


 Cite this: *RSC Adv.*, 2023, **13**, 3056

Design, synthesis, anticancer and *in silico* assessment of 8-caffeinyl-triazolymethoxy hybrid conjugates†

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In this research the synthesis, characterization, anticancer and the cytotoxicity assessments of novel 8-caffeinyl-triazolymethoxy hybrid conjugates have been described. These compounds are the first caffeine-1,2,3-triazolyl hybrid molecules that structurally are composed of three compartments comprising caffeinyl, 1,2,3-triazolyl and *N*-alkyl/aryl residues. The *in vitro* evaluations of synthesized compounds on cancer cell lines, including two breast cancer cell lines MDA-MB-468 (ATCC HTB-22), MCF-7 (ATCC HTB-22), melanoma cell line A-375 (ATCC CRL-1619) and normal cell line HEK-293 (ATCC CRL-11268) have determined that **22c** (IC₅₀ < 12.5 μM) demonstrated potent activity against A375 and its toxicity is even stronger than methotrexate (MTX) as a standard drug. Additionally, **22c** involves more selectivity than MTX regarding its non-toxicity for the HEK-293 cell line. Among the tested compounds against two breast cancer cell lines, **22f** (IC₅₀ = 136 ± 0.2 and 126 ± 0.6 μM for MCF-7 and MDA-MB-468, respectively) and **22i** (IC₅₀ = 165 ± 1.8 and 175 ± 1.4 μM for MCF-7 and MDA-MB-468, respectively) were the most potent compounds but their activities were less than MTX, moreover **22f** showed more selectivity regarding its lower toxicity against HEK-293. Overall, **22f** displayed general toxicity and selectivity on all tested cancer cell lines. The *in silico* physicochemical properties, pharmacokinetic profile, and drug likeness predictions were also carried out for all the studied compounds. Most new compounds exhibited zero violation of Lipinski's rule (RO5). A molecular docking study was also conducted to predict the binding mode and the interaction of **22c** as the most active anti-melanoma entry with B-RAF V600E kinase enzyme. The docking results determined that **22c** exhibited a strong binding affinity to the active site of the enzyme. These findings demonstrated **22c** and **22f** as potential future anticancer drug candidates.

 Received 2nd December 2022
 Accepted 12th January 2023

DOI: 10.1039/d2ra07683g

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1 Introduction

Perhaps, after infectious and cardiac diseases, cancer has been one of the most deadly diseases that has occurred throughout the history of human life. Cancer is a public health problem all over the world. This disease affects all people, young and old, rich and poor, men, women and children. Cancer is the uncontrollable growth and expansion of cells that can infect almost all tissues of the body. More than eleven million people are diagnosed with cancer every year. It is estimated that there will be 16 million new cases of cancer annually before 2020. The

treatment of cancer causes a high cost to governments and peoples, for example, the total cost of cancer was €199 billion in Europe (EU-27 plus Iceland, Norway, Switzerland, and the United Kingdom) in 2018.¹ The high expenditure of cancer treatment is due to various reasons; however, the most important reason for this high cost is the price of anti-cancer drugs. Despite the good advances in the discovery and use of anti-cancer drugs in recent years; nevertheless, these drugs have several shortcomings, such as low drug efficacy, high toxicity, lack of selectivity, high prices and so on. Therefore, there is still a constant need to design and generate new, cheap and effective anti-cancer drugs with specialized performance.²

It is not overstating that xanthine alkaloids are the most prominent naturally occurring alkaloids recognized throughout human life.³ Natural xanthines are purine-based scaffolds which possess widespread medicinal properties that are extensively exploited in the remedy of diverse diseases.⁴ Amongst xanthine alkaloids, methyl xanthines comprising caffeine, theophylline, theobromine and paraxanthine are amazing since these xanthine scaffolds are well-recognized by diverse

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d2ra07683g>



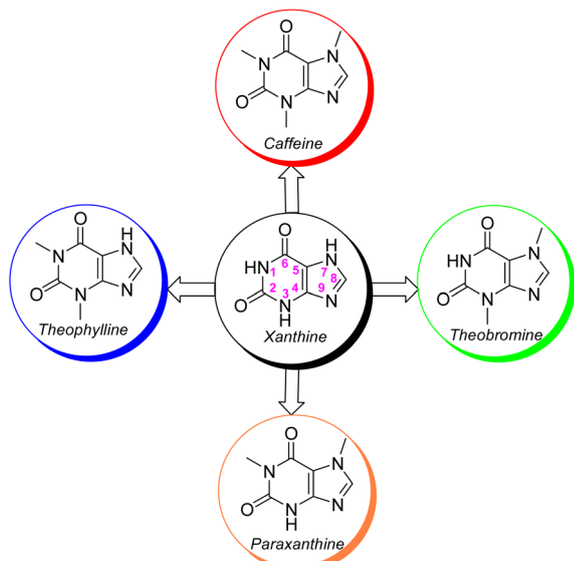


Fig. 1 The structures of xanthine and its natural methylated derivatives.

receptors or enzymes in animal cells that caused their magnificent clinical and therapeutic activity (Fig. 1).⁵ They are abundantly found in miscellaneous seeds, fruits, and leaves of a variety of plants, like tea, cacao beans, guarana berries and yerba matte leaves.^{6,7} However, caffeine is more pronounced than the other methylxanthines owing to its widespread popularity and extreme social consumption as tea, coffee and chocolate. Up to now, the parental caffeine and its derivatives are known as super nootropic molecules with remarkable clinical activity like anti-Alzheimer, anti-diabetes, anti-Parkinson, anti-cancer, anti-inflammatory, antitussives and stimulating agents for the central nervous system (CNS).⁸

