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# Symmetrical di-substituted phenylamino-s-triazine derivatives as anticancer agents: in vitro and in silico approach†

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A series of symmetrical tri-substituted s-triazine derivatives were designed and synthesized by two different methods (reflux and microwave-assisted methods). The structures of compounds were determined by infrared (IR), nuclear magnetic resonance (<sup>1</sup>H NMR and <sup>13</sup>C NMR), and mass spectrometry. The yield of the microwave-assisted method (91-98%) was significantly higher (about 10%) than that of the reflux method (80-88%) meanwhile the reaction time was significantly shorter (15-30 min). Compound 3b showed good cytotoxic activity against the MCF7 (human breast cancer) cell line with an  $IC_{50}$  value of 6.19 µM. Compounds 3a and 2e showed strong cytotoxic activity against the C26 (colon carcinoma) cell line with  $IC_{50}$  values of 1.21 and 8.28  $\mu$ M, respectively. Compound 3e showed good cytotoxic activity against both MCF7 and C26 cell lines with IC<sub>50</sub> values of 13.74, and 14.66 μM respectively. In particular, compound 2d exhibited the best potent cytotoxic activity among the synthesized compounds against both MCF7 and C26 cell lines with IC $_{50}$  values of 6.54 and 0.38  $\mu$ M, respectively. Moreover, compounds 2e, 3a, and 3e showed higher selectivity on cancer cell lines and lower toxicity on BAEC (bovine aorta endothelial) normal cells compared to compounds 2d, 3a, paclitaxel, and doxorubicin. In silico studies revealed five potent compounds with good physicochemical and ADMET profiles and potent interactions with key anticancer targets (EGFR, DHFR, VEGFR2, CDK2, mTOR, and PI3K) compared to reference drugs. This work paved the way for the synthesis of more potent compounds based on the phenylamino-s-triazine scaffold and the exploration of their diverse and potential biological activities as well as their mechanisms of action.

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#### Introduction

Cancer is a dangerous disease that greatly affects human health because it is the second leading cause of death in the world.¹ According to Globocan 2022 statistics, there are about 9.7 million deaths from cancer worldwide annually, second only to deaths from cardiovascular disease, and about 20 million new cases of cancer.² Breast cancer and colorectal cancer are two common and aggressive cancers. Breast cancer is the most common malignant cancer in women, while colorectal cancer is the third most common cancer worldwide, accounting for about 10% of all cancer cases.³.⁴ Furthermore, the incidence and mortality from breast and colon cancers are projected to

Five-membered and six-membered heterocyclic nuclei exhibit a wide range of potential pharmacological activities.5-9 The s-triazine (1,3,5-triazine) nucleus is an important pharmacophore in the development of new drugs with diverse biologantiviral,10 antibacterial,11,12 antifungal, 11,12 anti-inflammatory, 13,14 antimalarial, 15,16 and anticancer. 17-24 However, the anticancer activity of s-triazine derivatives has received the most research attention because these derivatives have shown great potential in the treatment of many different types of cancer including leukemia, breast cancer, colon cancer, cervical cancer, etc.25-27 In particular, many cancer drugs containing s-triazine nucleus have been developed and have made great contributions to cancer treatment worldwide such as Altretamine (anti-ovarian cancer),28 Tretamine (antineoplastic),29 Enasidenib (antileukemia),30 Gedatolisib (treatment of neoplasm, ovary cancer, breast

increase rapidly at a rate of millions of cases per year in the future.<sup>3,4</sup> This is a huge socio-economic burden for the world, especially low- and middle-income countries. Therefore, new cancer drug development for breast and colorectal cancer treatment needs more attention with small molecule orientation to reduce costs and increase treatment efficacy.

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Some marketed anticancer drugs contain s-triazine.

cancer, advanced cancer, and endometrial cancer),31 Decitabine and Azacitidine (treat myelodysplastic syndrome and acute myeloid leukemia),32,33 and Bimiralisib (anti-breast cancer)26 (Fig. 1).

Potential s-triazine derivatives have demonstrated mechanisms of action for anticancer activity by inhibiting various enzymes, such as dihydrofolate reductase (DHFR),34,35 vascular endothelial growth factor receptor 2 (VEGFR2),36 phosphoinositide 3-kinase (PI3K),37 cyclin-dependent kinase (CDK),38 epidermal growth factor receptor (EGFR), 39,40 mammalian target of rapamycin (mTOR),41 DNA topoisomerase,42 and carbonic anhydrase.43 Many drug discovery studies have also screened through these targets to find s-triazine derivatives with potent anticancer activity.25-27

There are various methods for the synthesis of s-triazine derivatives, but the direct synthesis from simple starting materials such as cyanuric chloride (2,4,6-trichlorotriazine) is a popular and simple approach with cost-effective, and easily applicable at an industrial scale.44 The three chlorine

substituents of cyanuric chloride can be sequentially substituted at different temperatures by different N-nucleophiles (saturated and aromatic amines) to construct new drug candidates with optimum physicochemical properties and potential pharmacological activities.45

#### 1.1 Rational drug design

Tretamine, Altretamine, Dioxadet, Bimiralisib, and Gedatolisib drugs are symmetrical s-triazine derivatives with potent antitumor activity. The designed s-triazine derivatives showed common characteristics with these drugs by containing an striazine nucleus with two saturated cyclic amino groups (Fig. 2). The present study converted the amino group (cyclic-NH-) of Dioxadet to the aromatic amino groups (4-substituted aniline). Furthermore, the remaining two positions of the s-triazine nucleus were supplemented with 6-membered cyclic aliphatic amino groups (morpholine and piperidine) similar to Bimiralisib and Gedatolisib (dual inhibition of PI3K/mTOR).

Fig. 2 Rational study design of phenylamino-s-triazine derivatives from anticancer drugs.

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The aim of this study was to synthesize different symmetrical phenylamino-s-triazine derivatives containing one aromatic amine group and two hexagonal saturated cyclic amine groups and evaluate their anticancer activity against two cancer cell lines (MCF7 - human breast cancer cell line and C26 - colon carcinoma cell line) and one normal cell line (BAEC - bovine aorta endothelial cell line). The potential compounds will be further in silico molecular docking and ADMET (absorption, distribution, metabolism, excretion, and toxicity) studies to understand their potential drug-receptor interactions and pharmacokinetic parameters.

## Results and discussion

#### 2.1. Chemistry

Mono-substituted s-triazine derivatives were synthesized by a nucleophilic substitution reaction between 4-substituted aniline derivatives and cyanuric chloride in THF solvent with the presence of solid K2CO3 at low temperature of 0-5 °C to ensure the substitution of only the first chlorine atom and control the reaction selectivity (Scheme 1). The reaction occurred rapidly within 30-40 min and the reaction yield was high (91-96%). In addition, the reaction yield depended on the basicity of the aniline derivatives. More basic aniline derivatives with electron-donating substituents (4-OCH<sub>3</sub> and 4-CH<sub>3</sub>, 95-96%) showed higher yields than less basic aniline derivatives with electron-withdrawing substituents (4-F and 4-NO2, 91-92%). The yields of the mono-substituted s-triazine derivatives in this study showed similarities with some published studies (4-COOH - 95.4% (CH<sub>2</sub>Cl<sub>2</sub>/solid Na<sub>2</sub>CO<sub>3</sub>); <sup>46</sup> 4-Br, 4-Cl, 4-OCH<sub>3</sub> -92-95% (acetone/aqueous solution of NaHCO<sub>3</sub>)<sup>47</sup>) despite different reaction solvents and alkalines. However, the reaction time was recorded to be shorter than other studies (1-3 h).46,47

The tri-substituted s-triazine derivatives were prepared by nucleophilic reaction of the remaining two chlorine groups of mono-substituted s-triazine derivatives with saturated cyclic amines (piperidine and morpholine) in the presence of 1,4dioxane/solid K2CO3 by two different methods including reflux (RF) and microwave-assisted (MW) methods. The reaction yields, presented in Table 1, showed a difference between the two methods RF and MW. In general, the synthesis reaction of tri-substituted s-triazine derivatives exhibited high yields (>80%). The MW method (300 W, 15-30 min, 91-98%) exhibited shorter reaction times and about 10% higher yields than the RF method (12-24 h, 80-88%). Furthermore, the MW method used

less 1,4-dioxane solvent than the RF method. These results are similar to those in the study of Al-Zaydi et al., 2017 with the starting material of 4-COOH aniline for a tri-substituted reaction with saturated cyclic amines (piperidine and morpholine) in 1,4-dioxane/water (1:1)/Na<sub>2</sub>CO<sub>3</sub> (RF - 74.5-85.8%, 8-10 h; MW (400 W) - 88-93.1%, 10 min).46 This demonstrated that MW as the "green" method has great advantages in terms of time and yield in the synthesis of tri-substituted s-triazine derivatives, especially in limiting the use of solvents and protecting the environment.

The chemical structures of all the s-triazine derivatives were suitably elucidated by IR, 1H NMR, 13C NMR, and MS spectroscopy. The IR spectra of all compounds showed a medium absorbance band in the  $\nu$  3381–2900 cm<sup>-1</sup> region characteristic of the -NH- group and a strong absorbance band in the  $\nu$  1539-1389 cm<sup>-1</sup> region characteristic of the C=N groups of the striazine ring. In addition, the <sup>1</sup>H NMR spectra of the trisubstituted s-triazine derivatives showed a characteristic -NHproton of Ar-NH-s-triazine as a singlet in the  $\delta$  9.85-8.48 ppm region and four aromatic proton signals with two types of doublet proton (ortho coupling) in the  $\delta$  8.16–6.83 ppm region. Besides, the proton signals of the piperidine and morpholinesaturated rings showed good agreement including Hpiperidine  $\delta$  3.73–1.48 ppm (8H  $\delta$  3.73–3.68 ppm, 4H  $\delta$  1.65–1.59 ppm, and 8H  $\delta$  1.56–1.48 ppm) and H<sub>morpholine</sub>  $\delta$  3.71–3.61 ppm (8H  $\delta$  3.71–3.68 ppm and 8H  $\delta$  3.64–3.61 ppm). Furthermore,  $C_{Ar}$  ( $\delta$ 164.7–113.5 ppm),  $C_{piperidine}$   $\delta$  43.6–23.8 ppm (1C  $\delta$  43.6– 43.3 ppm, 1C  $\delta$  25.4–24.9 ppm and 1C  $\delta$  24.4–23.8 ppm) and  $C_{\text{morpholine}}$   $\delta$  66.0–43.2 ppm (1C  $\delta$  66.0–65.6 ppm and 1C  $\delta$  43.4– 43.2 ppm) were identified in the <sup>13</sup>C NMR spectrum. The molecular ion peak M (m/z) of compounds 2-3 was observed in the mass spectrum, confirming the hypothesized structure. In particular, all tri-substituted s-triazine derivatives showed physicochemical properties of fragments (MWt (molecular weight) < 500) that follow Lipinski's rules which could lead to potent compounds for further development (Table 1).

