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Design, synthesis, molecular docking and in vitro anticancer activities of 1-(4-(benzamido)phenyl)-3-arylurea derivatives†

Prafulla Sabale, **D** **a Nusrat Sayyad, **a Abuzer Ali, **D** Vidya Sabale, **D** Mohammed Kaleem, **D** **d** Turky Omar Asar, **e Amena Ali, **D** Md. Ali Mujtaba **D** and Md. Khalid Anwer **D** h

In both premenopausal and postmenopausal women, oestrogens play a critical role in the development of breast cancer. Aromatase is an enzyme that catalyses the final step in the biosynthesis of estrogen and has emerged as a promising target for therapeutic intervention. This study aimed to design and evaluate novel 1-(4-(benzamido)phenyl)-3-arylurea derivatives as potential aromatase inhibitors. Through molecular docking, promising leads were identified and synthesized. Spectroscopic techniques confirmed their structural integrity. Cytotoxicity against various cancer cell lines was assessed using MTT assay. Docking investigations against the aromatase enzyme (3s7s) elucidated binding interactions and energies. Compound 6g, exhibiting a binding energy of -8.6 kcal mol $^{-1}$ and interacting with ALA306 and THR310 residues, showed the most promising activity. It demonstrated GI₅₀ values ranging from 14.46 μ M, 13.97 μ M, 11.35 μ M, 11.58 μ M, and 15.77 μ M against A-498, NCI-H23, MDAMB-231, MCF-7, and A-549 respectively. Lastly, the physicochemical, and ADMET properties of the compound were predicted. These findings highlight the potential of 1-(4-(benzamido)phenyl)-3-arylureas as a new class of antitumor agents targeting aromatase. Their versatility and superior activity compared to standard chemotherapeutic agents, like doxorubicin, warrant further investigation for the development of broader-spectrum anticancer drugs.

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Introduction

Cancer is a complicated disease with some distinct characteristics, one of them is dysregulated cell production. ¹⁻³ It is caused

^aDepartment of Pharmaceutical Sciences, Rashtrasant Tukadoji Maharaj Nagpur University, Mahatma Jyotiba Fuley Shaikshanik Parisar, Nagpur-440033, India. E-mail: prafullasable@yahoo.com; nusratsayyad11@gmail.com; Tel: +919158537050

^bDepartment of Pharmacognosy, College of Pharmacy, Taif University, P.O. Box 11099, Taif 21944, Saudi Arabia. E-mail: abuali@tu.edu.sa

Department of Pharmaceutics, Dadasaheb Balpande College of Pharmacy, Rashtrasant Tukadoji Maharaj Nagpur University, Nagpur, Maharashtra, 440037, India. E-mail: vidyasabale@gmail.com

^dDepartment of Pharmacology, Dadasaheb Balpande College of Pharmacy, Rashtrasant Tukadoji Maharaj Nagpur University, Nagpur, Maharashtra, 440037, India. E-mail: kaleemmubin88@gmail.com

*Department of Biology, College of Science and Arts at Alkamil, University of Jeddah, Saudi Arabia. E-mail: tasar@uj.edu.sa

Department of Pharmaceutical Chemistry, College of Pharmacy, Taif University, P.O. Box 11099, Taif 21944, Saudi Arabia. E-mail: amrathore@tu.edu.sa

*Department of Pharmaceutics, Faculty of Pharmacy, Northern Border University, Arar, Saudi Arabia. E-mail: M.Mujtaba@nbu.edu.sa

^hDepartment of Pharmaceutics, College of Pharmacy, Prince Sattam Bin Abdulaziz University, P.O. Box 173, Al-Kharj 11942, Saudi Arabia. E-mail: m.anwer@psau. edu.sa

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by an imbalance between the expression of oncogenes and proto-oncogenes, which is caused by genetic and epigenetic abnormalities.4,5 Globally, the number of cancer diagnoses has climbed to 19.3 million, and each year, 10 million deaths related to cancer are reported. The most common type of cancer among women is breast cancer (BC), globally.6 BC is a highly complex disease with distinct biological subtypes and several targeted prognostic markers of therapeutic significance.7 Approximately 2.3 million (11.7%) new instances of breast cancer were reported in 2020, and 684 996 (6.9%) fatalities were linked to the disease. Furthermore, It is predicted that by 2040, there will be around 3 million new cases of breast cancer and one million more deaths from the disease.8 The prognosis and clinical manifestations vary considerably amongst patients. BC develops through multiple pathways, including aromatase enzyme-regulated estrogen synthesis that primarily affects the expression of aromatase enzyme is highest in or near breast tumor sites and is known to catalyze the last step of conversion of androgen to estrogen.9 Elevated expressions of estrogen and aromatase in breast cancer tissues than non-cancerous have been reported by several studies.10 Tamoxifen has been used as standard endocrine therapy for a decade, however, the development of third-generation aromatase inhibitors (Anastrozole, Letrozole, Exemestane) has greater efficacy and improved disease-free survival compared to tamoxifen.11 Although many