The xanthine as a starting material encompasses multiple active nucleophilic sites (N1, N3, N7 and N9 sites) for the enormous diversification reaction through the nucleophilic substitution. Nevertheless, methyl xanthines like theophylline, theobromine and paraxanthine can merely contribute to the nucleophilic substitution by N7, N1 and N3 atoms, respectively. Conversely, the nucleophilic sites in caffeine are masked by methylated nitrogens and hence the caffeine is unable to undertake the substitution reaction through its masked nitrogens. Indeed, the caffeine can be chemically altered merely *via* its C8 site with rare electrophiles or active species.^{9–12} It is worth mentioning that the alteration of methylxanthines at all available sites has emerged the impressive and brilliant pharmacological profiles reviewed in many literature.^{13,14} In particular, C8-modification of caffeine derivatives extensively improves the biological activities designated in Fig. 2. The most C8-modified caffeine derivatives can be acquired by two main strategies. First, the halogenation (*i.e.*: bromination and chlorination) followed by S_NAr -type reaction *via* diverse nucleophiles. Second, the coupling of 1,3-dimethyl-5,6-diaminouracil with various carboxylic acids. In general, the reported caffeine's C8-modified adducts can be categorized on the basis of atom's type linked to C8. In this regard, they can be sorted into C8-C, C8-O, C8-S, C8-N caffeine derivatives. The structures of a few examples of C8-C, C8-O, C8-S, C8-N derivatives beside their corresponding biological profiles are summarized in Fig. 3. The C8-C derivatives were mainly prepared through the reaction of 1,3-dimethyl-5,6-diaminouracil with carboxylic acids bearing diverse functionalities.^{15–20} General structures 1–3 and 5 (Fig. 3) were prepared by this method, however for 3 with $n = 0$ (*i.e.* 8-aryl caffeine), the aryl halide and 8-caffeinyly cobalt(I) complex was used.¹¹ Moreover, C8-C derivatives 4 can be obtained by direct reaction of caffeine, alcohol and *t*-butyl hydroperoxide (TBHP) under microwave irradiation.¹⁰ Regarding compounds 6–10 (C8-S),^{12,21–28} 11–19 (C8-N),^{11,29–34} 20 and 21 (C8-O)^{35–37}

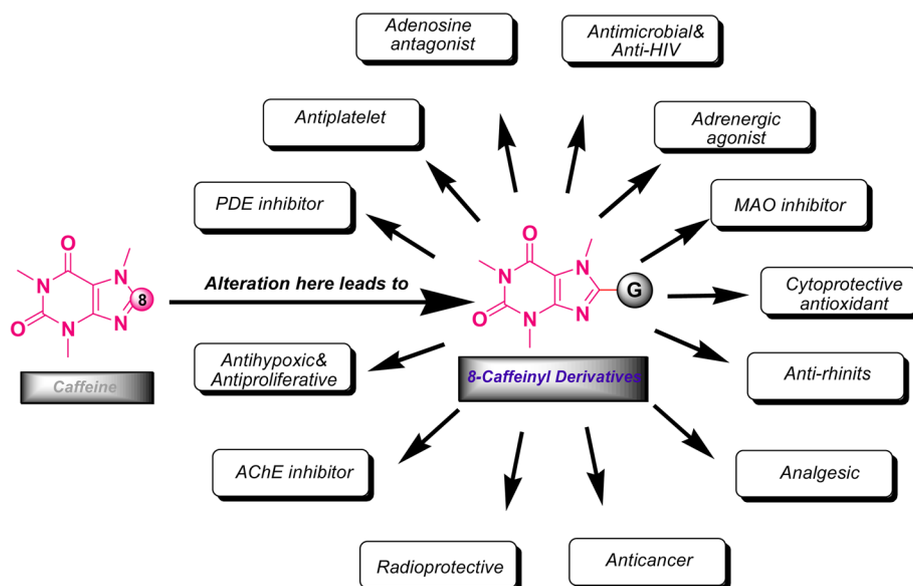


Fig. 2 The resulting biological properties by the C8 modification of caffeine.



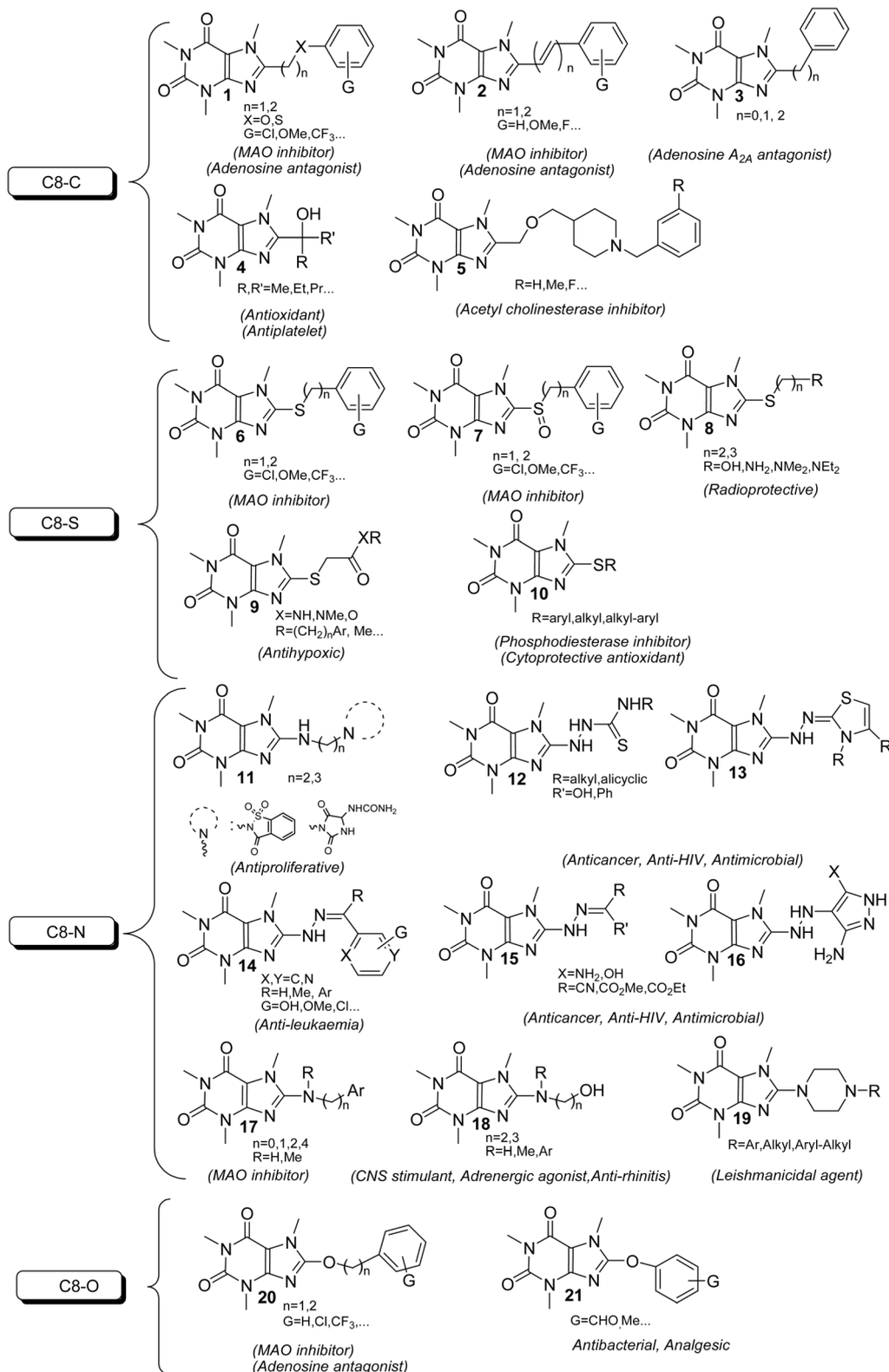


Fig. 3 Classification of the C8-modified caffeine derivatives and their corresponding biological activity.