#### 2.2. In vitro anticancer activity

The IC<sub>50</sub> results (μM) on two cancer cell lines (MCF7 and C26) and one normal cell line (BAEC) are presented in Table 2. The comparison of IC50 values of the s-triazine derivatives with potential anticancer activity on the cancer cell lines MCF7 and C26 compared to the reference drugs Paclitaxel (PTX) and Doxorubicin (DOX) is shown in Fig. 3. The mono-substituted striazine derivatives (1a-1c) showed strong anticancer activity on

Scheme 1 Construction of phenylamino-s-triazine derivatives (MW: microwave irradiation, RF: reflux, THF: tetrahydrofuran).

Table 1 Yields and physicochemical parameters of 4-substituted phenylamino-s-triazine derivatives (2a-2e and 3a-3e)

	R Group					Yield	
Entry	R	$R^2$	Code	Physicochemical parame	RF	MW	
1	-Cl	Piper	2a	MWt: 372.90 nHA: 3	MR: 113.28	83	97
				nHD: 1	Log P: 5.08 TPSA: 57.18		
				nRot: 4	$\log S$ : -5.43		
2	-F	Piper	2b	MWt: 356.44 nHA: 4	MR: 108.23	85	92
				nHD: 1	Log P: 4.55		
				nRot: 4	TPSA: 57.18		
					$\log S$ : -4.99		
3	-OCH <sub>3</sub>	Piper	2c	MWt: 368.48 nHA: 4	MR: 114.76	81	98
				nHD: 1	Log P: 4.42		
				nRot: 5	TPSA: 66.41		
					Log S: -4.91		
4	$-CH_3$	Piper	2d	MWt: 352.48 nHA: 3	MR: 113.23	82	97
				nHD: 1	Log P: 4.82		
				nRot: 4	TPSA: 57.18		
					$\log S$ : $-5.14$		
5	$-NO_2$	Piper	2e	MWt: 383.45 nHA: 5	MR: 117.09	88	96
				nHD: 1	Log P: 4.28		
				nRot: 5	TPSA: 100.32		
					$\log S$ : -4.90		
6	-Cl	Mor	3a	MWt: 376.84 nHA: 5	MR: 105.83	81	98
				nHD: 1	Log P: 2.64		
				nRot: 4	TPSA: 75.64		
					$\log S$ : -3.92		
7	-F	Mor	3 <b>b</b>	MWt: 360.39 nHA: 6	MR: 100.78	84	91
				nHD: 1	Log P: 2.11		
				nRot: 4	TPSA: 75.64		
					Log S: -3.48		
8	$-OCH_3$	Mor	3 <b>c</b>	MWt: 372.42 nHA: 6	MR: 107.32	80	98
				nHD: 1	Log P: 1.99		
				nRot: 5	TPSA: 84.87		
					$\log S$ : -3.40		
9	$-CH_3$	Mor	3 <b>d</b>	MWt: 356.42 nHA: 5	MR: 105.79	81	97
	Ü			nHD: 1	Log P: 2.38		
				nRot: 4	TPSA: 75.64		
					$\log S$ : -3.63		
10	$-NO_2$	Mor	3e	MWt: 387.39 nHA: 7	MR: 109.65	86	93
	2	-		nHD: 1	Log P: 1.84		
				nRot: 5	TPSA: 118.78		
					$\log S: -3.39$		

a calculated using SwissADME, Piper - piperidinyl, Mor - morpholino, RF - reflux method (/conventional heating), MW - microwave-assisted methods, MWt - molecular weight, nHA - number of hydrogen bond acceptor, nHD - number of hydrogen bond donor, nRot - number rotatable bond, TPSA – polar surface area (Angstroms squared), MR – Molar Refractivity,  $\log P - \log P_{\text{o/w}}$  (XLOGP3),  $\log S - \log S$  (ESOL).

two cell lines MCF7 and C26 with  $IC_{50}$  values in the range of 1.77-13.46  $\mu M$  compared to the reference drugs PTX (IC<sub>50</sub> =  $2.35-4.32 \mu M$ ) and DOX (IC<sub>50</sub> =  $8.06-10.52 \mu M$ ). Compound **1e** exhibited strong anticancer activity on the MCF7 cell line (IC50 = 10.28  $\mu$ M) but weak anticancer activity on the C26 cell line (IC<sub>50</sub> = 48.34  $\mu$ M). In particular, compound 1d showed the strongest anticancer activity among the synthesized monosubstituted s-triazine derivatives with  $IC_{50} = 0.2 \mu M$  on the MCF7 cell line and is stronger than the reference drugs PTX and DOX. However, the strong toxicity on cancer cell lines of monosubstituted s-triazine derivatives may be due to the chemical reaction of the Cl group on the s-triazine nucleus. The disubstituted s-triazine derivatives can be formed at room temperature and even at low temperatures in the presence of

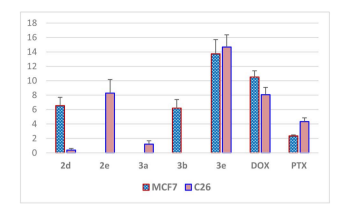
**Table 2** Anticancer activity of phenylamino-s-triazine derivatives  $(IC_{50}, \mu M)^a$ 

		Cancer cell line		Normal cell line
Entry	Code	MCF7	C26	BAEC
1	1a	$5.35 \pm 1.05$	$13.46 \pm 0.24$	_
2	1b	$7.89 \pm 1.80$	$9.10 \pm 0.61$	_
3	1c	$\textbf{1.77} \pm \textbf{0.05}$	$9.89 \pm 0.17$	_
4	1d	$0.20\pm0.11$	$9.99\pm0.59$	_
5	1e	$10.28\pm1.51$	$48.34 \pm 1.94$	_
6	2a	$23.60 \pm 2.56$	$47.52 \pm 2.86$	$48.99 \pm 3.16$
7	2b	$32.93 \pm 4.23$	$33.63 \pm 1.86$	$21.24\pm2.40$
8	2c	$50.93 \pm 8.54$	$99.40 \pm 14.07$	$15.09\pm1.98$
9	2d	$6.54 \pm 1.16 *$	$0.38 \pm 0.23 *$	$1.84 \pm 0.64 *$
10	2e	$26.11 \pm 1.86$	$8.28 \pm 1.90 *$	$315.1 \pm 9.53$
11	3a	$42.40\pm4.48$	$1.21 \pm 0.47*$	$41.23 \pm 2.55$
12	3b	$6.19 \pm 1.20 *$	$24.53\pm0.86$	$7.98 \pm 0.97 *$
13	3 <b>c</b>	$43.52 \pm 4.23$	$76.82 \pm 8.45$	$2.01\pm0.59$
14	3 <b>d</b>	$22.32\pm1.49$	$16.34\pm2.22$	$0.57\pm0.24$
15	3e	$13.74 \pm 1.96 *$	$14.66 \!\pm 1.70$	$86.34 \pm 6.79$
16	DOX	$10.52\pm0.86$	$8.06\pm1.02$	$4.79\pm0.53$
17	PTX	$2.35\pm0.14$	$4.32\pm0.52$	$\textbf{2.67} \pm \textbf{0.41}$

 $^a$  DOX – Doxorubicin, PTX – paclitaxel, MCF7 – human breast cancer cell line, C26 – colon carcinoma cell line, BAEC – bovine aorta endothelial cell line, IC $_{50}$   $\pm$  SEM ( $\mu$ M, SEM – standard error of the mean). The values in bold highlight the best compounds with the best IC $_{50}$  values compared to positive controls, \* – statistically significant (p < 0.05) compared to reference drug PTX.

base groups (-NH<sub>2</sub>) in the cell.<sup>48</sup> Therefore, mono-substituted striazine derivatives are likely to have non-selective anticancer activity and high toxicity to normal cells and the human body.

The tri-substituted s-triazine derivatives (2a, 3a, 2b, 2c, and 3c) exhibited weak to moderate anticancer activity against the MCF7 cell line with IC<sub>50</sub> ranging from 23.60 to 50.93  $\mu$ M. Compound 3**b** (4-F-Ar, dimorpholino) exhibited strong anticancer activity with  $IC_{50} = 6.19 \mu M$  against the MCF7 cell line compared to the reference drugs PTX (IC<sub>50</sub> MCF7 =  $2.35 \mu M$ ) and DOX (IC<sub>50</sub> MCF7 = 10.52  $\mu$ M). Meanwhile, compounds 2a, 2b, 3b, 2c, and 3c exhibited weak to moderate anticancer activity against the C26 cell line with IC<sub>50</sub> ranging from 24.53 to 99.40 μM compared to the reference drugs PTX (IC<sub>50</sub> C26 = 4.32  $\mu$ M) and DOX (IC<sub>50</sub> C26 = 8.06  $\mu$ M). Compounds 2e (4-NO<sub>2</sub>-Ar, dipiperidinyl) and 3a (4-Cl-Ar, dimorpholino) showed potent anticancer activity on the C26 cell line with IC<sub>50</sub> values of 8.28 and 1.21  $\mu$ M, respectively. Notably, compounds 2d (4-CH<sub>3</sub>-Ar, dipiperidinyl) and 3e (4-NO<sub>2</sub>-Ar, dimorpholino) showed strong anticancer activity on both MCF7 and C26 cell lines compared to the reference drugs PTX  $(IC_{50} = 2.35-4.32 \mu M)$  and DOX  $(IC_{50} = 8.06-10.52 \mu M)$ . Compound 2d had IC<sub>50</sub> values on the MCF7 and C26 cell lines of 6.54 and 0.38 μM, respectively, and exhibited stronger anticancer activity on the C26 cell line than PTX and DOX. In contrast, compound 3e had IC50 values on MCF7 and C26 cell lines of 13.74 and 14.66 µM, respectively, and exhibited weaker anticancer activity on the C26 cell line than PTX and DOX (Fig. 3A). Therefore, the study results showed that tri-substituted striazines 2d and 3e have the most potential among the synthesized s-triazine derivatives due to their strong anticancer activity against both MCF7 and C26 cell lines.