potential aromatase inhibitors have been published. A few numbers are undergoing clinical studies to treat breast cancer but there are possible serious side effects such as loss of bone density, cardiac events, joint and muscle pain, and hot flashes. Many novel series of compounds have been synthesized and their anticancer efficacy was assessed using sorafenib as parent structure.12 The Diaryl urea is a prominent pharmacophore in building anticancer moiety.13 This activity due to pharmacophore screening shows its near-perfect binding with certain acceptors.14 The protons on two nitrogen and oxygen atoms present in urea moiety act as hydrogen bond donors and hydrogen bond acceptors respectively.¹⁵ For drug design and development molecular hybridization is an attractive area in medicinal chemistry that aims to combine two or more pharmacophores for the development of a single compound with improved efficacy.16 From a synthetic view, it was of interest to combine heterocycles to benzamide through a urea linker bridge to develop more potent antiproliferative agents. 17,18 Concurrently sorafenib and imatinib, as a potent antiproliferative drug, have structural similarities. 19 Dasatinib, a USFDA-approved drug has been accepted as a chemotherapeutic anticancer agent.20 They all include a diphenyl skeleton linked by an amide or urea, and aromatic heterocyclic rings.21 In medicinal chemistry, the amide group is immensely essential for biological molecules due to their improved solubility in the biological environment and bioavailability.22-24 Amide derivatives were associated with a variety of biological activities like anticonvulsant, anti-tuberculosis, insecticidal, antifungal, antimicrobial, anti-HCV, and antiproliferative properties.²⁵⁻²⁷ The intermolecular forces have increased importance in medicinal chemistry and allow them to interact with a variety of enzymes and receptors due to their high solubility, ease, and efficiency in reaching target sites.28-30

2. Material and methods

2.1. Molecular docking

The structures of each designed novel derivative and native ligand have been drawn using ChemDraw Ultra 16.0. (mole. File format). Universal Force Field (UFF) was utilized to carry out the energy minimization or optimization. All the compounds were docked to the crystal structure of human placental aromatase complexed with the breast cancer medication exemestane, using autodock vina 1.1.2 in PyRx 0.8. The crystal structure of the exemestane was procured from the RCSB Protein Data Bank (PDB) as entry 3s7s (https://www.rcsb.org/structure/3S7S).³¹

With the aid of Discovery Studio Visualizer 2019,³² The protein structure of human placental aromatase complexed with the breast cancer drug exemestane was cleaned up, optimized, and made ready for docking. This involved removing unnecessary water molecules and associated ligands. The Vina Wizard Tool in PyRx 0.8 was used to carry out the binding energy investigations. The docking investigation included all ligand molecules (PDBQT Files) and target molecules (aromatase enzymes). To specify the area for interactions around the RBD (to occupy), a three-dimensional grid box of known size (size_x = 51.602628352Ao; size_y = 62.2935613655Ao; size_z =

41.7297385081Ao) was calculated for MD simulation using Autodock tool 1.5.6 with an exhaustiveness value of 8. BIOVIA Discovery Studio Visualizer (version 19.1.0.18287) was employed to locate and record the active amino acid residues in the protein. The entire process of molecular docking has been described in our previous publication by Sayyad *et al.*³³

2.2. Chemistry

2.2.1. General chemistry. The data for the FT-IR measurements were expressed in cm⁻¹ and were obtained using IR Spirit FT-IR spectrophotometers from Shimadzu. Mass spectra were recorded with Agilent Q-TOS. The IR measurements were conducted with an Agilent Technologies spectrophotometer, and the results are reported in cm⁻¹. The ¹H and ¹³C NMR spectra in CDCl₃ and DMSO solution were determined using spectrometers at 500 and 125 MHz frequencies. Chemical shift data was expressed in parts per million using the Varian-VXR-500S operating and Bruker Avance Neo at 500 MHz. Tetramethylsilane (TMS, $\delta = 0.00$) as an internal standard and expressed in ppm. A melting point apparatus of the VEEGO MODEL VMP-D was used to determine the melting points and are uncorrected. Thin-layer chromatography (TLC) was employed to monitor the progress of the reaction using preparatory TLC plates (Merck precoated silica GF 254).

2.2.2. Synthesis of *N***-(4-chlorophenyl)benzamide** (3). 4-Chloroaniline (2) (1 eq.) was added to a solution of benzoic acid (1) (1 eq.) and DCC (1.1 eq.) in DMF and left to agitate at room temperature for eight hours. Following the reaction's conclusion, the reaction mass was put to crushed ice. A crude product was first produced, which was filtered, cleaned with ice-cold water, and then dried. Using ethanol, the crude product is recrystallized to produce *N*-(4-chlorophenyl)benzamide (3) **https://chem.libretexts.org/@go/page/37192** (accessed Oct 7, 2021).