(Fig. 3), they were often prepared by the S_NAr -type reaction of 8-bromocaffeine (8-BC) or 8-chlorocaffeine (8-CC) with the corresponding nucleophiles in a single or multistep processes.

Undoubtedly, the 1*H*-1,2,3-triazolyl cores are unique and significant scaffolds in medicinal chemistry. Owing to the incorporation of the 1*H*-1,2,3-triazolyl core into the structure of



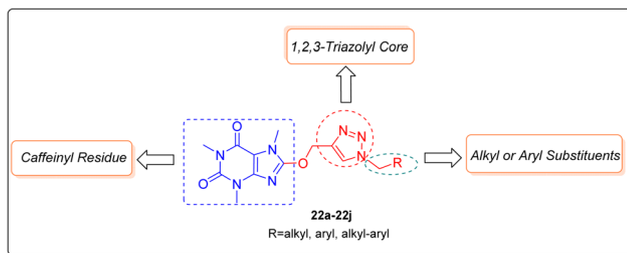


Fig. 4 General structure of novel 8-caffeinylyl-triazolylmethoxy hybrid conjugates.

a bioactive molecule, it extensively increases its pharmacophoric strength by establishing a variety of chemical interactions.^{38,39} Additionally, the 1,2,3-triazolyl core is known as a main non-hydrolysable bioisostere of the amide bond regarding to its significant electronic and topological similarities.⁴⁰ In this connection, the 'Click' Huisgen azide-alkyne cycloaddition is a reliable and efficient tool for incorporating the 1*H*-1,2,3-triazolyl cores into the drugs scaffold and also the conjugation of two organic molecules to acquire a hybrid molecule.⁴¹

As mentioned earlier and shown in Fig. 3, the C8-modification of caffeine with diverse organic moieties extensively improves its biological activity, however to the best of our knowledge no molecule comprising 1*H*-1,2,3-triazolyl core was considered for conjugating with caffeinylyl moiety. In this regard, hereby we would like to report the synthesis, characterization, anticancer and *in silico* assessment of novel 8-caffeinylyl-triazolylmethoxy hybrid conjugates **22a–22j** (Fig. 4).

2 Results and discussion

2.1. Chemistry

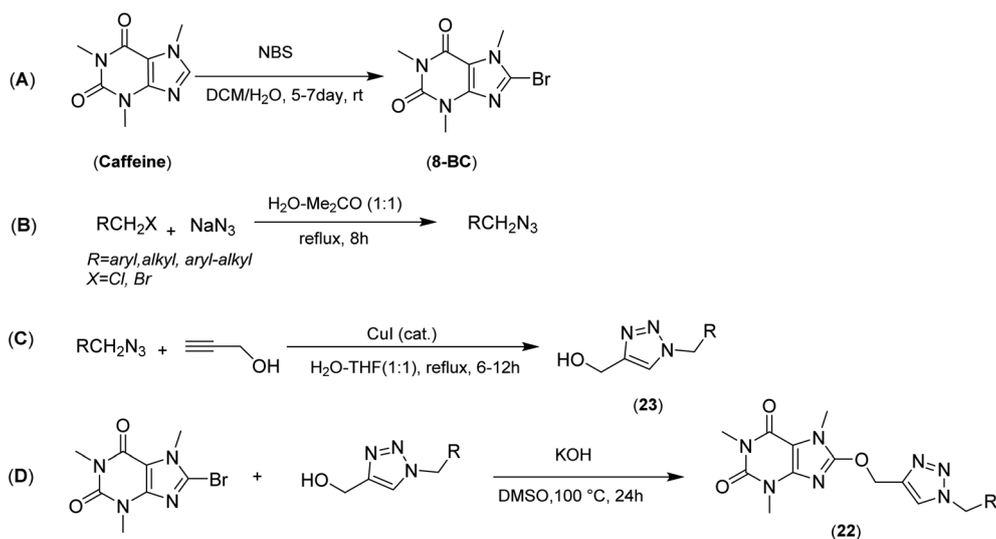
The synthesis of 8-caffeinylyl-triazolylmethoxy hybrid conjugates **22** was carried out due to Scheme 1A–D. As indicated in Scheme 1A–D, this synthesis was started by the bromination of caffeine to

increase the positive charge density on C(8) for the subsequent S_NAr -type reaction (Scheme 1A). The bromination of caffeine was carried out due to our previously reported procedure using NBS as a brominating reagent in a mixture of DCM and water at room temperature.²⁷ Also the bromination of caffeine can be conducted using Br_2/H_2O_2 in glacial HOAc²⁵ or HBr/H_2O_2 ;²⁴ however, none of these two current procedures were efficient for synthesis of the 8-BC. Applying the NBS method almost yielded the pure 8-BC, quantitatively. In another reaction the require alkyl azides were readily synthesized using alkyl chlorides or bromides and excess amounts of sodium azide in a mixture of acetone–water at reflux to yield the corresponding alkyl azides almost, quantitatively (Scheme 1B).⁴² Subsequently, to afford the 8-proparyloxy caffeine as a require alkyne, we attempted to tether the propargyl moiety to caffeinylyl residue through the reaction of 8-BC and propargyl alcohol using KOH as a base in DMSO at 100 °C; however, the obtained yield was not better than 25%. The use of other organic solvents, different bases and catalysts had no considerable effect on improving the obtained yield. Thus, **23** was prepared through the 'Click' Huisgen alkyl-azide cycloaddition using the propargyl alcohol and desired alkyl azides in the presence of cuprous catalyst and in a mixture of THF–H₂O at reflux condition for 6–12 h (Scheme 1C). Ultimately, **23** was coupled with 8-BC through S_NAr -type reaction using KOH in DMSO at 100 °C to afford the title products in good yields (Scheme 1D).