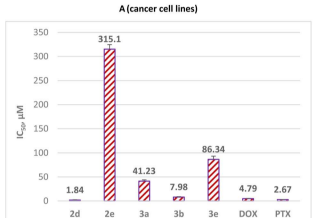


Fig. 3 The  $IC_{50}$  values ( $\mu$ M) of potential *s*-triazine derivatives compared to Paclitaxel (PTX) and Doxorubicin (DOX) (MCF7 – human breast cancer cell line, C26 – colon carcinoma cell line, BAEC – bovine aorta endothelial cell line)

In addition, all s-triazine derivatives were also tested for

B (BAEC - normal cell line)

cytotoxicity against the BAEC normal cell line to evaluate their selectivity towards cancer cell lines and systemic toxicity. The potentially active compounds 2d (IC<sub>50</sub> = 1.84  $\mu$ M) and 3b (IC<sub>50</sub> = 7.98  $\mu$ M) exhibited high cytotoxicity against BAEC normal cells similar to the reference drugs PTX (IC<sub>50</sub> = 2.67  $\mu$ M) and DOX (IC<sub>50</sub> = 4.79  $\mu$ M). Meanwhile, the potentially active *s*-triazines 2e (IC<sub>50</sub> = 315.1  $\mu$ M), 3a (IC<sub>50</sub> = 41.23  $\mu$ M), and 3e (IC<sub>50</sub> = 86.24  $\mu$ M) exhibited low cytotoxicity against BAEC normal cells compared to PTX and DOX. These results demonstrated that compounds 2e, 3a, and 3e have higher selectivity on cancer cell lines and lower toxicity to the human body compared to compounds 2d and 3b

(Fig. 3B). However, the development of targeted drug delivery

systems (liposome, transferosome, niosomes, gold nano-

particles, nanotube, etc) has significantly reduced the side effects

and toxicity of anticancer drugs.49 Therefore, good anticancer s-

triazine derivatives are potential candidates for the development

#### 2.3. Structure-activity relationships (SAR)

of new anticancer drugs.

The s-triazine derivatives with low molecular weight (MWt < 500) according to Lipinski's rule are receiving research interest

worldwide. 4-Bromophenylamino-s-triazine derivatives combined with mono-pyrazolyl and morpholino groups on the s-triazine ring showed good anticancer activity against MCF7  $(IC_{50} = 4.53 \mu M)$ , colon carcinoma (HCT-116,  $IC_{50} = 0.50 \mu M)$ , and liver carcinoma (HepG2,  $IC_{50} = 3.01 \mu M$ ).<sup>47</sup> Similarly, 4bromo/4-chlorophenylamino-s-triazine derivatives combined with indol-3-ylpyrazolyl and morpholino groups exhibited strong anticancer activity against MCF7, A549 (non-small cell lung cancer) and HDFs (non-cancerous human dermal fibroblasts) with IC<sub>50</sub> values in the range of 2-4  $\mu$ M. The IC<sub>50</sub> values of these compounds were significantly higher than doxorubicin (IC<sub>50</sub> = 0.42  $\mu$ M). This result indicates that the toxicity of these derivatives is significantly lower than that of doxorubicin towards healthy cells and may exhibit a better therapeutic index.20 Meanwhile, the sulfaguanidine-triazine derivatives with 4-sulfamoylphenylamino group showed IC50 values ranging from 14.8 to 33.2 µM against MCF7 and A549 cancer cell lines, especially the s-triazine derivative containing 4-sulfamoylphenylamino, diethylamine and morpholine could be a promising lead compound for developing new target-selected anticancer compounds (% inhibition against PI3Kα enzyme = 68%).23 Besides, the 4,6-dimorpholino-s-triazine derivatives with a 4acylphenylamino group (bearing a pyridyl group) exhibited potent cytotoxic activity against SW620 (human colon cancer cells), A549, HeLa (human cervical cancer cells), and MCF7 with IC<sub>50</sub> values of 8.71, 9.55, 15.67, and 21.77 μM, respectively.<sup>22</sup> Similar to these studies, most of the potential active s-triazines contained electron-withdrawing groups such as 4-halogeno (4-Cl and 4-F) and 4-nitro (4-NO2) on the benzene ring of the phenylamino-s-triazine scaffold. In addition, the morpholino group was also a common substituent present in the structure of lead compounds for strong anticancer activity. Furthermore, substituting dipiperidinyl and dimorpholino groups on the 4substituted phenylamino-s-triazine scaffold in the present study may enhance the anticancer activity, increase selectivity, and limit toxicity against normal cells. The symmetrical s-triazine compounds 2e, 3a, and 3e were shown to be approximately 15-118 times less toxic than PTX and indol-3-yl-pyrazolyl-striazine,20 and approximately 8.6-65.8 times less toxic than DOX (Fig. 3B). These compounds have great potential in the development of anticancer drugs with low toxicity, high selectivity,

#### 2.4. In silico ADMET profile

and better therapeutic index.

The ADMET profile of the potent compounds and reference drug Gedatolisib (Ged) is shown in Table 3.

**2.4.1 Adsorption.** The computational study of the five active compounds **2d**, **2e**, **3a**, **3b**, and **3e** showed that the physicochemical properties agreed with the directions of druglikeness rules including Lipinski (MWt  $\leq$  500,  $M \log P \leq$  4.15, nHA  $\leq$  10, and nHD  $\leq$  5), Ghose (160  $\leq$  MWt  $\leq$  480,  $-0.4 \leq$  W  $\log P \leq$  5.6,  $40 \leq$  MR  $\leq$  130, and  $20 \leq$  atoms  $\leq$  70), Veber (nRot  $\leq$  10 and TPSA  $\leq$  140), Egan ( $W \log P \leq$  5.88 and TPSA  $\leq$  131.6), and Muegge ( $200 \leq$  MWt  $\leq$  600,  $-2 \leq$   $X \log P \leq$  5, TPSA  $\leq$  150, num. rings  $\leq$  7, num. carbon > 4, num. heteroatoms > 1, nRot  $\leq$  15, nHA  $\leq$  10, and nHD  $\leq$  5) (Table 1). Moreover, the absorption

of these compounds is predicted to be good to excellent through parameters such as high MDCK permeability (>20  $\times$   $10^{-6}$  cm s $^{-1}$ ), PAMPA (the experimental data for Peff was logarithmically transformed), Pgp-substrate, HIA (human intestinal absorption),  $F_{20\%}$ , and  $F_{30\%}$  compared to the reference drug Ged. In particular, all compounds showed Caco-2 permeability higher than the reference drug Ged and higher than the -5.15 log unit. All compounds are most likely not Pgp inhibitors similar to Ged. The study results predicted that these compounds exhibited high gastrointestinal absorption and have great potential for new drug development.

**2.4.2 Distribution.** Besides, compounds **2d**, **2e**, **3a**, **3b**, and **3e** showed good plasma protein binding capacity with PPB > 94% and not as good as Ged (PPB = 76.5% < optimal value = 90%) because drugs with high protein-bound may have a low therapeutic index. Five potent compounds showed good volume distribution (optimal: 0.04–20 L kg<sup>-1</sup>) and did not have BBB (Blood–Brain Barrier) penetration similar to Ged. Brain tumor treatment is a drawback of these active compounds but the lack of BBB penetration may decrease the influx of peripheral toxins into the brain and limit side effects on the brain or central nervous system. Moreover, all compounds were non-inhibitors of OATP1B1, OATP1B3, BCRP, MRP1, and BSEP. The fraction unbound in plasms of **3b**, **3e**, and Ged (Fu = 5–20%) showed better than that of compounds **2d**, **2e**, and **3a** (Fu < 5%).

**2.4.3 Metabolism.** Regarding metabolic properties, all potent compounds showed strong inhibition of the CYP1A2 enzyme similar to Ged. Compounds **2d** and **2e** showed strong inhibition of CYP2C19 and CYP2B6 enzymes, while **3a**, **3b**, and **3e** showed moderate inhibition of CYP3A4 enzyme.

**2.4.4 Excretion.** Compounds **3a** and **3b** (5.307–5.329 mL min<sup>-1</sup> kg<sup>-1</sup>) were classified as a moderate clearance level ranging between 5 to 15 mL min<sup>-1</sup> kg<sup>-1</sup> similar to Ged (5.329 mL min<sup>-1</sup> kg<sup>-1</sup>). In contrast, compounds **2d** (4.995 mL min<sup>-1</sup> kg<sup>-1</sup>), **2e** (4.479 mL min<sup>-1</sup> kg<sup>-1</sup>), and **3e** (4.902 mL min<sup>-1</sup> kg<sup>-1</sup>) showed lower CL values and were classified as low clearance levels (CL < 5 mL min<sup>-1</sup> kg<sup>-1</sup>). Moreover, all compounds are classified as ultra-short half-life drugs ( $T_{1/2}$  < 1 h).

**2.4.5 Toxicity.** Finally, five potent compounds showed hERG blockers, hERG blockers ( $10~\mu m$ ), DILI (drug-induced liver injury), skin sensitization, carcinogenicity, human hepatotoxicity, drug-induced nephrotoxicity, ototoxicity, genotoxicity, and Hek293 cytotoxicity similar to the reference drug Ged. Hematotoxicity was possible for compounds **2e** and Ged, while compounds **2d**, **3a**, **3b**, and **3e** were predicted not to exhibit this type of toxicity. However, all compounds did not show rat oral acute toxicity, eye corrosion, and RPMI-8226 immunotoxicity. Furthermore, all compounds exhibited good "Toxicophore rules" and "Tox21 pathway" profiles as compared to Ged.