2.2.3. Synthesis of N-(4-ureidophenyl)benzamide (4). A mixture of N-(4-chlorophenyl)benzamide (3)(1 eq.), urea (1.1 eq.), and K_2CO_3 (2 eq.) in DMF was refluxed to obtain N-(4-ureidophenyl)benzamide. Reaction development was observed with the help of TLC. After completion of reaction material is poured on ice-water to obtain solid mass collected using filtration. Further crude is recrystallized with the help of ethanol.³⁴

2.2.4. General procedure for the synthesis of N-(4-(3-phenylureido)phenyl)benzamide derivatives (6a–g). A mixture of N-(4-ureidophenyl)benzamide (4)(1 eq.), appropriate alkyl/aryl chloride (5a–g) (1.2 eq.), and K_2CO_3 (2 eq.) in 5 ml DMF was refluxed to obtained N-(4-(3-phenylureido)phenyl)benzamide. After the reaction was finished, the reaction mixture was poured into ice water to produce a solid mass, which was subsequently collected using filtering. TLC was used to monitor the reaction condition.³⁴

2.2.4.1. 1-(4-(Benzamido)phenyl)-3-(pyridin-3-yl)urea (6a). Obtained as off white solid, yield: 69%; molecular formula: $C_{19}H_{16}N_4O_2$; molecular weight: 332 g mol⁻¹; m.p.: 258-261 °C; R_f value: 0.76; IR (cm⁻¹): 3324.8 (N-H stretching), 1241.2 (C-N stretching (Ar)) 1520.8 (N-H bending) 1672.9 (C=N stretching) 1625.1 (C=O stretching (amide)), 823.7, 689.6, 894.6 (aromatic

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C–H bending); ¹H NMR (500 MHz, CDCl₃) (chemical shift δ): 6.089 (s, –NH, 2H), 7.336, 7.353, 7.491, 7.504, 7.521, 7.558, 7.572, 7.587, 7.601, 7.619, 7.800, 7.860 (m, Ar–CH)., 7.875(s, –NH, 1H); MS: m/z exact mass: 332.13, mass found: 333.73 (M + 1).

2.2.4.2. 1-(4-(Benzamido)phenyl)-3-(thiazol-2-yl)urea (6b). Yielded as a solid off-white, yield: 71%; molecular formula: $C_{17}H_{14}N_4O_2S$; molecular weight: 338 g mol⁻¹; m.p.: 371–374 °C; R_f value: 0.53; IR (neat cm⁻¹): 3324.8 (N-H stretching), 1312.0 (C-N stretching (Ar)) 1576.7 (N-H bending) 1672.9 (C=N stretching) 1625.1 (C=O stretching (amide)), 820.0, 715.6, 894.6 (aromatic C-H bending); ¹H NMR (500 MHz, CDCl₃) (chemical shift δ): 6.029 (s, 2H, -NH urea), 7.93(s, -NH, 1H benzamide), 7.88, 7.84, 7.50, 7.39, 7.36, 7.31, 7.28, 7.25, 6.701, 6.691 (m, Ar-CH); MS: m/z exact mass: 338.08, mass found: 339.01 (M + 1).

2.2.4.3. 1-(4-(Benzamido)phenyl)-3-(thiophen-2-yl)urea (6c). Yielded as a solid off-white, yield: 60%; molecular formula: $C1_8H_{15}N_3O_2S$; molecular weight: 337 g mol $^{-1}$; m.p.: 343–346 °C; R_f value: 0.81; IR (neat cm $^{-1}$) 3324.8 (N–H stretching), 1241.2 (C–N stretching (Ar)) 1520.8 (N–H bending) 1672.9 (C–N stretching) 1625.1 (C–O stretching (amide)), 823.7, 689.6, 894.6 (aromatic C–H bending); 1 H NMR (500 MHz, CDCl $_3$) (chemical shift δ): 5.99 (s, 1H, –NH urea), 6.00 (s, 1H, –NH urea), 6.39, 6.41, 6.56, 6.59, 6.6., 6.71, 7.22, 7.25, 7.27, 7.30, 7.32, 7.36, 7.58, 7.90, 7.95(m, Ar–CH), 8.00(s, –NH, 1H benzamide); MS: m/z exact mass: 337.09, mass found: 338.41 (M + 1).