The structures of all target molecules as well as corresponding yields and melting points are demonstrated in Fig. 5. As shown in Fig. 5, compounds are common in caffeinylyl and triazolylmethoxy parts but they are different in alkyl residues. Compounds bearing alkyl–aryl (**22a–22e**) and diverse aliphatic residues having differences in carbon chain length (**22f–22j**) were synthesized.

2.2. *In vitro* anticancer and cytotoxic activity

The World Health Organization (WHO) reported Melanoma as the most aggressive form of skin cancer that is increasing



Scheme 1 General synthetic pathway (A–D) for preparing 8-caffeinylyl-triazolylmethoxy hybrid conjugates (**22**).



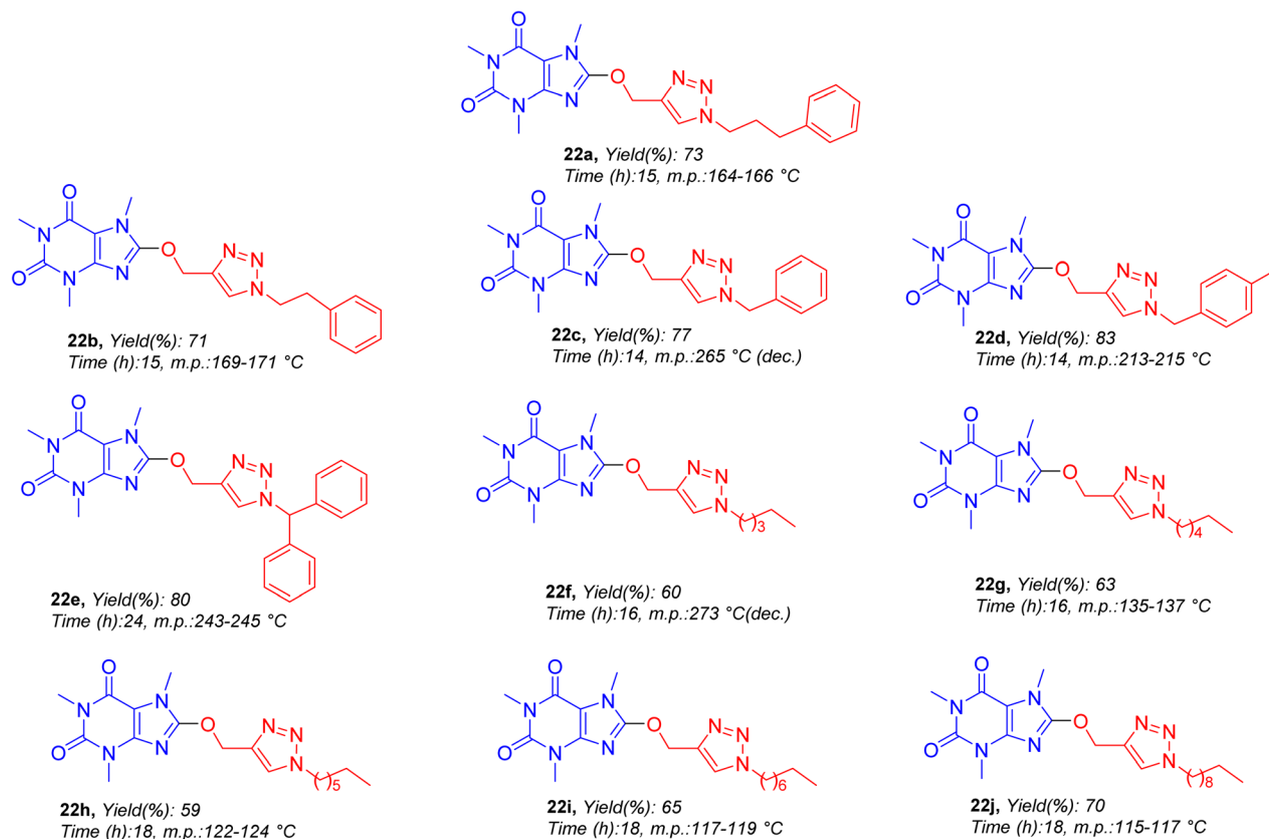


Fig. 5 The structures, isolated yields and melting points of compounds 22a–22j.

worldwide. The A375 cell line is a known human melanoma cell line derived from a metastatic melanoma.⁴³ Breast carcinoma is the most frequent malignancy among women and the second leading cause of cancer death in them. The MDA-MB-468 cell line was isolated from women patient with metastatic adenocarcinoma of the breast⁴⁴ The MCF-7 cell line was established from a pleural effusion from a 69 years-old woman at the Michigan Cancer Foundation in 1973 which is the most common model used for breast cancer. Despite its origin from the metastases of an advanced tumor, the cell line is noninvasive and represents a model of early-stage disease.⁴⁵ The human embryonic kidney 293 cells (HEK 293) are a normal cell line originally derived from human embryonic kidney cells grown in tissue culture taken from a female fetus.⁴⁶ The IC_{50} values pertaining to toxicity tests for all compounds were carried out in three replicates and the final values were reported as mean \pm SD. The final values for all samples were compared with SPSS software using one-way ANOVA with a confidence interval of 95% (Table 1). As depicted in Table 1, against normal cell line (HEK-293) all compounds showed IC_{50} values higher than 300 μ M except those of 22e, 22i, 22j and MTX. In the case of A375, 22c ($IC_{50} < 12.5 \mu$ M) demonstrated potent activity against A375 and its toxicity is even stronger than MTX as a standard drug. Regarding the ineffectiveness of 22c ($IC_{50} < 300 \mu$ M) on HEK-293 and also the remarkable toxicity of MTX ($IC_{50} = 38 \pm 1.6 \mu$ M) against HEK-293, thus 22c proved to have more selectivity

Table 1 The cytotoxic activity of 22a–22j and MTX against A-375, MCF7 and MDA-MB-468 and HEK-293 cell lines

Compound	Cell line [IC_{50} (μ M)]			
	A375	MCF7	MDA-MB-468	HEK-293
22a	179 \pm 1.2	268 \pm 0.2	256 \pm 1.4	>300
22b	>300	>300	>300	>300
22c	<12.5	>300	>300	>300
22d	194 \pm 1.6	>300	>300	>300
22e	286 \pm 0.2	284 \pm 0.2	>300	284 \pm 1.4
22f	160 \pm 1.5	136 \pm 0.2	126 \pm 0.6	>300
22g	>300	>300	>300	>300
22h	>300	>300	>300	>300
22i	146 \pm 1.5	165 \pm 1.8	175 \pm 1.4	194 \pm 1.2
22j	>300	>300	>300	194 \pm 1.2
MTX ^a	32 \pm 2.2	30 \pm 2.9	37 \pm 1.9	38 \pm 1.6