#### 2.5. In silico molecular docking study

Molecular docking is a powerful computational tool for predicting protein-ligand interaction energy scores. However, the reliability of predictions regarding non-covalent interactions varies depending on several factors such as scoring function, <sup>50,51</sup> protein and ligand flexibility, <sup>52,53</sup> water molecules and solvent

Table 3 ADMET profile of the active compounds and reference  $drug^a$ 

Parameter	2d		2e		3a		3 <b>b</b>		3e		Ged	
Absorption												
Caco-2 permeability	-4.745	E	-4.713	E	-4.769	E	-4.756	E	-4.801	E	-5.322	P
MDCK permeability	0.0	E	0.0	$\mathbf{E}$	0.0	E	0.0	$\mathbf{E}$	0.0	E	0.0	$\mathbf{E}$
PAMPA		E		$\mathbf{E}$		E		$\mathbf{E}$		E		$\mathbf{E}$
Pgp-inhibitor	+++	M	+++	P	+++	P	+++	P	++	P	+++	P
Pgp-substrate		$\mathbf{E}$		$\mathbf{E}$		$\mathbf{E}$		$\mathbf{E}$		$\mathbf{E}$		$\mathbf{E}$
HIA		E		$\mathbf{E}$		E		$\mathbf{E}$		E		$\mathbf{E}$
F <sub>20%</sub>		E		E		E		$\mathbf{E}$		E		E
F <sub>30%</sub>		E		E		E		$\mathbf{E}$		E		E
F <sub>50%</sub>		P		E	++	P	+	M	++	P	+++	P
Distribution												
PPB (%)	99.2	P	99.2	P	97.1	P	94.6	P	94.7	P	76.5	E
VDss (L kg <sup>-1</sup> )	4.428	E	3.151	$\mathbf{E}$	1.342	E	2.032	E	1.636	E	2.66	E
BBB penetration		E		E		E		E		E		E
Fu (%)	0.4	P	0.4	P	3.0	P	6.2	E	6.4	E	18.0	E
OATP1B1 inhibitor	+++	P	+++	P	+++	P	+++	P	+++	P	+++	P
OATP1B3 inhibitor	+++	P	+++	P	+++	P	+++	P	+++	P	+++	P
BCRP inhibitor		P		P		P		P		P		P
MRP1 inhibitor	+++	P	+++	P	+	M	+	M	+++	P		M
BSEP inhibitor	+++	P	+++	P	+++	P	+++	P	+++	P	+++	P
Metabolism												
CYP1A2 inhibitor	+++		+++		+++		+++		+++		+++	
CYP1A2 minutor CYP1A2 substrate	+++		+++		+++		++		+++			
CYP2C19 inhibitor	+++		+++		++				+			
CYP2C19 minibitor CYP2C19 substrate	+++		+++								+++	
	+++				++		++		+			
CYP2C9 inhibitor CYP2C9 substrate	++				++		++		+			
CYP2D6 inhibitor			+++									
CYP2D6 substrate	++		+						+			
CYP3A4 inhibitor					++		+		++			
CYP3A4 substrate	+++		+++								++	
CYP2B6 inhibitor	+++		+++								+++	
CYP2B6 substrate												
CYP2C8 inhibitor	+++		+		+++	ъ	+++	ъ.	++	ъ		ъ
HLM stability	+++	E	+++	E		P		P		P		P
Excretion		_		_						_		
CL <sub>plasma</sub> (mL min <sup>-1</sup> kg <sup>-1</sup> )	4.995	E	4.479	E	5.307	M	5.254	M	4.902	E	5.329	M
$T_{1/2}$	0.226		0.464		0.229		0.254		0.54		0.297	
Toxicity												
hERG blockers	0.555	M	0.569	M	0.564	M	0.411	M	0.43	M	0.949	P
hERG blockers (10 μm)	0.801	P	0.795	P	0.717	P	0.576	M	0.599	M	0.337	M
DILI	0.647	M	0.974	P	0.991	P	0.958	P	0.999	P	0.998	P
AMES toxicity	0.19	$\mathbf{E}$	0.697	M	0.3	M	0.517	M	0.883	P	0.545	M
Rat oral acute toxicity	0.119	$\mathbf{E}$	0.242	$\mathbf{E}$	0.117	$\mathbf{E}$	0.189	$\mathbf{E}$	0.179	$\mathbf{E}$	0.289	E
FDAMDD	0.525	M	0.624	M	0.067	$\mathbf{E}$	0.128	$\mathbf{E}$	0.09	$\mathbf{E}$	0.457	M
Skin sensitization	0.359	M	0.72	P	0.775	P	0.467	M	0.926	P	0.635	M
Carcinogenicity	0.658	M	0.71	P	0.966	P	0.978	P	0.98	P	0.947	P
Eye corrosion	0.001	$\mathbf{E}$	0.001	$\mathbf{E}$	0.0	$\mathbf{E}$	0.002	$\mathbf{E}$	0.001	$\mathbf{E}$	0.0	$\mathbf{E}$
Eye irritation	0.771	P	0.873	P	0.625	M	0.763	P	0.915	P	0.0	$\mathbf{E}$
Respiratory toxicity	0.538	M	0.748	P	0.287	$\mathbf{E}$	0.358	M	0.594	M	0.791	P
Human hepatotoxicity	0.773	P	0.829	P	0.906	P	0.916	P	0.936	P	0.986	P
Drug-induced nephrotoxicity	0.615	M	0.5	M	0.977	P	0.988	P	0.917	P	0.997	P
Drug-induced neurotoxicity	0.781	P	0.178	$\mathbf{E}$	0.937	P	0.968	P	0.439	M	0.991	P
Ototoxicity	0.354	M	0.314	M	0.524	M	0.492	M	0.431	M	0.864	P
Hematotoxicity	0.227	E	0.364	M	0.169	E	0.183	$\mathbf{E}$	0.267	E	0.475	M
Genotoxicity	0.672	M	1.0	P	0.955	P	0.958	P	1.0	P	1.0	P
										-		
•	0.072	$\mathbf{E}$	0.078	E	0.138	Е	0.159	E	0.162	$\mathbf{E}$	0.624	M
RPMI-8226 immunotoxicity A549 cytotoxicity	0.072 $0.451$	E M	0.078 0.57	E M	0.138 0.162	E E	0.159 0.076	E E	0.162 $0.103$	E E	0.624 $0.051$	M E

Table 3 (Contd.)

Parameter	2d	2e	3a	3b	3e	Ged
BCF	1.679	1.168	1.387	0.532	0.482	0.755
IGC50	3.759	3.772	3.765	3.06	3.274	3.379
LC50DM	4.558	4.802	5.118	4.709	4.307	5.31
LC50FM	4.432	4.479	4.65	3,774	3,943	4,206

<sup>a</sup> DOX – doxorubicin (MWt: 543.17, nHA: 12, nHD: 7, nRot: 5, Flex: 0.167, log P: 1.208, TPSA: 206.07, log S: −3.326), PTX – paclitaxel (MWt: 853.33, nHA: 15, nHD: 4, nRot: 15, Flex: 0.341, log P: 2.85, TPSA: 221.29, log S: −5.133), Ged – Gedatolisib (MW: 615.33, nHA: 13, nHD: 2, nRot: 10, Flex: 0.263, log P: 2.305, TPSA: 128.29, log S: −2.824), Caco-2 permeability (optimal: higher than −5.15 log unit), MDCK permeability (low permeability:  $2 \times 10^{-6}$  cm s<sup>-1</sup>, medium permeability:  $2 \times 20 \times 10^{-6}$  cm s<sup>-1</sup>, high passive permeability:  $2 \times 20 \times 10^{-6}$  cm s<sup>-1</sup>), PAMPA – the experimental data for Peff was logarithmically transformed (log  $P_{\rm eff} < 2$ : low-permeability, log  $P_{\rm eff} > 2.5$ : high-permeability), Pgp – P-glycoprotein, HIA – Human Intestinal Absorption ( $\div$ : ≥ 30%,  $\div$ : <30%), F: Bioavailability ( $\div$ : < percent value,  $\div$ : ≥ percent value), PPB: Plasma Protein Binding (optimal: <90%), VD: Volume Distribution (optimal: 0.04–20 L kg<sup>-1</sup>), BBB: Blood–Brain Barrier Penetration, Fu: The fraction unbound in plasms (low: <5%, middle: 5-20%, high: >20%), CL: Clearance (low: <5 mL min<sup>-1</sup> kg<sup>-1</sup>, moderate: 5-15 mL min<sup>-1</sup> kg<sup>-1</sup>, high: >15 mL min<sup>-1</sup> kg<sup>-1</sup>),  $T_{1/2}$  (ultra-short half-life drugs: 0.5 − <1 h; short half-life drugs: 1-4 h; intermediate short half-life drugs: 4-8 h; long half-life drugs: > 8 h), hERG blockers (IC<sub>50</sub> ≤ 10 μM or ≥ 50% inhibition at 10 μM were classified as hERG  $\div$ , DLI: Drug-Induced Liver Injury, **Rat Oral Acute Toxicity** (0: low-toxicity > 500 mg kg<sup>-1</sup>, 1: high-toxicity < 500 mg kg<sup>-1</sup>), FDAMDD – Maximum Recommended Daily Dose, BCF – Bioconcentration Factors, IGC50 – Tetrahymena pyriformis 50 percent lethal concentration. The output value is the probability of being inhibitor/substrate/active/positive/high-toxicity/sensitizer/carcinogens/corrosives/irritants (category 1) or non-inhibitor/non-substrate/inactive/negative/low-toxicity/non-sensitizer/non-carcinogens/noncorrosives/nonirritants (category 1) or non-inhibitor/substrate/lactive/positive/lon-

effects, <sup>54,55</sup> and ligand binding diversity. <sup>52,56</sup> The Autodock Vina tool can address these aspects to significantly improve the accuracy of molecular docking predictions. Specifically, molecular docking using AutoDock Vina offers several key benefits that make it a widely used tool in computational drug discovery including high speed, accuracy, automatic calculation of the grid maps, flexibility, support for multiple ligands and receptors, compatibility, and support for different force fields. <sup>57</sup>

The binding affinity and hydrogen bond formation of the *s*-triazine derivatives with 6 targets including dihydrofolate reductase (DHFR), vascular endothelial growth factor receptor 2 (VEGFR2), phosphoinositide 3-kinase (PI3K), cyclin-dependent

kinase 2 (CDK2), epidermal growth factor receptor (EGFR), mammalian target of rapamycin (mTOR) are shown in Table 4.<sup>25–27</sup> The interactions of the symmetrical tri-substituted *s*-triazine with amino acid residues at the active site of anticancer targets are shown in Fig. 4–6.