2.2.4.4. 1-(4-(Benzamido)phenyl)-3-(1H-benzo[d]imidazole-2-yl)urea (6d). Obtained as an off white solid, yield:74%; molecular formula: $C_{21}H_{17}N_5O_2$; molecular weight: 371 g mol $^{-1}$; m.p.: 415–417 °C; R_f value: 0.39; IR (neat cm $^{-1}$): 3324.8 (N–H stretching), 1285.9 (C–N stretching (Ar)) 1580.4 (N–H bending) 1677.3 (C=N stretching) 1625.1 (C=O stretching (amide)), 820.0, 708.2, 894.6 (aromatic C–H bending); 1 H NMR (500 MHz, CDCl $_3$) (chemical shift δ): 5.00 (s, 1H, –NH benzimidazole), 6.18 (s, 2H, –NH urea), 7.17, 7.19, 7.20, 7.39, 7.41, 7.44, 7.48, 7.50, 7.64, 7.70, 7.73, 7.76, 8.02, 8.05(m, Ar–CH), 8.10(s, –NH, 1H benzamide); MS: m/z exact mass: 371.14, mass found: 372.52 (M + 1).

2.2.4.5. 1-(4-(Benzamido)phenyl)-3-(1H-benzo[d]imidazole-5-yl)urea (6e). Yielded as a solid off-white, yield:76%; molecular formula: $C_{21}H_{17}N_5O_2$; molecular weight: 371 g mol $^{-1}$; m.p.: 407–409 °C; R_f value: 0.67; IR (neat cm $^{-1}$): 3324.8 (N–H stretching), 1312.0 (C–N stretching (Ar)) 1576.7 (N–H bending) 1677.3 (C=N stretching) 1625.1 (C=O stretching (amide)), 820.0, 704.5, 894.6 (aromatic C–H bending); 1H NMR (500 MHz, CDCl $_3$) (chemical shift δ): 4.92 (s, 1H, –NH benzimidazole), 6.03 (s, 2H, –NH urea), 7.21, 7.31, 7.32, 7.38, 7.40, 7.44, 7.58, 7.60, 7.62, 7.65, 7.92, 7.96, 8.01(m, Ar–CH), 8.10(s, –NH, 1H benzamide); MS: m/z exact mass: 371.14, mass found: 372.02 (M + 1).

2.2.4.6. 1-(4-(Benzamido)phenyl)-3-methylurea (6f). Yielded as a solid off-white, yield: 69%; molecular formula: $C_{15}H_{15}N_3O_2$; molecular weight: 269 g mol⁻¹; m.p.: 342–344; R_f value: 0.68; IR (neat cm⁻¹): 3324.8 (N–H stretching), 1312.0 (C–N stretching (Ar)) 1576.7 (N–H bending) 1672.9 (C—N stretching) 1625.1 (C—O stretching (amide)), 823.7, 715.6, 894.6 (aromatic C–H bending); ¹H NMR (500 MHz, CDCl₃) (chemical shift δ): 2.68 (s, 1H, –CH₃ methyl), 6.01 (s, 2H, –NH urea), 7.32, 7.35, 7.38, 7.40,

7.44, 7.59, 7.90, 7.93(m, Ar–CH), 8.00(s, –NH, 1H benzamide); MS: m/z exact mass: 269.12, mass found: 270.14 (M + 1).

2.2.4.7. 1-(4-(Benzamido)phenyl)-3-o-tolylurea (6g). Yielded as a solid off-white, yield: 73%; molecular formula: C₂₁H₁₉N₃O₂; molecular weight: 345 g mol^{-1} ; m.p.: 445-447; R_f value: 0.59; IR (neat cm⁻¹): 3324.8 (N-H stretching), 1312.0 (C-N stretching (Ar)) 1576.7 (N-H bending) 1654.9 (C=N stretching) 1625.1 (C= O stretching (amide)), 820.0, 715.6, 894.6 (aromatic C-H bending); ¹H NMR (500 MHz, CDCl₃) (chemical shift δ): 3.15 (s, 1H, -CH₃ methyl), 6.00 (s, 1H, -NH urea), 6.03 (s, 1H, -NH urea), 6.72, 6.76, 6.80, 6.92, 6.97, 6.99, 7.30, 7.33, 7.37, 7.40, 7.45, 7.57, 7.92, 7.97(m, Ar-CH), 8.01(s, -NH, 1H benzamide); ¹³C NMR (125 MHz, DMSO) 21.0(-CH₃), 111.56(-CH), 117.54(-CH), 121.77(-CH), 123.97(-CH), 125.18(-CH), 126.39(-CH), 127.17(-CH), 128.33(-CH), 128.42(-CH), 127.58(-CH), 128.65(-CH), 128.83(-CH), 128.97(-CH), 129.20(-CH), 131.61(-CH), 134.0(-CH), 134.78(-CH), 138.06(-CH), 155.36(-CO urea), 165.57(-CONH). MS: m/z exact mass: 345.15, mass found: 346.19 (M + 1).

2.3. Biological evaluation

The biological research has been conducted at the "National Centre for Cell Science" in Pune, India. All cell lines were purchased from the National Centre for Cell Science in Pune, Maharashtra, India. The cancer cell lines *viz* A-498 (renal carcinoma), NCI-H23 (lung carcinoma), A549 (lung adenocarcinoma), MCF-7 (breast adenocarcinoma), and MDA-MB-231 (breast adenocarcinoma), were used for study at the National Centre for Cell Science (Pune, India).

