation of new blood vessels that provide nutrients and oxygen to growing tumors, relies on VEGFR2. Blocking VEGFR2 leads to restricted growth and dissemination capability by starving tumors of essential nutrients, along with decreased angiogenesis.^{107,108} The identified interaction between 7 and VEGFR2 suggests that it could possess antiangiogenic characteristics, thereby aiding not just in the decrease of primary tumor growth but also in the avoidance of metastasis, which is vital for aggressive cancers like hepatocellular carcinoma and lung adenocarcinoma. This aligns with interaction investigations of established VEGFR2 medications like sorafenib, which additionally associates with ASN923 and CYS1045, supporting 7's expected antiangiogenic effectiveness.¹⁰⁹

3.5 Correlation with experimental cytotoxicity

Through hydrogen bonding, π -cation interactions, and hydrophobic contacts, the comprehensive docking analysis indicates



Table 2 Docking interactions and binding energies of compound 7 with FGFR4 (4XCU), Aurora A (3H0Z), CDK1 (6GU6), EGFR (1M17), and VEGFR2 (3VHE)

Protein	PDB ID	Ligand	Receptor	Interaction	Distance	E (kcal mol ⁻¹)	S (energy score)
FGFR4	4XCU	N 44	OD2 ASP 630	H-donor	3.15	-4.7	-8.18
		O 71	NH2 ARG 483	H-acceptor	2.88	-2.1	
		O 107	NZ LYS 503	H-acceptor	3.21	-2.5	
		6-Ring	NH2 ARG 483	Pi-cation	3.86	-0.5	
		6-Ring	ND2 ASN 557	Pi-H	4.71	-0.8	
Aurora A	3H0Z	O 70	N THR 292	H-acceptor	2.93	-13.6	-8.04
		O 107	NZ LYS 143	H-acceptor	2.90	-7.2	
		6-Ring	CG2 THR 292	Pi-H	4.16	-0.5	
		C 34	OD2 ASP 86	H-donor	3.10	-0.6	
CDK1	6GU6	O 70	N GLU 12	H-acceptor	3.23	-2.2	-7.51
		O 97	N LEU 83	H-acceptor	3.11	-1.6	
		C 99	6-Ring PHE 80	H-pi	4.27	-0.6	
		O 97	NZ LYS 721	H-acceptor	2.84	-12.1	
VEGFR2	3VHE	N 4	SG CYS 1045	H-donor	4.28	-0.7	-7.60
		C 13	O ASP 1046	H-donor	3.26	-0.5	
		O 70	N ASN 923	H-acceptor	3.20	-1.0	
		6-Ring	CG GLU 885	Pi-H	3.73	-0.6	
		6-Ring	CG2 VAL 899	Pi-H	4.19	-0.6	

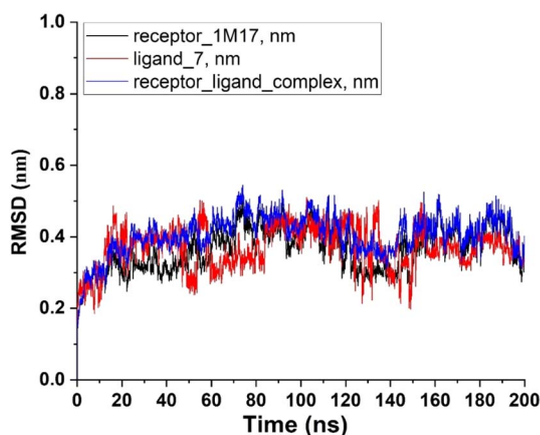


Fig. 5 The root means square deviation (RMSD) of solvated ligand (7), receptor, and protein ligand complex (7-1M17) during 200 ns MD simulation time.

affected them. The average RMSF and the standard deviation of the average RMSF values were 1.787 and 1.574 Å, respectively. Fig. S6 shows how the residues in both systems, protein–ligand complexes, change over time. The terminal residues exhibited greater variations compared to the core residues, which is anticipated due to the flexible characteristics of the bimolecular terminals. The system's average RMSF is less than 5 Å, which means that the complex is quite stable and the ligand molecule has a strong affinity for the receptors.

We used an angle cut = 20° and rcut = 3.0 Å to figure out how many hydrogen bonds formed between the solvated protein and ligands. We then charted the results in contrast to time (200 ns) in Fig. S7a and b. For the solvated protein–ligand complex, the average number of hydrogen bonds per timeframe was 3.1, and for the protein–ligand complex systems, it was 0.26. The total

study revealed that ligand–protein interaction dramatically altered the amount of hydrogen bonds, increasing from 1 to 9 HBs per trajectory analysis in the instance of the solvated protein–ligand complex, and from 1 to 2 in the case of protein–ligand complex systems. These findings show that the system becomes relatively stable for interactions between hydrogen bonds.