^a Methotrexate.

compared to MTX. Additionally, except those of 22b, 22g, 22h and 22j, other compounds were toxic to A375 but their toxicities were considerably less than 22c and MTX. In the case of MCF7 and MDA-MB-468, 22a, 22f and 22i displayed toxicity against both cell lines while 22e was merely effective on MCF7. Among tested compounds against breast cancer cell lines 22f and 22i were the most potent compounds, however 22f showed more selectivity regarding its less toxicity against HEK-293. In



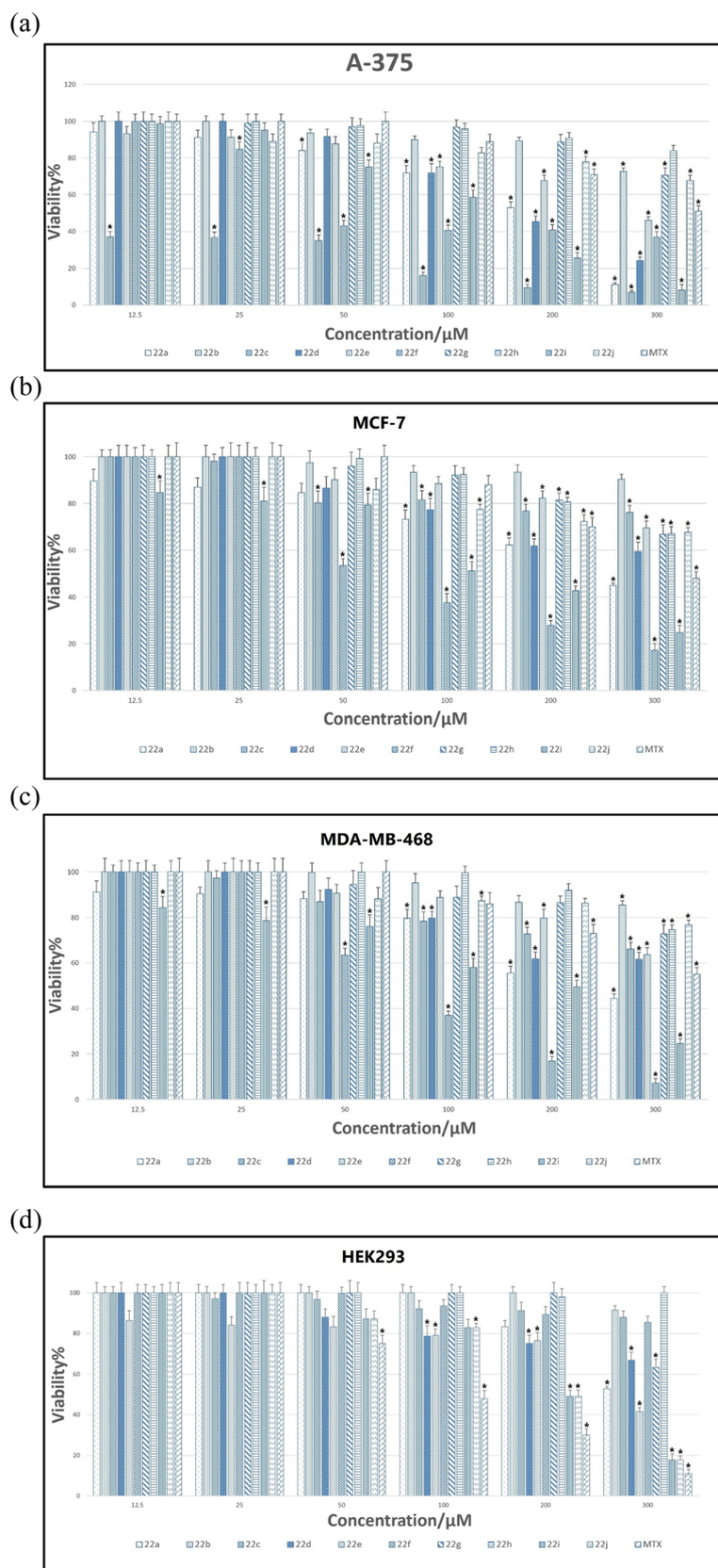


Fig. 6 (A) The viability assessment for 22a–22j and MTX against A375. (B) The viability assessment for 22a–22j and MTX against MCF7. (C) The viability assessment for 22a–22j and MTX against MDA-MB-468. (D) The viability assessment for 22a–22j and MTX against HEK-293.



general, all compounds were less toxic than MTX against both tested breast cancer cell lines. In overall, **22f** displayed general toxicity and selectivity on all tested cancer cell lines.

The viability percentages for examined cell lines *versus* different concentrations of **22a–22j** as well as MTX are shown in Fig. 6A–D. A *p*-value less than 0.05 (typically ≤ 0.05) is statistically significant that shown with stars in Fig. 6A–D.

As it is indicated in Fig. 6A, for A375, the lowest viability values and hence the highest toxicity are related to **22c** compared to all synthesized compounds. It is also worthy to mention that the viability percentage of **22c** at lowest concentration (*i.e.*: 12.5 μM) is even lower than that of MTX which endorses the higher toxicity of **22c** against A375 cancer cell line.

The viability of all compounds and MTX was also assessed for both MCF7 and MDA-MB-468 cell lines (Fig. 6B and C). As determined, among tested compounds, **22f** demonstrated lower viability at higher concentrations compared to all entries but its viability was higher than MTX at all tested concentrations. Fig. 6D indicates the viability percentage for HEK-293. As it is well indicated the lowest viability was pertained to MTX at all tested concentrations; however, for two most effective compounds **22c** and **22f**, the viability was higher than MTX, particularly at lower concentrations which indicate the selectivity of **22c** and **22f** against HEK-293.