The cell lines were processed in a lab as soon as they were delivered, and the first cells to be processed were cryopreserved in liquid nitrogen for subsequent use. After being meticulously cared for by guidelines, the cultivated cell lines reached maturity in less than eight weeks. Approximately 70–80% of the cells from grown tissues that were sown during the tests were confluent. The cell lines were preserved in complete media containing "Dulbecco's modified Eagle medium (Cell Clone Genetix brand, Catalogue No.: CC3004) along with penicillinstreptomycin (50 U ml⁻¹, 50 mg ml⁻¹; HiMedia, Catalogue No.: A002) and 10% fetal bovine serum by maintaining air (95%) CO₂ (5%) at a temperature of 37 °C.

2.3.1. In vitro anticancer activity (MTT assay). The cytotoxicity of all the synthesized compounds was evaluated using the MTT (3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide) assay against different cell lines.35 A-498 (renal carcinoma), NCI-H23 (lung carcinoma), A549 (lung adenocarcinoma), MDA-MB-231 (breast adenocarcinoma), and MCF-7 (breast adenocarcinoma) cell line were used, doxorubicin served as a good control. At the National Centre for Cell Science in Pune, Maharashtra, India, all cell line analyses were carried out. In brief, 96-well plates were filled with cells that were growing logarithmically at the following densities: A-498: 5000 cells per well, NCI-H23: 5000 cells per well, MDA-MB-231: 10 000 cells per well, MCF-7: 5000 cells per well, and A549: 5000 cells per well. The cells were then kept at 37 °C for 24 h in a humid atmosphere with 5% CO₂. The appropriate DMSO dilution was used as the vehicle control before test chemicals were added to wells

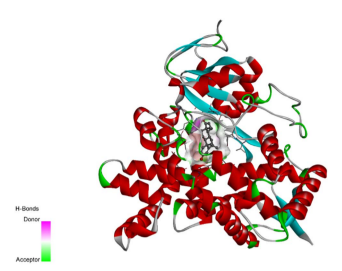


Fig. 1 The binding pocket of human aromatase enzyme with the cocrystallized ligand.

in triplicate. Following a 72 hours exposure to the test chemical in a humidified environment at 37 °C with 5% CO₂, at the end of the incubation time, twenty microliters of MTT reagent (5 mg

ml $^{-1}$) were added to each well. It was then incubated for another two to four hours at 37 °C in a 5% CO $_2$ environment. After the supernatant was removed, the cells were lysed and the formazan needles were dissolved using 200 µl of DMSO. The absorbance was then calculated using an epoch microplate reader at 570 nm. All cell lines were initially screened using test chemical concentrations of 10 and 25 µM. The ones that had an inhibition level of greater than 50% were moved on for GI $_{50}$ analysis (test chemical concentration inhibiting 50% of the cell population). Nine different concentrations (0.05, 0.1, 5, 1, 5, 10, 30, 50, and 100 M) were tested three times to ascertain the test substances' GI $_{50}$ value for the respective cell lines. Regression analysis was then used to determine the GI $_{50}$, which was then expressed in mM using a mean of three replicates. 36,37

2.4. In silico physicochemical, and ADMET predictions

By predicting *in silico* physicochemical characteristics, adsorption, distribution, metabolism, excretion, and toxicity (ADMET) properties, one can assess a drug candidate's potential. The standard approach suggested in SwissADME (http://swissadme.ch) was used to carry out the ADMET analysis.³⁸ The molecule has been uploaded to the 'ADMET Evaluation'

Table 1 Docking data of designed benzimidazole derivatives showing the ligand energy and the binding energy towards the aromatase (3s7s) binding site

Compound code	Structure	Ligand energy (kcal mol ^{–1})	Binding energy $(kcal mol^{-1})$	rmsd/ub	rmsd/lb
Native ligand	O H H	1113.62	-8.9	0	0
D1		250.47	-8.4	0	0
D2	O H O N O N O N O N O N O N O N O N O N	277.44	-8.4	0	0
D3 (6a)	O H O N O N O N O N O N O N O N O N O N	287.84	-8.2	0	0
D4		275.03	-8	0	0
D5 (6b)		458.41	-8	0	0

Table 1 (Contd.)