The radius of gyration (R_g) shows how tightly packed the protein structure is and how stable the simulation is. For all frames of protein EGFR TKD, compound 7, and ligand–protein complex systems as displayed in Fig. S8, the average R_g values were 2.054, 0.776, and 2.049 nm, respectively. The standard deviation of the average R_g values was 0.208, 0.675, and 0.197 Å, respectively. This is clear because of how ligands and bimolecular receptor complexes work. The overall change in the average R_g of the protein–ligand complex shows that it reaches equilibrium quickly, which means that the binding mode and interactions in the system are stable.

The solvent-accessible surface area (SASA) of the protein was computed during molecular dynamics simulation under ligand-bound circumstances, attributable to ligand binding to the protein. The average SASA values for all frames of protein EGFR TKD, compound 7, and ligand–protein complex systems shown in Fig. S9 were 162.541, 12.889, and 163.004 nm², respectively. The standard deviations of the average SASA values were 3.446, 0.553, and 3.729 nm², respectively. The study shows the folding configuration of the protein and how stable it is when it binds to a ligand, based on the nature of the complex and the outcomes that show a little change in the protein and its complexes. TKD complex equilibrium rapidly stabilizes, exhibiting oscillations within a restricted range, accompanied by minimal conformational alterations and a stable SASA, which correlates with enhanced hydrophobic interactions.



Table 3 Summary of the binding free energy MM/PBSA and MM/GBSA calculated for the solvated protein ligand complex (7-1M17) during a 200 ns MD simulation time. All values are given in (kcal mol⁻¹)

Protein–ligand complex ^a	GBSA							PBSA			
	ΔE_{VDW}	ΔE_{EEL}	ΔG_{GAS}	ΔE_{GB}	ΔE_{SURF}	ΔG_{SOLV}	$\Delta TOTAL$	ΔE_{PB}	ΔE_{NPOLAR}	ΔG_{SOLV}	$\Delta TOTAL$
7-1M17	-50.11	-21.83	-71.94	47.09	-6.99	40.1	-31.84	50	-6.23	43.77	-28.17

^a ΔE_{VDW} = van der Waals energy; ΔE_{ELE} = electrostatic energy; ΔG_{GAS} = gas phase free energy; ΔG_{SOLV} = solvation free energy; ΔE_{SURF} = Surface energy; ΔE_{GB} = generalized Born energy; ΔE_{PB} = Poisson–Boltzmann energy; ΔE_{NPOLAR} = non-polar energy; ΔG_{bin} = calculated total binding free energy (kcal mol⁻¹). MM-GBSA molecular mechanics generalized Born surface area; MM-PBSA Molecular mechanics Poisson–Boltzmann surface area.

To quantify the binding between the EGFR TKD receptor and the ligand, a contact frequency (CF) study was conducted using the contact Freq.tcl module on VMD, with a cutoff of 4 Å, as presented in Fig. S10. In the simulated receptor–ligand complex, the following amino acid residues displayed elevated CF values ranging from 93.65% to 43.0%: GLY695, PHE699, SER696, VAL702, ARG817, THY766, CYS773, ASN818, LEU694, ALA719, ASP831, LEU820, LYS721, THR830, LYS830, THR830, and VAL693. These values indicate that the complex system demonstrates a strong binding affinity to TKD inhibition complexes.

The molecular mechanics energy methodology (MM/GBSA and PBSA) is a well-known way to find the binding free energies of receptor–ligand complexes. GBSA merges the generalized Born and surface area continuum solvation models, whereas the one-average molecular mechanics Poisson–Boltzmann surface area (MM-PBSA) method^{70,118–120} is path-independent and calculates the free energy of binding by examining ensembles of the initial and final states. Consequently, MM-PBSA demonstrates superior efficiency compared to MMGBSA. Both methods could be more dependable, less costly in terms of calculation, and more efficient than docking scores.^{117,121} Based on molecular dynamics (MD) simulated data, we calculated the binding free energy of the simulated complex to double-check the protein affinity that docking simulation studies had predicted for receptor–ligand complexes. Table 3 shows all the calculated contributions of each energy component in MMGBSA and MMPBSA for both complexes. Strong negative values show advantageous interactions and strong receptor–ligand affinity, whereas positive net binding energy shows low-docked stability. The net binding free energy of the study complex was determined to be -31.84 kcal mol⁻¹ for GBSA and -28.17 kcal mol⁻¹ for PBSA. A detailed analysis of the individual energy contributions indicates that the van der Waals, electrostatic, and free energies for the gas phase are -50.11, -21.83, and -71.94 kcal mol⁻¹, respectively, corresponding to the complex. Notably, the van der Waals and electrostatic energies significantly influenced the binding affinity that facilitated complex formation. Finally, the energy of the surface and non-polar components is very reasonable for complexes, ranging from -6.99 to -6.23 kcal mol⁻¹. The net binding free energy stabilizes the conformation of the receptor–ligand affinity complex, indicating a promising and high affinity between the ligand and receptor. This must be experimentally