Compound 2d (4-CH<sub>3</sub>-phenylamino and piperidinyl) exhibited binding affinities ranging from -9.1 to -10.2 kcal mol<sup>-1</sup> and formed strong hydrogen, or carbon-hydrogen bonds, with all six tested anticancer targets. In addition, this compound showed greater binding affinities than the reference drugs Erlotinib, Methotrexate, and Seliciclib on three targets, namely EGFR, DHFR, and CDK2, and exhibited a binding affinity close

Table 4 In silico molecular docking results of active s-triazines and reference drugs<sup>a</sup>

	EGFR		DHFR CD		CDK	CDK2 V		VEGFR2		mTOR		PI3K	
Entry	Compound	BA	Туре	BA	Туре	BA	Туре	BA	Туре	BA	Туре	BA	Туре
1	2d	-9.8	1 CHB	-9.1	1 CHB	-9.1	1 CHB	-9.5	1 HB, 1 CHB	-9.4	1 HB, 1 CHB	-10.2	1 HB
	2e	-9.7	1 HB	-9.6	2 HB, 1 CHB	-9.3	1 HB, 1 DHB	-9.5	1 HB, 1 CHB	-9.1	1 HB	-9.8	3 HB
2	3a	-9.2	2 HB, 1 CHB	-8.5	2 CHB	-8.8	_	-9	1 HB, 2 CHB	-8.9	1 HB, 2 CHB	-9.4	1 CHB
3	3b	-9.3	1 HB, 1 CHB	-8.5	1 HB, 1 CHB	-8.9	1 HB, 2 CHB	-8.8	1 HB, 4 CHB	-8.8	2 HB, 4 CHB	-9.6	2 HB, 1 CHB
	3e	-9.1	1 HB, 1 CHB	-9.3	3 HB, 3 CHB	-8.7	1 HB	-8.6	1 HB, 3 CHB	-8.7	1 HB, 2 CHB	-9.0	2 HB, 1 CHB
4	Co-crystallized	-7.4	_	-8.7	_	-8	_	-9.4	_	-7.3	_	-8.7	_
	ligand												
5	Erlotinib	-7.7	4 HB, 2 CHB	_	_	_	_	_	_	_	_	_	_
6	Methotrexate	_	_	-8.3	6 HB, 2 CHB	_	_	_	_	_	_	_	_
7	Seliciclib	_	_	_	_	-7.8	1 HB, 1 CHB	_	_	_	_	_	_
8	Axitinib	_	_	_	_	_	_	-9.7	1 HB, 1 CHB	_	_	_	_
9	Gedatolisib	_	_	_	_	_	_	_	_	-9.4	3 HB, 3 CHB	-10.6	4 HB, 5 CHB

 $<sup>^</sup>a$  BA – binding affinity (kcal mol $^{-1}$ ), HB–hydrogen bond (conventional/strong hydrogen bond), CHB – carbon–hydrogen bond, DHB –  $\pi$ –donor hydrogen bond, EGFR – epidermal growth factor receptor, DHFR – dihydrofolate reductase, CDK2 – cyclin-dependent kinase 2, VEGFR2 – vascular endothelial growth factor receptor 2, mTOR – mammalian target of rapamycin, PI3K – phosphoinositide 3-kinase.

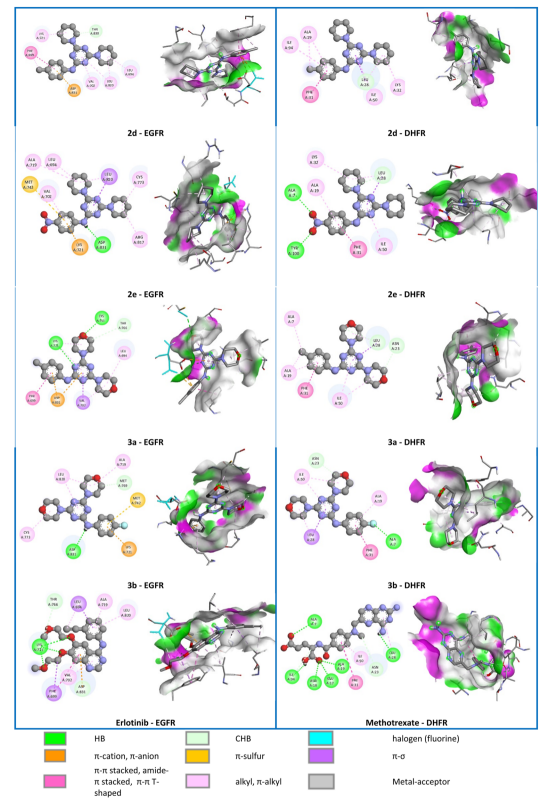


Fig. 4 2D and 3D representation of the interaction of s-triazine derivatives and reference drugs with epidermal growth factor receptor (EGFR) and dihydrofolate reductase (DHFR) targets.

to that of the reference drugs Axitinib and Gedatolisib on three targets, namely VEGFR2, mTOR, and PI3K. Meanwhile, compound **2e** (4-NO<sub>2</sub>-phenylamino and piperidinyl) exhibited

binding affinities ranging from -9.1 to -9.8 kcal mol<sup>-1</sup> and formed strong hydrogen, or  $\pi$ -donor hydrogen bonds, or carbon-hydrogen bonds with all six tested anticancer targets.

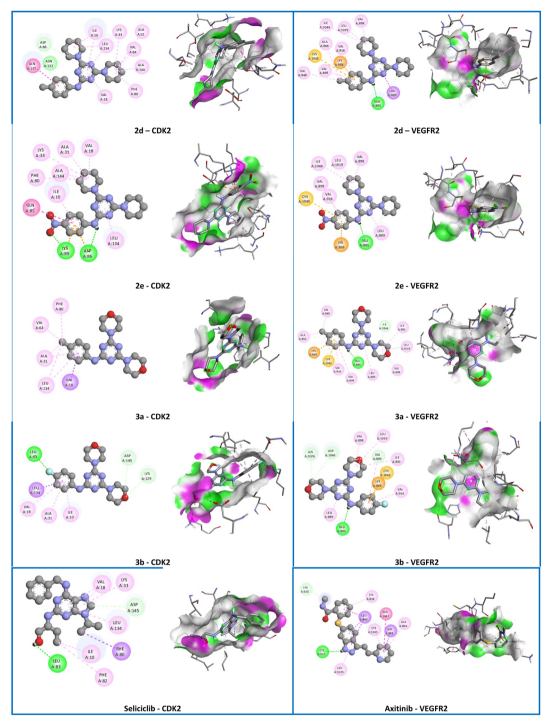


Fig. 5 2D and 3D representation of the interaction of s-triazine derivatives and reference drugs with cyclin-dependent kinase 2 (CDK2) and vascular endothelial growth factor receptor 2 (VEGFR2) targets.

Similar to compound **2d**, this compound exhibited greater binding affinity than reference drugs Erlotinib, Methotrexate, and Seliciclib on 3 targets such as EGFR, DHFR, and CDK2, and nearly equal to the binding affinity of reference drug Axitinib on VEGFR2 target. However, compound **2e** exhibited lower binding affinity than Gedatolisib and **2d** on two targets mTOR and PI3K. Compounds **3a** (4-Cl-phenylamino and morpholino), **3b** (4-Fphenylamino and morpholino) and **3e** (4-NO<sub>2</sub>-phenylamino

and morpholino) exhibited binding affinities ranging from -8.5 to -9.3 kcal mol<sup>-1</sup> and were greater than the reference drugs Erlotinib, Methotrexate and Seliciclib on the three targets EGFR, DHFR, and CDK2. In contrast, these compounds exhibited lower binding affinities than the reference drugs Axitinib and Gedatolisib on three targets VEGFR2, mTOR, and PI3K. Therefore, the study results suggested that all potent compounds have a multitarget anticancer mechanism of action

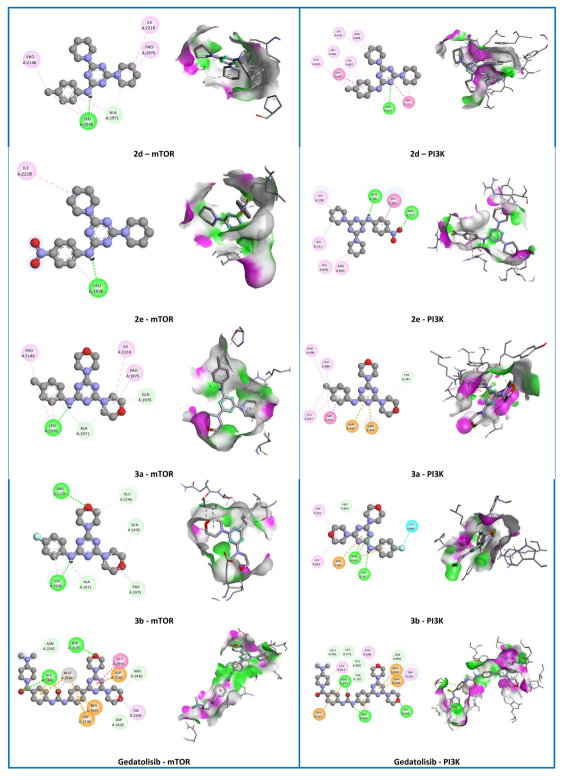


Fig. 6 2D and 3D representation of the interaction of s-triazine derivatives and reference drugs with mammalian target of rapamycin (mTOR) and phosphoinositide 3-kinase (PI3K) targets.

and have a great advantage in limiting resistance to current cancer drugs.

At the EGFR active site, reference drug Erlotinib formed four strong hydrogen bonds (HBs) with amino acid LYS721 (2.05–

2.92 Å), two carbon-hydrogen bonds (CHB) with amino acids THR766 (3.75 Å) and ASP831 (3.48 Å), one electrostatic ( $\pi$ -anion) interaction with ASP831 (3.85 Å), and hydrophobic interactions ( $\pi$ - $\sigma$ ,  $\pi$ - $\pi$  stacked, alkyl, and  $\pi$ -alkyl) with amino acid residues

LEU694, PHE699, LEU694, VAL702, ALA719, and LEU820 (3.47-5.56 Å). Like Erlotinib, compounds 3a and 3e formed HBs with amino acid LYS721 (2.42-2.53 Å). Meanwhile, compounds 3b and **2e** formed electrostatic ( $\pi$ -cation) interactions with bond lengths ranging from 4.14 to 4.37 Å, and 2d exhibited hydrophobic (alkyl) interaction with amino acid LYS721 with a bond length of 5.10 Å. In addition, compounds 2d, 3a and 3e exhibited electrostatic ( $\pi$ anion) interactions with ASP831 (3.37-4.24 Å) similar to Erlotinib. Furthermore, compounds 2d, 3a and 3e exhibited 6-7 hydrophobic interactions with amino acids similar to Erlotinib such as PHE699, LEU694, LYS721, LEU820, PHE699, VAL702, and LEU820 (3.67-5.28 Å). Docking analysis also revealed that LYS721 and ASP831 located in the EGFR binding pocket play a key role in the binding mode of the ligand to the receptor. Therefore, compounds 3a and 3e showed the most similar interactions compared to the reference drug Erlotinib at the EGFR active site.