Compound code	Structure	Ligand energy (kcal mol ⁻¹)	Binding energy (kcal mol ⁻¹)	rmsd/ub	rmsd/lb
D6	N N N CH3	575.4	-7.2	0	0
D7 (6c)	H O S	402.26	-8.1	0	0
D8	H N N N N N N N N N N N N N N N N N N N	420.02	-8.3	0	0
D9 (6d)		514.81	-7.9	0	0
D10 (6e)		528.5	-7.9	0	0
D11 (6f)	NH N	175.12	-7.8	0	0
D12	NH N	174.28	-7.4	0	0
D13	D H O N	172.5	-6.4	0	0
D14		177.15	-7.9	0	0
D15	HO HO	274.78	-7.9	0	0
D16	OH NOH	284.12	-8.6	0	0
D17	OH OH	300.1	-8.3	0	0

Table 1 (Contd.)

Compound code	Structure	Ligand energy (kcal mol ⁻¹)	Binding energy (kcal mol ⁻¹)	rmsd/ub	rmsd/lb
D18		276.5	-8.7	0	0
D19 (6g)	H H H H	275.96	-8.6	0	0
D20		227.72	-8.6	0	0

Scheme 1 Synthesis of N-(4-(3-phenylureido)phenyl)benzamide derivatives.

option of the 'services' section of the ADMETLab 2.0 server. Many of the ADMET attributes of a molecule were calculated and shown by the server after submission.

3. Results

3.1. Molecular docking

The crystal coordinates of the aromatase enzyme were taken from **rcsb.org** with PDB ID: 3s7s and the designed molecules (**D1–20**) were docked into the binding pocket of the enzyme (Fig. 1). The docking scores of the compounds were compared with the co-crystallized ligand where the designed molecules showed good binding scores with good binding affinities. The docking scores of the native ligand and all of its derivatives are listed in Table 1. The comprehensive docking data including residues of active amino acids, type of bond, bond length (Å), and bond class involved in the interactions are illustrated in table **D1**, **D2** and **D3** of the (ESI data†).

3.2. Chemistry

The synthesis of *N*-(4-(3-phenylureido)phenyl)benzamide (**6a–g**) derivatives is depicted in Scheme 1. Benzoic acid (**1**) was coupled with 4-chloroaniline (**2**) in the presence of DCC resulting in *N*-(4-chlorophenyl)benzamide (**3**), which was then

Table 2 Compound screening at% viability at $10 \mu M$

	% Viability at 10 μM						
Comp. code	A-498	NCI-H23	MDAMB-231	MCF-7	A-549		
6a (D3)	57.16	67.17	81.60	79.62	57.67		
6b (D5)	59.16	81.16	91.27	90.72	71.81		
6c (D7)	71.12	69.19	86.95	75.15	77.13		
6d (D9)	67.28	75.18	91.38	89.18	92.12		
6e (D10)	81.16	92.83	90.13	89.13	79.17		
6f (D11)	75.15	94.07	84.42	90.17	78.16		
6g (D19)	54.16	61.68	50.10	57.47	65.16		

Table 3 Determination of Gl₅₀ for **6g**

$ ext{GI}_{50} \pm ext{SD}$							
Entry	Comp. code	A498	NCI-H23	MDAMB-231	MCF-7	A-549	
1 2	6g (D19) Doxorubicin	$\begin{array}{c} 14.46~\mu M \pm 0.06079 \\ 3.45~\mu M \pm 0.09215 \end{array}$	$\begin{array}{c} 13.97 \; \mu M \pm 0.07182 \\ 3.05 \; \mu M \pm 0.04911 \end{array}$	$\begin{array}{c} 11.35~\mu M \pm 0.05929 \\ 3.06~\mu M \pm 0.04619 \end{array}$	$\begin{array}{c} 11.58 \; \mu M \pm 0.05363 \\ 2.70 \; \mu M \pm 0.02131 \end{array}$	$\begin{array}{c} 15.77~\mu\text{M} \pm 0.05613 \\ 2.97~\mu\text{M} \pm 0.04133 \end{array}$	

refluxed for 12 h with urea in dimethyl formamide in the presence of potassium carbonate to form N-(4-ureidophenyl) benzamide (4). This N-(4-ureidophenyl)benzamide (4) was then refluxed with substituted alkyl/aryl halides (5a–g) in the presence of potassium carbonate to afford the titled N-(4-(3-phenylureido)phenyl)benzamide (6a–g) derivatives.

3.3. Biological evaluation

3.3.1. MTT assay. The in vitro cytotoxicity study on synthesized compounds was done using doxorubicin as the reference standard. The cytotoxicity of the synthesized compounds was evaluated using the MTT assay against different cell lines as follows A-498 (renal carcinoma), NCI-H23 (lung carcinoma), MDA-MB-231 (breast adenocarcinoma), A549 (lung adenocarcinoma) and MCF-7(breast adenocarcinoma), tumor cell lines. Each molecule was first tested in two concentrations (10 µM and 25 µM). The screening results showed among the synthesized molecules, compound 6g exhibited >50% inhibition at 10 micromolar concentration as shown in Table 2. The obtained results demonstrated the 6g showing promising activity with a % viability of 54.16, 61.68, 50.10, 57.47 and 65.16 at 10 μM against A-498, NCI-H23, MDAMB-231, MCF-7, and A-549 respectively. The vehicle DMSO served as the negative control and doxorubicin served as the positive control.