Recently Shahzadi and coworkers have synthesized a series of theophylline-7-acetic acid (acefylline)-derived 1,2,4-triazole hybrids having *N*-phenyl acetamide moieties and tested for their inhibitory (*in vitro*) potential against two cancer cell lines, A549 (lung) and MCF-7 (breast), using MTT assay.⁴⁷ They determined that their most potent compound against MCF-7 breast cancer cell line had the viability value of 31.76 ± 3.16 at effective concentration ($100 \mu\text{g} \mu\text{L}^{-1}$), whereas current **22f** as the most potent entry against MCF-7, demonstrated the viability value of 17.0 ± 3.0 at the most effective concentration of $300 \mu\text{M}$ (*i.e.*: $0.12 \mu\text{g} \mu\text{L}^{-1}$). This result has clearly shown that **22f**

inhibited the MCF-7 cells at a much lower concentration than the most potent compound reported by Shahzadi *et al.*

Methotrexate (MTX, amethopterin) as well as aminopterin (4-aminopteronic acid) are known as 4-amino derivative of folic acid which are different only in a methyl moiety. In fact, these two compounds, which have a very close structure to folic acid, inhibit the dihydrofolate reductase (DHFR) enzyme in a competitive interaction with folic acid.⁴⁸ This enzyme plays a key role in the reduction of dihydrofolate to tetrahydrofolate (coenzyme F), and its inhibition prevents the biosynthesis of DNA and RNA and ultimately the protein.⁴⁹ Like folic acid, MTX and aminopterin are composing of three parts comprising pteridine, *p*-aminobenzoic acid (PABA) and glutamic acid residues. The main pharmacophoric residue in folic acid, MTX and aminopterin is pteridine core which is closely related to xanthine alkaloids and biologically is synthesized from guanine. In same circumstance to MTX and aminopterin, 8-caffeinyl-triazolylmethoxy hybrid conjugates are composed of three parts comprising (i) caffeinyl core as a methyl xanthine that structurally resembles to pteridine, (ii) 1,2,3-triazolyl core is a bioisostere of amide that can mimic the PABA-amide residue in MTX and aminopterin, and ultimately (iii) diverse alkyl or aryl-alkyl residues in 8-caffeinyl-triazolylmethoxy hybrid conjugates that can be considered as a successor for glutamic acid in MTX and aminopterin. Thus, owing to structural similarities of title compounds with MTX and aminopterin, it is proposed that 8-caffeinyl-triazolylmethoxy hybrid conjugates potentially inhibit DHFR enzyme and confine the biosynthesis of DNA (Fig. 7).

2.3. *In silico* assessments

2.3.1 *In silico* physicochemical and pharmacokinetic profile studies. Assessment of physicochemical parameters of a drug candidate based on Lipinski's rule of five (RO5),

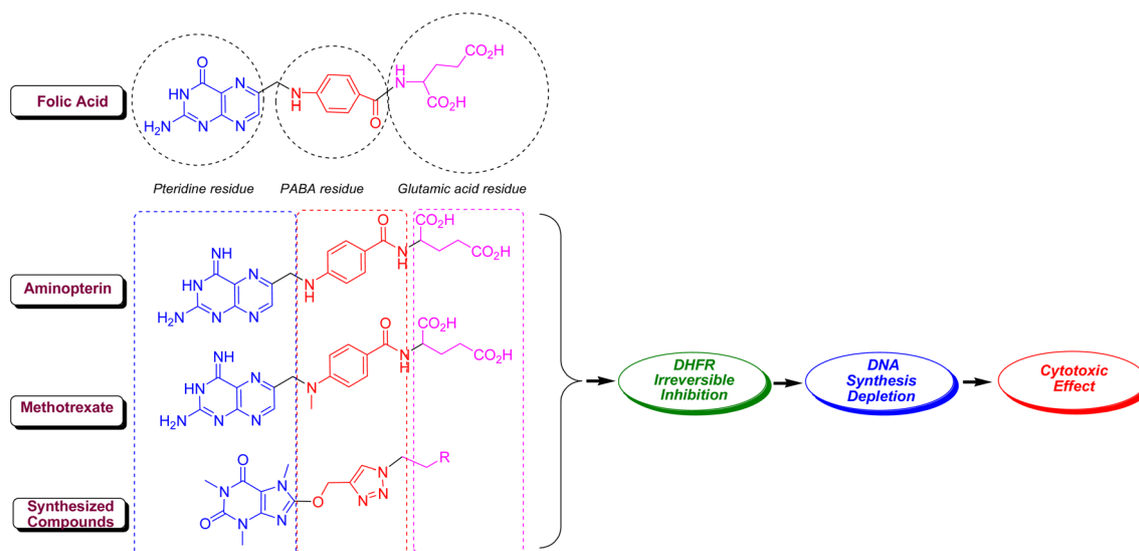


Fig. 7 The possible anticancer mechanism of 8-caffeinyl-triazolylmethoxy hybrid conjugates.



pharmacokinetics, and drug likeness is a well-known strategy for speculative evaluations of its physicochemical properties and prediction of its chance to be introduced as a drug.^{50,51} A drug candidate should conform to RO5 and require (1) molecular weight ≤ 500 Dalton, (2) rotatable bonds ≤ 10 , (3) hydrogen bond donor ≤ 5 , (4) hydrogen bond acceptor ≤ 10 , and (5) octanol-water partition coefficient ($\log P$) value ≤ 5 . The *in silico* physicochemical and pharmacokinetic profile for **22a–22j** and MTX was assessed using SwissADME online software,⁵² OSIRIS Property Explorer,⁵³ and preADMET online server.⁵⁴ The results of these studies are indicated in Table 2 and 3.

As shown in Table 2, the molecular weights of **22a–22j** are between 361.40 and 457.48 Dalton (<500 Dalton). The numbers of rotatable bonds (nRB) for all synthesized compounds except **22j** are between 5–10. The number of hydrogen bond donors ($nHBD$) and hydrogen bond acceptors ($nHBA$) for **22a–22j** are also obeyed from RO5. The lipophilicity parameter of a molecule as one of the most significant features of RO5 is characterized by $\text{clog } P$. The calculated $\text{clog } P$ for compounds **22a–22j** are between 0.66–3.19. Therefore, all title compounds conform to RO5. The drug transport character can be predicted by total polar surface area (TPSA) which is the summation of surfaces of

polar atoms in a molecule. The TPSA contents of most approved drugs are less than 140 \AA^2 . The calculated TPSA contents for all synthesized compounds are less than 140 \AA^2 which express likely appropriate oral absorption or membrane permeability. A parameter computed by OSIRIS Property Explorer software is drug likenesses which are negative for all compounds as well as MTX. Using the above calculated parameters, the software is able to give the drug score which predicts the chance of a molecule to be selected as a drug candidate. The analysis of computed data indicated that the minimum and maximum drug score values of synthesized compounds belong to **22e** and **22b**, respectively.