At the DHFR active site, the reference drug Methotrexate formed many interactions including 6 HBs with amino acid residues ALA7 (2.80 Å), GLU17 (2.18 Å), ASN18 (2.08 Å), ALA19 (2.41 Å), ILE94 (2.79 Å), and LEU28 (3.05 Å); 2 CHBs with amino acids ASN23 (2.89 Å) and ILE94 (3.43 Å) and hydrophobic interactions ( $\pi$ – $\pi$  T-shaped and  $\pi$ –alkyl) with amino acids PHE31 (5.54 Å), ILE50 (4.39 Å) and LEU28 (5.49 Å). Similar to Methotrexate, all five compounds formed CHBs with LEU28 or/and ASN23 (3.16-3.52 Å) and exhibited hydrophobic interactions with amino acids PHE31, ILE50, and LEU28 (3.80-5.48 Å). Additionally, compounds 2e and 3b formed one HB with ALA7 (2.61-2.81 Å), and 3e formed two HBs with ALA7 (2.71 and 2.84 Å). However, all five compounds only showed hydrophobic interactions (alkyl and  $\pi$ -alkyl) with ALA19 (3.83–5.04 Å) and these interactions were weaker than the HB of Methotrexate at ALA19.

At the CDK2 active site, the reference drug Seliciclib exhibited some interactions such as one HB (LEU83, 1.93 Å), one CHB (ASP145, 3.38 Å), and hydrophobic interactions ( $\pi$ - $\sigma$ , alkyl, and  $\pi$ -alkyl) with amino acids PHE80, ILE10, PHE82, VAL18, LEU134, VAL18, and LYS33 (3.70-5.09 Å). Compounds 2d, 2e, 3d, and 3e exhibited hydrophobic interactions ( $\pi$ - $\sigma$  or  $\pi$ -alkyl) with amino acids VAL18, ILE10, and LEU134 (3.76-5.36 Å), whereas compound 3b exhibited hydrophobic interactions (alkyl and  $\pi$ -alkyl) with amino acids LEU134 and PHE80 (4.62– 5.40 Å) similar to Seliciclib. Additionally, compound 2d exhibited the highest number of hydrophobic interactions similar to Seliciclib at amino acids such as PHE80, ILE10, VAL18, LEU134, VAL18, and LYS33 (4.24–5.46 Å). However, this compound did not form HB with LEU83 compared to 3b, 3e, and Seliciclib.

At the VEGFR2 active site, the reference drug Axitinib showed some interactions such as one HB (CYS919, 2.22 Å), one CHB (LYS920, 3.59 Å), and hydrophobic interactions ( $\pi$ - $\sigma$ ,  $\pi$ - $\pi$  Tshaped, and  $\pi$ -alkyl) with the amino acids LEU840, VAL848, PHE1047, LEU1035, ALA866, CYS1045, and LYS838 (3.58-5.42 Å). Compounds 2d, 2e, 3a, and 3b showed  $\pi$ -sulfur hydrophobic interactions with CYS1045 (5.51-5.79 Å) similar to the reference drug Axitinib. Moreover, compound 3e exhibited the most similar interactions with amino acid residues against VEGFR2 compared to Axitinib including one HB (CYS919, 2.41 Å), 1 CHB (LEU840, 3.76 Å), and hydrophobic interactions ( $\pi$ - $\sigma$ 

and  $\pi$ -alkyl) with LEU840, ALA866, CYS919, and LEU1035 (3.81-5.38 Å).

At the PI3K active site, the reference drug Gedatolisib formed many interactions including four HBs with amino acids ARG277 (2.56 and 2.41 Å), HIS658 (2.48 Å), and GLU880 (2.28 Å); five CHBs with GLN846 (3.50 Å), CYS869 (3.56 Å), TYR787 (3.35 Å), GLU301 (3.71 Å), and CYS275 (3.56 Å); five electrostatic ( $\pi$ cation and  $\pi$ -anion) interactions with amino acids ARG277 (4.64 Å), ARG690 (4.21 Å), ARG849 (3.85 Å), GLU826 (4.82 Å), and GLU880 (4.15 Å) and hydrophobic interactions (alkyl and  $\pi$ alkyl) with amino acids LEU823, TRP201, HIS658 and PHE694 (4.50-5.31 Å). Compounds 2d, 3b, and 3e exhibited the highest number of interactions (6-8 interactions) with amino acids similar to Gedatolisib such as PHE694, TYR787, CYS869, ARG690, TRP201, GLU880, ARG849, and HIS658 (2.07-5.45 Å). In addition, compound 2e formed two HBs with ARG277 with bond lengths ranging from 2.67 to 2.94 Å similar to Gedatolisib, however, this compound only formed hydrophobic interactions (alkyl and  $\pi$ -alkyl) with amino acids ARG849 and HIS658 (4.43-4.80 Å) compared to Gedatolisib. Meanwhile, all tested compounds did not show interactions with amino acids similar to Gedatolisib at the mTOR active site.

In summary, in terms of hydrogen bond formation, compound 3b formed the most hydrogen bonds with 5-6 hydrogen bonds (1-2 HBs and 4 CHBs) with bond lengths ranging from 2.17 to 3.76 Å on the VEGFR2 and mTOR targets compared to other compounds and reference drugs. Compound 2e formed three HBs (2.52-2.94 Å) at the PI3K active site. Notably, compound 3e formed three HBs (2.39-2.84 Å) and 3 CHBs (3.17-3.57 Å) on the DHFR target compared to other compounds (1-3 HBs, 2.35-5.48 Å) and reference drug Methotrexate (6 HBs and 1 CHB, 2.08-3.43 Å). Furthermore, all the potential derivatives exhibited interactions (hydrogen bond, electrostatic ( $\pi$ -ion), hydrophobic interactions ( $\pi$ - $\sigma$ ,  $\pi$ - $\pi$  Tshaped, alkyl,  $\pi$ -alkyl)) with some crucial amino acid residues similar to the reference drugs at the action site of the five targets, except for mTOR. In addition, the 4-phenylamino group of the s-triazine nucleus showed many strong interactions with the 6 tested anticancer targets. Therefore, this group is an important structural part that enhances the anticancer activity of s-triazine derivatives and exhibits multitarget interaction properties. In conclusion, five compounds 2d, 2e, 3a, 3b, and 3e exhibited potential anticancer activity as demonstrated through in vitro and in silico studies.

#### Conclusion 3.

In summary, ten tri-substituted s-triazine derivatives have been designed, synthesized, and evaluated for in vitro and in silico anticancer activities. The microwave-assisted method showed excellent yields (>90%) and significantly reduced reaction time compared to the reflux method. In addition, five compounds showed potent anticancer activity with IC<sub>50</sub> < 15  $\mu$ M against two tested cell lines (MCF7 and C26) compared to reference drugs PTX and DOX. From the structure-activity relationship, the most potent s-triazine derivatives contained electronwithdrawing groups such as 4-halogeno and 4-nitro on the benzene ring and morpholino group of the phenylamino-striazine scaffold. Three symmetrical s-triazines **2e**, **3a**, and **3e** showed low toxicity on BAEC normal cell lines compared to PTX and DOX. Molecular docking demonstrated that five potent compounds showed good hydrogen and hydrophobic interactions against multi-targets such as DHFR, VEGFR2, PI3K, CDK2, EGFR, and mTOR that resemble the co-crystallization ligand, and reference drugs. Furthermore, the obtained *in silico* results predicted that these compounds may show a good ADMET profile. Therefore, these s-triazines are potential candidates for further research in the development of new anticancer drugs.

## 4. Experimental section

#### 4.1. Materials

All chemicals and solvents were purchased from commercial suppliers Merck and Acros. All the reactions were carried out under an inert atmosphere of nitrogen. TLC was performed on pre-coated aluminum sheets of silica (60  $F_{254}$  nm) and visualized by shortwave UV light at  $\lambda$  254. Column chromatography uses 0.040–0.063 mm granular silica gel (Merck).

The microwave reactor used was the Microwave synthesizer – CEM Discover, USA, fitted with a magnetic stirrer for continuous stirring and an infrared temperature sensor, which enabled and controlled the temperature. Melting points (mp) were determined using a Sanyo-Gallenkamp melting point apparatus. IR spectra were recorded on an IRAffinity-1S. NMR spectra were recorded on a Bruker Avance 500 NMR Spectrometer. Chemical shifts were measured in  $\delta$  (ppm). Mass spectrometry was measured on 1100 series LC-MS Trap Agilent.

#### 4.2. Experimental procedures

- 4.2.1 General procedure for the preparation of monosubstituted s-triazine derivatives (1a–1e). Cyanuric chloride (7.5 mmol) was dissolved in THF (15 mL) and cooled at 0–5 °C. Then, 4-substituted aniline derivatives (5 mmol) and  $\rm K_2CO_3$  (5 mmol) were added slowly and the mixture was stirred continuously until no aromatic amine material was present on TLC (about 30–60 min). After the reaction was completed, THF was removed by a Heidolph rotary evaporator. The crude solid was purified by recrystallization with ethanol:water (1:1, v/v) solvent. Reaction yields range from 91 to 96%.
- 4.2.2 4,6-Dichloro-*N*-(4-chlorophenyl)-1,3,5-triazin-2-amine (1a). White solid, yield 96%, mp 146–148 °C. IR (v cm<sup>-1</sup>): 3254 (NH), 1383 (C=N), 2978 (ArCH), 806 (C-N, s-triazine), 791 (ArCl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 10.91 (1H, s, NH), 7.60 (2H, d, J = 9 Hz, H<sub>Ar</sub>), 7.42 (2H, d, J = 8.5 Hz, H<sub>Ar</sub>). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 163.8, 154.1, 136.0, 135.9, 128.9, 128.8, 128.4, 123.1, 122.9. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>9</sub>H<sub>6</sub>Cl<sub>3</sub>N<sub>4</sub> 274.9653, found 274.9405.
- **4.2.3 4,6-Dichloro-***N***-(4-fluorophenyl)-1,3,5-triazin-2-amine (1b).** White solid, yield 92%, mp 168–170 °C. IR (v cm $^{-1}$ ): 3385 (NH), 1541 (C=N), 1207 (Ar–F), 1011 (C–N, *s*-triazine), 791 (Ar–Cl).  $^{1}$ H NMR (500 MHz, DMSO-d $_{6}$ ,  $\delta$  ppm): 11.13 (1H, s, NH), 7.58 (2H, d, J=9.0 Hz, H $_{Ar}$ ), 7.20 (2H, d, J=9.0 Hz, H $_{Ar}$ ).  $^{13}$ C NMR (125 MHz, DMSO-d $_{6}$ ,  $\delta$  ppm): 164.0 (>C=); 161.19 và

159.26 (>C=,  $J_{C-F}$  = 241.25 Hz); 154.0 (>C=); 133.40 và 133.38 (>C=,  $J_{C-F}$  = 2.5 Hz); 125.53 và 125.47 (>C=,  $J_{C-F}$  = 7.5 Hz); 116.33 và 116.15 (>C=,  $J_{C-F}$  = 22.5 Hz). LC-MS (m/z) [M - H]<sup>-</sup> calcd for C<sub>9</sub>H<sub>4</sub>Cl<sub>2</sub>FN<sub>4</sub> 256.9803, found 257.0161.