3.3.2. Half maximal growth inhibition (GI_{50}) calculation. The potent compound 6g was further assessed to determine the GI_{50} which exhibited a significantly promising effect in the majority of the evaluated cancer cell lines. As depicted in Table 3. The vehicle DMSO served as the negative control and doxorubicin served as the positive control. Findings of the study showed the compound 6g when compared with doxorubicin showed a GI_{50} value of 14.46 μ M, 13.97 μ M, 11.35 μ M, and 11.58 μ M, 15.77 μ M against A-498, NCI-H23, MDAMB-231, MCF-7, and A-549 respectively.

3.4. In silico physicochemical and ADMET analysis

The compound's (D19) physicochemical and ADMET properties, which may be found in the ESI,† were predicted using SwissADME (refer to ESI materials†).

4. Discussion

The proliferation of estrogen-sensitive breast cancer cells and the progression of breast cancer can be attributed to the aromatase enzyme.^{39,40} Therefore, it is postulated that the reduction in aromatase enzyme expression in BC will prevent cell division and growth.⁴¹ Even though there are natural and

synthetic chemical moieties are available in the market due to various side effects and resistance toward drugs it is necessary to develop new molecules. 42 In this work, we have demonstrated that the synthesized molecules show promising percent viability at 10 and 25 μM on an array of five different cell lines. In that MCF7 breast cancer cell lines had shown good inhibitory effects. The downregulation of aromatase enzyme was linked to the inhibitory effects of on panel of cell lines, suggesting that synthesized compounds could be a bright prospect for treating breast cancer. The derivatives were made by applying reaction plans. These novel derivatives were characterized using their mass, ¹H NMR, and FTIR spectroscopic data. The ¹H NMR, MS, and FTIR spectra of the synthesized compounds were found in the anticipated range. Based on their mass spectra, every synthesized compound has a molecular ion [M + H]⁺ peak that corresponds to its molecular formula (refer to ESI materials†).

The native ligand binding mechanism of the aromatase enzymes crystal structure has been compared to the derivatives binding affinities (PDB ID: 3s7s). Exemestane, the native ligand of the enzyme, has a binding energy of -8.9 kcal mol^{-1} . The enzyme did not produce any hydrogen bonds of any kind. ILE133 (4.78 Å), CYS437 (3.59 Å), VAL370 (3.91 Å), ALA306 (4.12 Å), ILE133 (4.0 Å), ALA306 (5.25 Å), and TRP224 (5.23 Å) are all hydrophobic interaction. It was prevalent for multiple variants to form over two hydrogen bonds with the target. For synthesis and biological activity, only molecules with three or more hydrogen bonds have been chosen. The virtual screening revealed that compounds **D3**, **D5**, **D7**, **D9**, **D10**, **D11**, and **D19** were the most efficient molecules as presented in Table 4.

Compound **D3** exhibited a binding energy of -8.2 kcal mol⁻¹ and established three conventional hydrogen bonds with THR310 (2.99 Å, 2.0 Å), and ALA306 (2.52 Å). Three hydrophobic interactions have been formed with THR310 (3.71 Å), MET374 (5.10 Å), and VAL370 (5.23 Å).

Compound **D5** exhibited a binding energy of -8 kcal mol⁻¹ and established three conventional hydrogen bonds with THR310 (2.90 Å, 1.95 Å), and ALA306 (2.73 Å). It has displayed several hydrophobic interactions with THR310 (3.83 Å), MET446 (3.96 Å), MET374 (5.05 Å), VAL370 (5.24 Å), LEU152 (5.32 Å), ALA306 (4.51 Å), and ALA307 (4.24 Å). Compound **D7** has shown a binding energy of -8.1 kcal mol⁻¹ and substantiates three conventional hydrogen bonds with THR310 (2.96 Å, 1.87 Å), and ALA306 (2.62 Å). It has resulted in numerous hydrophobic interactions with the same amino acids that compound **D5** forms, including Pi-sigma, Pi-sulfur, and Pi-alkyl. Compound **D9** demonstrated -7.9 kcal mol⁻¹ binding energy and established three conventional hydrogen bonds with CYS124 (2.62 Å,

Table 4 3D Images of all synthesized derivatives with active amino acid residues in the protein

3D images D3D5**D**7 **D8** D10 D11 **MET303** D19 EU152

1.96 Å), and TYR241 (2.54 Å). Pi-sulfur, amide-Pi stacked, Pi-Pi stacked, and Pi-alkyl bonds have all been produced with CYS124 (5.56 Å), TYR241 (5.34 Å), PHE235; LYS236 (3.85 Å), VAL248 (5.46 Å), LYS249 (4.60 Å), and LYS236 (4.92 Å).