The pharmacokinetic profile of **22a–22j** including absorption, distribution, metabolism and toxicity properties are also depicted in Table 3. The aqueous solubility ($\log S$), human intestinal absorption (HIA), skin permeability ($\log K_p$), and P-glycoprotein (P-gp) substrate and/or inhibitor are important parameters for prediction of the absorption profile of a drug candidate. The aqueous solubility ($\log S$) of numerous approved and traded drugs is >-4 . Due to obtained results, all synthesized compounds except of **22e** and **22j** may exhibit the desirable water solubility. While the human intestinal absorption of MTX

Table 2 Physicochemical properties of **22a–22j** and MTX

Compd.	Mw. ^a	nRB ^b	$nHBD$ ^c	$nHBA$ ^d	$\text{cLog } P$ ^e	TPSA ^f	Drug likeness	Drug score
22a	409.44	7	0	6	1.45	101.76	-1.87	0.46
22b	395.42	6	0	6	1.00	101.76	-0.74	0.56
22c	381.39	5	0	6	0.66	101.76	-1.12	0.54
22d	395.42	5	0	6	1.00	101.76	-2.32	0.46
22e	457.48	6	0	6	2.45	101.76	-1.39	0.29
22f	361.40	7	0	6	0.92	101.76	-6.46	0.44
22g	375.43	8	0	6	1.37	101.76	-10.88	0.43
22h	389.45	9	0	6	1.83	101.76	-16.02	0.42
22i	403.48	10	0	6	2.28	101.76	-18.89	0.40
22j	431.53	12	0	6	3.19	101.76	-18.89	0.35
MTX ^g	454.44	10	5	9	1.23	210.54	-7.09	0.22

^a Molecular weight. ^b Number of rotatable bonds. ^c Number of hydrogen bond donors. ^d Number of hydrogen bond acceptors. ^e Logarithm of octanol-water partition coefficient ($\log P$). ^f Topological polar surface area (\AA^2). ^g Methotrexate.

Table 3 Pharmacokinetic profile of **22a–22j** and MTX

Compd.	$\log S$	HIA ^a	$\log K_p$ ^b	P-gp ^c	BBB permeability ^d	CYP2D6	CYP3A4	Carcino. ^e	hERG inhibition ^f
22a	-3.30	99.28	-4.13	NS/I	0.16	NS/NI	S/NI	NC	Medium
22b	-3.07	99.05	-4.23	S/I	0.12	NS/NI	S/NI	NC	Medium
22c	-2.88	98.77	-4.30	S/I	0.11	NS/NI	S/NI	NC	Medium
22d	-3.18	99.05	-4.31	NS/I	0.13	NS/NI	S/NI	NC	Medium
22e	-4.40	99.37	-3.51	S/I	0.19	NS/NI	S/NI	NC	Medium
22f	-2.58	95.94	-4.43	S/I	0.12	NS/NI	S/NI	NC	Medium
22g	-2.93	96.61	-4.37	NS/I	0.13	NS/NI	S/NI	NC	Medium
22h	-3.28	97.20	-4.30	NS/I	0.15	NS/NI	S/NI	NC	Medium
22i	-3.63	97.71	-4.22	NS/I	0.19	NS/NI	S/NI	NC	Medium
22j	-4.33	98.53	-4.03	NS/I	0.14	NS/NI	WS/NI	NC	Medium
MTX ^g	-1.19	36.61	-4.63	S/NI	0.04	NS/NI	WS/I	NC	High

^a Human intestinal absorption (%). ^b skin permeability ($\log K_p$, cm h^{-1}). ^c P-glycoprotein. ^d Blood-brain barrier permeability (C. brain/C. blood). ^e Carcinogenicity (mouse). ^f Human ether-a-go-go related gene channel. ^g Methotrexate. S = substrate. WS = weak-substrate. NS = non-substrate. I = inhibitor. NI = non-inhibitor. NC = non-carcinogenic.



is low; however, all compounds may involve more than 95% human intestinal absorption which would be very ideal. Skin permeability (also known as transdermal delivery) is an important factor in drug delivery which describes the significance of skin absorption. All compounds display skin permeability similar to MTX. P-glycoprotein (P-gp) is a drug transporter which excretes its substrates from the cells. While induction of P-gp function by its substrate reduces the bioavailability of that chemical; however, its inhibition may cause the enhancement in drug bioavailability. According to the obtained data, all compounds are P-gp inhibitors. Additionally, all compounds except those of **22b**, **22c**, **22e**, **22f** and MTX are predicted to be the non-substrate of P-gp.

The blood–brain barrier (BBB) permeability indicates the distribution of a compound between the blood (as the most hydrophilic) and the brain (as the most lipophilic) tissues. All compounds possess low BBB permeability which suggests that none of them can easily cross this barrier and thus does not lead to neurotoxicity. Cytochrome P450s, especially CYP2D6 and CYP3A4, play significant roles for drug metabolism. The results have indicated that **22a–22j** are non-inhibitor of either CYP2D6 or CYP3A4. MTX acts as non-inhibitor of CYP2D6 and inhibitor of CYP3A4. In addition, all compounds are substrates of CYP3A4 which indicates that they can be metabolized in the liver. Also, the results illustrated that **22a–22j** as well as MTX cannot lead to carcinogenicity in mice. Human ether-a-go-go related gene channel (hERG) inhibitors may lead to lengthening the QT and cardiac side effects. As depicted in Table 2, a medium risk of hERG inhibition potency is predicted for **22a–22j** while MTX showed high risk of hERG inhibitory activity. Taken all together, **22c** can be considered as a promising future drug candidate.

2.3.2 Docking study. The RAS/RAF/MEK/ERK signaling pathway (MAPK pathway, also known as ERK pathway) is one of the most well-known pathways involved in tumorigenesis and cancer progression.⁵⁵ It is well established that the B-RAF kinase of the MAPK pathway is responsible for 66% of melanomas.⁵⁶ Among more than 30 mutations of B-RAF kinase, substitution of glutamic acid for valine at amino acid 600 (B-RAF V600E) is the most frequent oncogenic mutation of B-RAF. Hence, B-RAF V600E is an attractive therapeutic target for treatment of melanoma.⁵⁷ In this connection, the binding mode and interactions of **22c** as the most active compound against melanoma A375 cell line were explored in the active site of B-RAF V600E. To this end, the docking study was achieved using the crystal structure of B-RAF kinase V600E oncogenic mutant. The three-dimensional (3D) crystal structure of B-RAF V600E (PDB code: 3OG7) with 2.45 Å resolution was selected from the protein Data Bank (<http://www.rcsb.org>). In 3OG7, B-RAF V600E is complexed with PLX4032 (vemurafenib). The docking process was performed with Molegro Virtual Docker (MVD 6.0) software using its default settings.⁵⁸ The enzyme was imported to the software. Then, the water molecules were removed and the missing hydrogens were added to refine its structure. The structure of B-RAF V600E contains two homo-dimeric chains A and B. To perform the docking study, the chain B and the bound ligand (vemurafenib) to the mutated chain A were removed.