- 4.2.4 4,6-Dichloro-*N*-(4-methoxyphenyl)-1,3,5-triazin-2-amine (1c). White solid, yield 96%, mp 150–151 °C. IR (v cm<sup>-1</sup>): 3247 (NH), 1389 (C=N), 2985 (ArCH), 802 (C-N, s-triazine), 790 (Ar–Cl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 10.97 (1H, s, NH), 7.42 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 6.95 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 3.10 (3H, s, -OCH<sub>3</sub>). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 157.8, 153.7, 127.8, 125.3, 123.6, 114.6, 114.2, 55.5. LC-MS (m/z) [M H]<sup>-</sup> calcd for C<sub>10</sub>H<sub>7</sub>Cl<sub>2</sub>N<sub>4</sub>O 269.0002, found 268.9938.
- 4.2.5 4,6-Dichloro-*N*-(p-tolyl)-1,3,5-triazin-2-amine (1d). White solid, yield 95%, mp 156–157 °C. IR (v cm<sup>-1</sup>): 3379 (NH), 1541 (C=N), 1013 (C-N, s-triazine), 791 (Ar–Cl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 11.03 (1H, s, NH), 7.46 (2H, d, J = 8.5 Hz, H<sub>Ar</sub>), 7.17 (2H, d, J = 8.5 Hz, H<sub>Ar</sub>), 2.27 (3H, s, –CH<sub>3</sub>). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 163.8, 153.8, 134.4, 130.4, 129.7, 121.8, 20.6. LC-MS (m/z) [M H]<sup>-</sup> calcd for C<sub>10</sub>H<sub>7</sub>Cl<sub>2</sub>N<sub>4</sub> 253.0053, found 253.0039.
- **4.2.6 4,6-Dichloro-N-(4-nitrophenyl)-1,3,5-triazin-2-amine (1e).** White solid, yield 91%, mp 350–352 °C. IR (v cm<sup>-1</sup>): 3389 (NH), 1497 (C=N), 1315 (N=O), 1113 (C-N, *s*-triazine), 800 (Ar-Cl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 10.89 (1H, s, NH), 8.25 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 7.91 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 160.7, 154.2, 144.2, 142.5, 125.0, 119.8. LC-MS (m/z) [M H]<sup>-</sup> calcd for C<sub>9</sub>H<sub>4</sub>Cl<sub>2</sub>N<sub>5</sub>O<sub>2</sub> 283.9748, found 283.9722.
- 4.2.7 General procedure for the preparation of 4-substituted phenylamino-s-triazine derivatives (2a-2e and 3a-3e)
- 4.2.7.1 The reflux method. A mixture of mono-substituted striazine derivatives **1a-e** (5 mmol), saturated cyclic amine (15 mmol), and potassium carbonate (10 mmol) in 1,4-dioxane (30 mL) was refluxed until the reaction was complete (10–12 h). Then, 1,4-dioxane was removed by a Heidolph rotary evaporator. The crude solid was purified by recrystallization with ethanol: water (2:8, v/v) solvent or column chromatography on silica gel using hexane/ethyl acetate as eluent. Reaction yields range from 80 to 88%.
- 4.2.7.2 The microwave-assisted method. A mixture of monosubstituted s-triazine derivatives 1a-e (5 mmol), saturated cyclic amine (15 mmol), and potassium carbonate (10 mmol) in 1,4-dioxane (10 mL) was placed in a microwave synthesizer and irradiated at a fixed power of 300 W at 107 °C until the reaction was complete (15–30 min). Then, 1,4-dioxane was removed by a Heidolph rotary evaporator. The crude solid was purified by recrystallization with ethanol:water (2:8, v/v) solvent or column chromatography on silica gel using hexane/ethyl acetate as eluent. Reaction yields range from 91 to 98%.
- 4.2.7.3 Solubility. The synthesized compounds are well soluble in DMSO, 1,4-dioxane, and methanol; moderately soluble in ethanol; poorly soluble in non-polar solvents (hexane and ethyl acetate) and water.
- 4.2.7.4 Stability. The tri-substituted s-triazine derivatives are stable at room temperature and can be stored long-term at low temperatures of 4–8  $^{\circ}$ C.

**4.2.8** *N*-(4-Chlorophenyl)-4,6-di(piperidin-1-yl)-1,3,5-triazin-2-amine (2a). White solid, mp 180–181 °C. IR (v cm<sup>-1</sup>): 2900 (NH), 1485 (C=N), 1240 (C-N, s-triazine), 804 (Ar–Cl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 8.96 (1H, s, NH), 7.69 (2H, d, J=8.5 Hz, H<sub>Ar</sub>), 7.07 (2H, d, J=8.5 Hz, H<sub>Ar</sub>), 3.69 (8H, t, J=4.5 Hz, –CH<sub>2</sub>–), 1.59 (4H, t, J=4.5 Hz, –CH<sub>2</sub>–), 1.48 (8H, s, –CH<sub>2</sub>–). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 164.39, 164.35, 164.01, 163.99, 157.8, 155.9, 139.6, 136.96, 136.94, 128.0, 124.6, 120.8, 120.7, 120.6, 114.7, 114.5, 43.5, 25.3, 24.3. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>19</sub>H<sub>26</sub>ClN<sub>6</sub> 373.1902, found 373.1898.

4.2.9 *N*-(4-Chlorophenyl)-4,6-dimorpholino-1,3,5-triazin-2-amine (3a). White solid, mp 192–193 °C. IR (v cm<sup>-1</sup>): 3350 (NH), 1539 (C=N), 1109 (C-N, *s*-triazine), 802 (Ar-Cl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 8.82 (1H, s, NH), 7.65 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 7.01 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 3.70 (8H, t, J = 5.5 Hz, -CH<sub>2</sub>-), 3.63 (8H, t, J = 5.0 Hz, -CH<sub>2</sub>-). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 164.7, 163.8, 158.0, 156.1, 136.3, 121.2, 121.1, 114.4, 114.2, 65.6, 43.2. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>17</sub>H<sub>22</sub>ClN<sub>6</sub>O<sub>2</sub> 377.1487, found 377.1477.

**4.2.10** *N*-(4-Fluorophenyl)-4,6-di(piperidin-1-yl)-1,3,5-triazin-2-amine (2b). White solid, mp 148–150 °C. IR (v cm<sup>-1</sup>): 2936 (NH), 1479 (C=N), 1207 (Ar–F), 1107 (C–N, *s*-triazine). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 8.66 (1H, s, –NH–), 7.68 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 7.04 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 3.70 (8H, t, J = 5.5 Hz, –CH<sub>2</sub>–), 1.62 (4H, t, J = 5.0 Hz, –CH<sub>2</sub>–), 1.52 (8H, t, J = 4.0 Hz, –CH<sub>2</sub>–). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 164.4, 164.0, 157.86 and 155.98 ( $J_{\rm C-F} = 235$  Hz), 136.99 and 136.98 ( $J_{\rm C-F} = 1.25$  Hz), 120.78 and 120.72 ( $J_{\rm C-F} = 7.5$  Hz), 114.78 and 114.61 ( $J_{\rm C-F} = 21.25$  Hz), 43.6, 25.3, 24.3. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>19</sub>H<sub>26</sub>FN<sub>6</sub> 357.2197, found 357.2124.

**4.2.11** *N*-(4-Fluorophenyl)-4,6-dimorpholino-1,3,5-triazin-2-amine (3b). White solid, mp 208–209 °C. IR (v cm $^{-1}$ ): 3300 (NH), 1479 (C=N), 1253 (Ar-F), 1109 (C-N, s-triazine).  $^{1}$ H NMR (500 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 9.11 (1H, s, -NH-), 7.66 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 7.08 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 3.68 (8H, t, J = 5.0 Hz, -CH<sub>2</sub>-), 3.61 (8H, t, J = 5.0 Hz, -CH<sub>2</sub>-).  $^{13}$ C NMR (125 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 164.7, 163.9, 158.09 and 156.20 (J<sub>C-F</sub> = 236.25 Hz), 136.6, 121.19 and 121.13 (J<sub>C-F</sub> = 7.5 Hz), 114.90 and 114.73 (J<sub>C-F</sub> = 21.25 Hz), 66.0, 43.3. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>17</sub>H<sub>22</sub>FN<sub>6</sub>O<sub>2</sub> 361.1783, found 361.1778.

4.2.12 *N*-(4-Methoxyphenyl)-4,6-di(piperidin-1-yl)-1,3,5-triazin-2-amine (2c). White solid, mp 204–205 °C. IR (v cm<sup>-1</sup>): 3360 (NH), 1506 (C=N), 1033 (C-N, *s*-triazine). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 8.72 (1H, s, NH), 7.57 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 6.83 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 3.70 (3H, s, -OCH<sub>3</sub>), 3.68 (8H, t, J = 5.0 Hz, -CH<sub>2</sub>-), 1.59 (4H, t, J = 4.5 Hz, -CH<sub>2</sub>-), 1.48 (8H, d, J = 3.5 Hz, -CH<sub>2</sub>). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 164.5, 164.0, 154.0, 133.8, 120.8, 113.5, 55.1, 43.5, 25.4, 24.4. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>20</sub>H<sub>29</sub>N<sub>6</sub>O 369.2397, found 369.2299.