Compound **D10** exhibits a binding energy of -7.9 kcal mol⁻¹ and has shown four conventional hydrogen bonds with GLY117 (2.03 Å, 2.67 Å), GLU129 (2.53 Å), and ASN136 (2.55 Å). Additionally, it has produced one Pi-donor and one Pi-anion

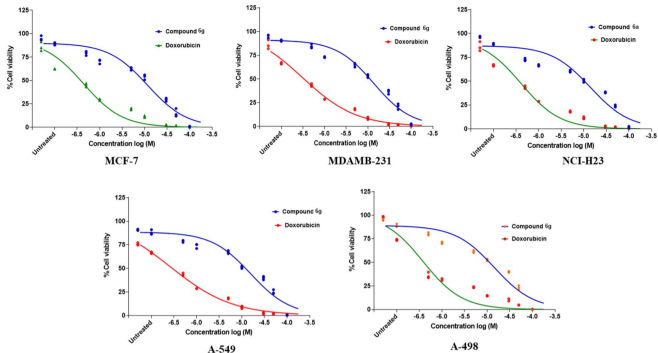


Fig. 2 Concentration-response curve for compound 6g.

hydrogen bond with GLU92 (4.05 Å) and LYS119 (3.08 Å), respectively. Compound **D11** exhibited a binding energy of -7.8 kcal mol⁻¹ and formed 4 conventional hydrogen bonds with THR310 (2.73 Å), PRO429 (2.16 Å, 2.33 Å), and VAL370 (2.77 Å). It included many Pi-sigma, Pi-sulfur, and Pi-alkyl interactions which contributed to his potency. Compound **D19** has shown a binding energy of -8.6 kcal mol⁻¹. Three conventional hydrogen bonds have been established through THR310 (3.02 Å, 1.94 Å), and ALA306 (2.77 Å). Multiple hydrophobic interactions have been developed with THR310 (3.65 Å), MET374 (5.15 Å), ALA443 (3.38 Å), MET446 (5.45 Å), VAL370 (5.18 Å), LEU152 (5.14 Å), ALA306 (4.52 Å), and ALA307 (4.29 Å).

The spectroscopic characterization of data of synthesized compounds from 6a-g can be summarized. The common core of NH stretching 3324.8 cm⁻¹ C-N aryl stretching at 1312 cm⁻¹ NH bending at 1576.7 cm⁻¹ C=O stretching at 1625.1 cm⁻¹ and aromatic C-H bending 820.0 cm⁻¹, 715.6 cm⁻¹, 894.6 cm⁻¹. The two NH present in the urea bridge come around 5.99 and 6.00 ppm and amide NH comes around 8.101 ppm. Aromatic region around 6-7 ppm. The ¹³C NMR spectrum was recorded at 125 MHz in DMSO solution, revealing distinctive chemical shifts for various carbon environments within the molecule. The spectrum features signals at 21.0 ppm, indicative of a methyl group (-CH₃), along with multiple signals between 111.56 and 138.06 ppm corresponding to different types of methylene (-CH) present in aromatic rings. Notably, at 155.36 ppm, there is a peak attributed to a carbonyl carbon (-CO) of urea functional group, and at 165.57 ppm, another peak suggests a carbon atom in an amide (-CONH) group. Biological results are shown in the Table 3 along with the concentrationresponse curve in Fig. 2. From these findings, we believe that further research is necessary to create broader-spectrum anticancer medications because of their adaptability and increased efficacy over conventional chemotherapeutic medicines like doxorubicin. To express the cell-carcinogenic selectivity *in vitro* tests were conducted to determine the compound's potential to inhibit cell proliferation on a panel of cell lines and compound **6g** (**D19**) shows promising anti-cancer activity with notable potency against MDAMB-231 cells, further optimization may be necessary to enhance its efficacy compared to the positive control, doxorubicin. These results lay the groundwork for future research to explore the therapeutic potential of compound **6g** (**D19**) and optimize its effectiveness as a potential anti-cancer agent.

5. Conclusion

Finally, a novel class of N-(4-(3-phenylureido) phenyl) benzamide ($\mathbf{6a-g}$) derivatives was prepared and tested for antitumor activity against a variety of tumor cell lines. The compound $\mathbf{6g}$ demonstrated good docking scores with noteworthy GI_{50} value and was found to have potential anti-cancer activity amongst synthesized molecules. Molecular docking analyses indicated the potential molecular interactions that underlie inhibition, which aligns with the findings of experiments. The best compounds revealed in this study have the potential as beginning points and could lead to the development of novel medication candidates for BC treatment. We have evaluated molecules on five different cell lines and in the future could be evaluated for aromatase inhibition activity.

Data availability

All the data supporting this article have been presented in the article and the ESI file. \dagger

Conflicts of interest

The authors declare no conflicts of interest.

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