evaluated to confirm the biological efficacy of the proposed pharmacological inhibition.

These results provide valuable mechanistic insight into the molecular basis of compound 7's activity. Docking studies indicated favorable initial binding, which was corroborated by MD simulations showing sustained stability and reduced conformational flexibility of the EGFR catalytic domain.^{122,123} The favorable binding free energy, together with its decomposition by MM/PBSA and MM/GBSA, further supports the robustness of this interaction and offers both structural and energetic justification for the experimentally observed anticancer potential of compound 7.¹²⁴ The simulations highlight a significant inhibitory capacity, reinforcing compound 7 as a promising framework for the development of novel EGFR-targeted anticancer agents.^{125,126}

3.7 EGFR inhibition assay of compound 7

Following molecular docking and molecular dynamics analyses, compound 7 was selected for biochemical evaluation due to its strong predicted affinity toward EGFR, a well-established therapeutic target in several malignancies.¹²⁷ *In silico* studies revealed that compound 7 displayed superior docking scores with EGFR compared to the other proteins investigated, suggesting its potential as a novel EGFR inhibitor.

To experimentally validate this prediction, compound 7 was tested using an EGFR kinase inhibition assay at concentrations ranging from 7.81 to 1000 µg mL⁻¹ and compared to sorafenib, a clinically approved multikinase inhibitor.¹²⁸ As shown in Fig. 6, compound 7 inhibited EGFR in a dose-dependent manner, with an IC₅₀ value of 21.45 µg mL⁻¹, whereas sorafenib exhibited a stronger effect with an IC₅₀ of 7.52 µg mL⁻¹ under the same conditions. Although less potent than sorafenib, compound 7 demonstrated clear inhibitory activity, supporting its ability to interact with the EGFR active site in a biologically relevant manner. The observed inhibition is consistent with docking predictions, which indicated that compound 7 engages key residues within the EGFR binding cleft through hydrogen bonding and hydrophobic interactions critical for kinase inhibition. The alignment between computational and experimental findings further supports EGFR as a plausible molecular target of compound 7. Moreover, the antiproliferative effects observed in cellular assays align with the enzymatic inhibition results, suggesting that EGFR suppression contributes to the cytotoxic activity of compound 7.



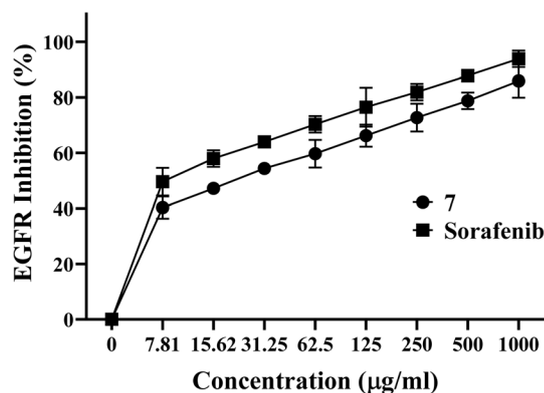


Fig. 6 Dose–response curves of compound 7 and sorafenib against EGFR.

While less potent than sorafenib, compound 7 possesses a distinct scaffold that may confer unique selectivity or pharmacological properties. Structural optimization and kinase panel screening will be important next steps to refine its potency and broaden its therapeutic relevance. Collectively, these results identify compound 7 as a promising EGFR-targeted scaffold warranting further development as a potential anticancer agent.

3.8 ADMET analysis of 7

Initial chemical assessment in current anticancer drug development heavily depends on monitoring ADMET (absorption, distribution, metabolism, excretion, and toxicity). It assists in pinpointing candidates with outstanding pharmacokinetic attributes and acceptable safety thresholds before significant *in vivo* or clinical evaluations.^{129,130} Many of the most promising cytotoxic drugs fail in subsequent phases due to low bioavailability, high toxicity, or metabolic instability.^{129,130} Employing ADMET analysis^{129,131} ensures that potential drug candidates like 7 meet necessary efficacy and safety criteria, while also enhancing lead optimization, reducing late-stage attrition, and accelerating development timelines.