**4.2.13** *N*-(4-Methoxyphenyl)-4,6-dimorpholino-1,3,5-triazin-2-amine (3c). White solid, mp 210–211 °C. IR (v cm<sup>-1</sup>): 3381 (NH), 1504 (C=N), 1109 (C-N, *s*-triazine), 800 (Ar–Cl). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 8.90 (1H, s, NH), 7.55 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 6.84 (2H, d, J = 9.0 Hz, H<sub>Ar</sub>), 3.70 (3H, s, –OCH<sub>3</sub>), 3.68 (8H, t, J = 5.0 Hz, –CH<sub>2</sub>–), 3.61 (8H, t, J = 4.5 Hz, –CH<sub>2</sub>–). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 164.7, 163.9, 154.2, 133.3,

121.1, 113.6, 66.3, 66.0, 55.1, 43.3. LC-MS (m/z)  $[M + H]^+$  calcd for  $C_{18}H_{25}N_6O_3$  373.1983, found 373.1981.

**4.2.14 4,6-Di(piperidin-1-yl)-***N***-(p-tolyl)-1,3,5-triazin-2-amine (2d).** White solid, mp 153–155 °C. IR (v cm<sup>-1</sup>): 3350 (NH), 1483 (C=N), 1101 (C-N, *s*-triazine). <sup>1</sup>H NMR (500 MHz, DMSOde,  $\delta$  ppm): 8.48 (1H, s, NH), 7.55 (2H, d, J = 8.5 Hz, H<sub>Ar</sub>), 7.04 (2H, d, J = 8.0 Hz, H<sub>Ar</sub>), 3.70 (8H, t, J = 5.5 Hz, -CH<sub>2</sub>-), 2.24 (3H, s, -CH<sub>3</sub>), 1.63 (4H, t, J = 5.5 Hz, -CH<sub>2</sub>-), 1.52 (8H, t, J = 5.5 Hz, -CH<sub>2</sub>-). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 164.4, 163.9, 137.7, 129.6, 128.2, 119.3, 43.3, 24.9, 24.0, 19.8. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>20</sub>H<sub>29</sub>N<sub>6</sub> 353.2448, found 353.2369.

**4.2.15 4,6-Dimorpholino-***N***-(p-tolyl)-1,3,5-triazin-2-amine (3d)**. White solid, mp 212–213 °C. IR (v cm<sup>-1</sup>): 3360 (NH), 1504 (C=N), 1109 (C-N, *s*-triazine). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 8.95 (1H, s, NH), 7.53 (2H, d, J = 8.5 Hz, H<sub>Ar</sub>), 7.05 (2H, d, J = 8.0 Hz, H<sub>Ar</sub>), 3.69 (8H, t, J = 5.0 Hz, -CH<sub>2</sub>-), 3.61 (8H, t, J = 4.5 Hz, -CH<sub>2</sub>-), 2.23 (3H, s, -CH<sub>3</sub>). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 164.7, 163.9, 137.6, 130.2, 128.7, 119.7, 66.0, 43.3, 20.3. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>18</sub>H<sub>25</sub>N<sub>6</sub>O<sub>2</sub> 357.2304, found 357.2304.

**4.2.16** *N*-(**4-Nitrophenyl**)-**4,6-di(piperidin-1-yl**)-**1,3,5-triazin-2-amine** (**2e**). Yellow solid, mp 236–238 °C. IR (v cm<sup>-1</sup>): 3335 (NH), 1485 (C=N), 1109 (C-N, s-triazine). <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 9.46 (1H, s, NH), 8.13 (2H, d, J = 9.5 Hz, H<sub>Ar</sub>), 7.95 (2H, d, J = 9.5 Hz, H<sub>Ar</sub>), 3.73 (8H, t, J = 5.0 Hz, -CH<sub>2</sub>-), 1.65 (4H, t, J = 5.0 Hz, -CH<sub>2</sub>-), 1.56–1.51 (8H, m, -CH<sub>2</sub>-). <sup>13</sup>C NMR (125 MHz, DMSO-d<sub>6</sub>,  $\delta$  ppm): 164.2, 163.7, 147.0, 140.2, 124.1, 118.0, 43.5, 24.9, 23.8. LC-MS (m/z) [M + H]<sup>+</sup> calcd for C<sub>19</sub>H<sub>26</sub>N<sub>7</sub>O<sub>2</sub> 384.2142, found 384.2064.

4.2.17 4,6-Dimorpholino-*N*-(4-nitrophenyl)-1,3,5-triazin-2-amine (3e). Yellow solid, mp 273–275 °C. IR (v cm $^{-1}$ ): 3350 (NH), 1491 (C=N), 1109 (C-N, *s*-triazine).  $^{1}$ H NMR (500 MHz, DMSO-d<sub>6</sub>, δ ppm): 9.85 (1H, s, NH), 8.16 (2H, d, J=9.5 Hz, H<sub>Ar</sub>), 7.93 (2H, d, J=9.5 Hz, H<sub>Ar</sub>), 3.71 (8H, t, J=4.5 Hz, -CH<sub>2</sub>-), 3.64 (8H, t, J=4.5 Hz, -CH<sub>2</sub>-).  $^{13}$ C NMR (125 MHz, DMSO-d<sub>6</sub>, δ ppm): 164.6, 163.9, 147.1, 140.4, 124.8, 118.5, 65.9, 43.4. LC-MS (m/z) [M + H] $^{+}$  calcd for C<sub>17</sub>H<sub>20</sub>N<sub>7</sub>O<sub>4</sub> 386.1582, found 386.1509.

#### 4.3. In vitro anticancer activity

The cytotoxic activity of the s-triazine derivatives was evaluated against the two cancer cell lines (MCF7 - ATCC HTB-22 and C26 - ATCC CRL-2638) and one normal cell line (BAEC - CSC 2B2, Cell Systems) using the methyl thiazolyl tetrazolium (MTT) method conducted according to the MTT assay protocol. PTX and DOX were used as the positive control. The assay detects the reduction of yellow tetrazolium by metabolically active cells to purple formazan, which is measured using spectrophotometry. The cell lines were seeded into 96-well plates at a density of 5000 cells per well and incubated at 37 °C in 5% CO<sub>2</sub> for 24 h with growth media consisting of 2 mM L-glutamine, 100 IU mL<sup>-1</sup> penicillin, 100 μg mL<sup>-1</sup> streptomycin, Eagle's minimum essential medium, and 10% fetal calf serum. After that, a series of concentrations of the tested compounds and the reference drugs (PTX and DOX) in DMSO was added to each well of the plate and incubated for 24 h. The 10 µL fresh solution of MTT reagent was added to each well, and the plate was incubated in

Table 5 Targets for in silico molecular docking studies

Entry	Target	Symbol	PDB ID	Organism	Expression system	Ref drug
1	Dihydrofolate reductase	DHFR	1RG7	Escherichia coli	Escherichia coli	Methotrexate
2	Vascular endothelial growth factor receptor 2	VEGFR2	4AG8	Homo sapiens	Spodoptera frugiperda	Axitinib
3	Phosphoinositide 3-kinase	PI3K	5JHB	Homo sapiens	Spodoptera frugiperda	Gedatolisib
4	Cyclin-dependent kinase 2	CDK2	2 A4L	Homo sapiens	Spodoptera frugiperda	Seliciclib
5	Epidermal growth factor receptor	EGFR	1M17	Homo sapiens	Spodoptera frugiperda	Erlotinib
6	Mammalian target of rapamycin	mTOR	4JSV	Homo sapiens	Homo sapiens	Gedatolisib

a CO<sub>2</sub> incubator at 37 °C for 4 h until a purple precipitate appeared. Finally, the cells were solubilized in ethanol and their optical density was recorded at 570 nm.<sup>5,7</sup> The percent of proliferation inhibition was calculated using the following formula:

Viability cells inhibition (%) = 
$$100 - \left[\frac{(A_t - A_b)}{(A_c - A_b)}\right] \times 100\%$$

 $A_t$  = absorption of test compound,  $A_b$  = absorption of blank,  $A_c$  = absorption of control.

#### 4.4. ADME-Tox predictions

The physicochemical and *in silico* ADMET properties were performed using ADMETlab 3.0 descriptors algorithm protocol.<sup>58</sup>

#### 4.5. In silico molecular docking studies

The structure of ligands and reference drugs were drawn in ChemBioDraw Ultra 19.0 and the energy of each ligand was minimized using ChemBio3D Ultra 19.0.5-9 The ligands with minimized energy were then used as input for AutoDock Vina, in order to carry out the docking simulation. Anticancer targets of DHFR, VEGFR2, PI3K, CDK2, EGFR, and mTOR were retrieved from the protein data bank (Table 5). The targets were removed all the water molecules and added only polar hydrogen and Kollman charges. The AutoDock Tools was used to set the grid box for docking simulations. In addition, the docking protocol was validated by extracting and re-docking the cocrystallized ligand into the active site. Then, synthesized compounds and reference drugs were docked with the target to determine the docking parameters with the help of grid-based ligand docking (Table 6). Finally, the interaction information (bond type, length, amino acid residue, etc.) and the pictorial representation of the interaction between the ligands and the target were processed using Discovery Studio 2021 software.

Table 6 Grid box parameters for anticancer targets

	Size			Center					
Target	x y		z	x	У	z			
DHFR	40	40	40	-1.65679	22.0285	23.0763			
VEGFR2	40	40	40	20.8237	25.5351	39.4596			
PI3K	40	40	40	21.3492	-4.07429	20.8696			
CDK2	40	40	40	100.865	101.747	79.8926			
EGFR	40	40	40	22.0137	0.252828	52.794			
mTOR	40	40	40	50.1244	-1.7953	-45.9034			

#### 4.6. Statistical analysis

All values are expressed in Mean  $\pm$  SEM (Standard Error of Mean). The difference in IC<sub>50</sub> value between tested compounds and reference drug was analyzed by one-way analysis of variance (ANOVA) with Tukey's Honestly Significant Difference (Tukey HSD) post hoc test using SPSS 26 software. The results were considered statistically significant if p < 0.05. The chart is drawn using Microsoft Excel software.

## Data availability statements

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

## Author contributions

Em Canh Pham: conceptualization, methodology, investigation, data curation, supervision, writing-original draft preparation, writing – review & editing. Bich-Ngoc Thi Le: investigation. Anh Minh Ngo: investigation. Long Binh Vong: investigation, supervision. Tuyen Ngoc Truong: data curation, supervision, writing-original draft preparation, writing – review & editing.

## Conflicts of interest

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

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