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Compound 7 demonstrated favorable absorption characteristics, supporting its potential oral bioavailability. As shown in Table 4, it exhibited a high predicted intestinal absorption rate (97.87%), consistent with a moderate Caco-2 permeability value (0.489 log Papp in 10⁻⁶ cm s⁻¹), suggesting efficient uptake

across intestinal epithelial cells. The compound was also predicted to be both a substrate and dual inhibitor of *P*-glycoprotein (*P*-gp), indicating the possibility of efflux-mediated transport and transporter-related drug interactions. This dual behavior may influence intracellular drug accumulation and could play a role in overcoming resistance mechanisms commonly associated with *P*-gp overexpression in cancer cells. In addition, the predicted skin permeability (−2.735 log *K*_p) indicated negligible dermal absorption, while the low passive PAMPA permeability suggested that active transporters may facilitate uptake in specific tissues. Collectively, these findings highlight a favorable absorption profile for compound 7, supporting its advancement into further preclinical evaluation.

Distribution assessments indicated a relatively low volume of distribution (0.098 log L kg⁻¹) and significant plasma protein binding (percent unbound of 0.015), suggesting restricted tissue infiltration and a low free drug level in plasma. Minimal values for blood–brain barrier permeability (−0.611 log BB) and CNS penetration (−3.077 log PS) suggested insufficient exposure to the central nervous system. This is good for non-CNS anticancer agents because it lowers the risk of neurotoxicity while increasing the drug's exposure to peripheral tumors.¹³² These traits might lead to a more targeted and safer anticancer profile, especially for solid tumors outside the central nervous system.

Concerning metabolism, compound 7 was identified as a CYP3A4 substrate, indicating liver metabolism through this primary enzymatic pathway.¹³³ Despite the recognized variability among individuals in this enzyme, CYP2D6 was not a substrate. Concerning potential drug–drug interactions that require attention in combination treatment situations, compound 7 showed inhibitory effects on several CYP isoforms, such as CYP3A4, CYP2C19, and CYP2C9.¹³⁴ This inhibitory behavior requires careful consideration when used with other CYP-metabolized drugs, since it might change their plasma levels and how well they work. The molecule, however, did not inhibit CYP1A2 or CYP2D6, which somewhat lowered its overall metabolic liability. This type of targeted inhibition implies the possibility of preventing extensive metabolic changes and may provide safer multiple therapy regimens.

While compound 7 is not a substrate for renal OCT2, it displayed a low total clearance rate (−0.499 log mL min⁻¹), indicating prolonged systemic retention, which could enhance therapeutic exposure through this transporter.¹³⁵ This persistence may lead to an extended dosage interval or prolonged pharmacological action, provided that hazardous thresholds are not surpassed.

From a toxicological perspective, compound 7 exhibited a very strong safety profile. The AMES test showed no mutagenicity, minimal genotoxicity, and no hERG I inhibition,¹³⁶ suggesting a low risk for cardiotoxicity or QT interval prolongation. But its projected suppression of hERG II means that cardiomyocytes or animal models need to be carefully tested for safety using electrophysiology. The highest dosage tolerated in humans was modest (0.075 log mg kg⁻¹ per day); in rats, both acute and chronic toxicity measures (LD₅₀: 3.128 mol kg⁻¹; LOAEL: 1.422 log mg kg⁻¹ per day) remained within acceptable



Table 4 Predicted ADMET properties of 7 using the pkCSM tool. This table presents key pharmacokinetic and toxicity parameters relevant to its potential as an anticancer agent

Property	Model name	Predicted value	Unit
Absorption	Water solubility	-4.662	Numeric (log mol L ⁻¹)
	Caco2 permeability	0.489	Numeric (log Papp in 10 ⁻⁶ cm s ⁻¹)
	Intestinal absorption (human)	97.873	Numeric (% absorbed)
	Skin permeability	-2.735	Numeric (log K _p)
	P-Glycoprotein substrate	Yes	Categorical (yes/no)
	P-Glycoprotein I inhibitor	Yes	Categorical (yes/no)
	P-Glycoprotein II inhibitor	Yes	Categorical (yes/no)
Distribution	VDss (human)	0.098	Numeric (log L kg ⁻¹)
	Fraction unbound (human)	0.015	Numeric (Fu)
	BBB permeability	-0.611	Numeric (log BB)
	CNS permeability	-3.077	Numeric (log PS)
Metabolism	CYP2D6 substrate	No	Categorical (yes/no)
	CYP3A4 substrate	Yes	Categorical (yes/no)
	CYP1A2 inhibitor	No	Categorical (yes/no)
	CYP2C19 inhibitor	Yes	Categorical (yes/no)
	CYP2C9 inhibitor	Yes	Categorical (yes/no)
	CYP2D6 inhibitor	No	Categorical (yes/no)
	CYP3A4 inhibitor	Yes	Categorical (yes/no)
Excretion	Total clearance	-0.499	Numeric (log mL min ⁻¹ kg ⁻¹)
	Renal OCT2 substrate	No	Categorical (yes/no)
Toxicity	AMES toxicity	No	Categorical (yes/no)
	Max. tolerated dose (human)	0.075	Numeric (log mg kg ⁻¹ per day)
	hERG I inhibitor	No	Categorical (yes/no)
	hERG II inhibitor	Yes	Categorical (yes/no)
	Oral rat acute toxicity (LD ₅₀)	3.128	Numeric (mol kg ⁻¹)
	Oral rat chronic toxicity (LOAEL)	1.422	Numeric (log mg kg ⁻¹ _bw per day)
	Hepatotoxicity	Yes	Categorical (yes/no)
	Skin sensitisation	No	Categorical (yes/no)
<i>T. Pyriformis</i> toxicity	0.285	Numeric (log ug L ⁻¹)	
Minnow toxicity	-2.124	Numeric (log mM)	

limits. Hepatotoxicity forecasts notably recommend rigorous liver surveillance during the preliminary *in vivo* studies. Even though hepatotoxicity appeared to require liver attention during development, the substance was not anticipated to cause skin sensitivity.

Ultimately, information regarding environmental toxicity indicated slight to significant effects on aquatic organisms, implying that ecological consequences must be evaluated in subsequent stages. This consideration becomes particularly important when addressing large-scale production and minimizing the environmental impact of manufacturing waste. Overall, compound 7 exhibits a drug-like ADMET profile, characterized by high intestinal absorption, selective metabolic interactions, regulated tissue distribution, and manageable toxicity risks. These integrated pharmacokinetic and toxicological properties support its continued investigation as a promising multi-target anticancer candidate, warranting further validation through experimental pharmacology and *in vivo* studies.

4 Conclusion

The increasing occurrence of cancer, especially in developing regions, highlights the pressing demand for new therapeutic agents that can tackle resistance to existing treatments. In this

study, we created a range of bis-heterocyclic structures connected by a pyridine unit, illustrating a new category of hybrid substances with significant pharmaceutical promise. These derivatives can be produced effectively under gentle conditions using easily accessible starting materials, allowing for the generation of adequate amounts for biological assessment. The synthesized compounds' cytotoxic activities were evaluated on A549 (lung cancer), HepG2 (hepatocellular carcinoma), and MCF7 (breast cancer) cell lines, using human fibroblasts (HSF) as a normal control. Out of the compounds evaluated, compound 7 demonstrated the most potent anticancer effects, significant selectivity for cancerous cells, and low toxicity to normal fibroblasts. Molecular docking analyses indicated that compound 7 could function *via* various oncogenic pathways, such as EGFR and FGFR4, essential for the survival of cancer cells. Toxicological profiling additionally revealed its beneficial safety features, such as non-mutagenicity and a low risk of cardiotoxicity. Moreover, ADMET evaluations validated its pharmaceutical characteristics, demonstrating promising absorption and tolerable safety measures. Combined, these findings emphasize compound 7 as a potential multi-target anticancer candidate. Subsequent research must encompass thorough *in vivo* assessments to analyze its pharmacodynamic effectiveness, bioavailability, metabolic stability, and overall safety. These studies will offer crucial translational insights and



enhance its prospects as a feasible candidate for preclinical development.

Conflicts of interest

There are no conflicts to declare.

Data availability

All data generated or analyzed during this study are included in the article and its supplementary information (SI). Supplementary information: molecular docking interactions of compound 7 with FGFR4, Aurora A, CDK1, and VEGFR2; molecular dynamics simulation analyses (RMSD, RMSF, hydrogen bonding, radius of gyration, SASA, and contact frequency); ^1H and ^{13}C NMR and IR spectra of compounds 7, 8, 11, 15, 16, and 18; and *in vitro* cytotoxicity (IC_{50} values (μM) and selectivity index) data against HepG2, A549, MCF7, and HFB4 cell lines.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d6ra00243a>.

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