Then, the optimized structure of chain A was used for docking study. The program setting was achieved to display the amino acid residues in a 7 Å radius around the inhibitor. The structure of **22c** was optimized with the Gaussian 09 program using the DFT method at the B3LYP/6-31+G** level of theory.⁵⁹ Initially, the co-crystallized ligand (*i.e.*: vemurafenib) was redocked at the active site of enzyme to endorse the docking process. The root-mean-square deviation (RMSD) value between the docked and co-crystallized vemurafenib was found to be 1.25 Å which is adequate. Next, the affinity, orientation, interactions, and binding mode of **22c** were studied based on the validated docking process and 50 independent runs were analyzed. All interactions were visualized and evaluated on the basis of the docking results using Discovery Studio visualizer.

It is reported that vemurafenib is stabilized in the binding site through the hydrogen bonds, π – π^* and π –cation interactions as well as the hydrophobic and van der Waals interactions with the residues of amino acids present in the active site of B-RAF V600E kinase enzyme.⁶⁰ The analysis of the docking protocol revealed that there are four conventional hydrogen bonds between vemurafenib and enzyme through Gln530, Cys532, Phe595, and Gly596. The π – π^* stacking interactions were observed between Trp531 and Phe583 with a heterocyclic ring of vemurafenib. The side chain of Lys483 is involved in π –cation interaction with the ligand. Additionally, the hydrophobic and van der Waals interactions of vemurafenib with the enzyme were created by the side chains of Ile463, Ala481, Lys483, Leu505, Leu514, Trp531, Cys532, Phe583, and Phe595.⁶⁰ As like vemurafenib, **22c** is delightfully located in the same binding site and mimics the key interactions with B-RAF V600E kinase enzyme. An overlay view of conformations of co-crystallized vemurafenib, redocked vemurafenib, and **22c** at the binding site of B-RAF V600E is shown in Fig. 8.

Compound **22c** provides four hydrogen bond interactions with the active site of B-RAF V600E kinase enzyme including one carbon–hydrogen bond between Gln530 and *N*(3)-methyl of caffeine, one conventional hydrogen bond between Cys532 and the oxygen atom of **22c** which is located on the C(2) atom of

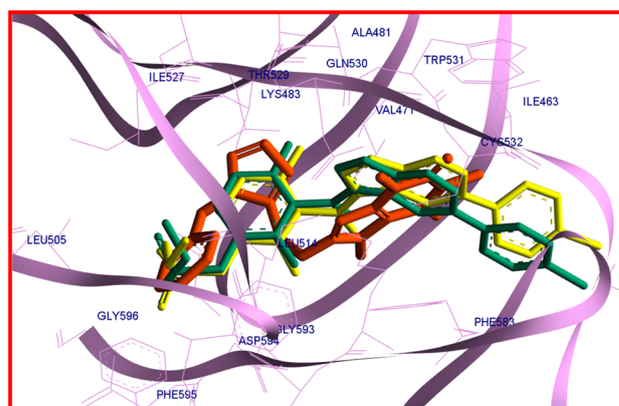
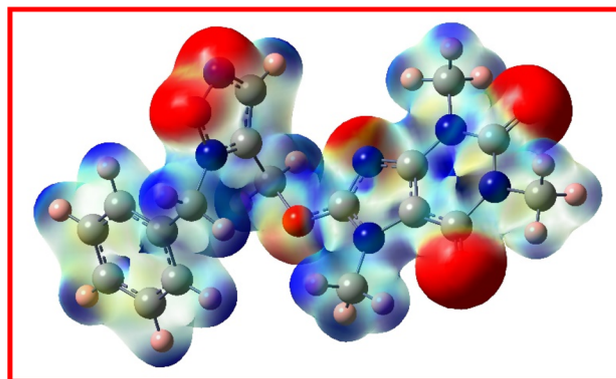
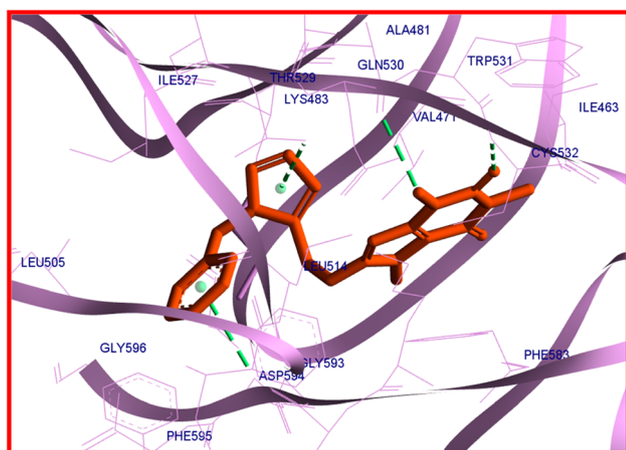
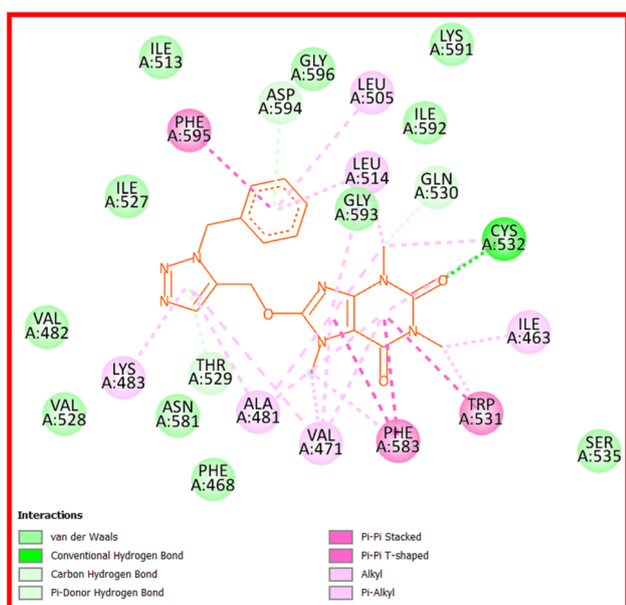


Fig. 8 The overlay view of conformations of co-crystallized vemurafenib (green), redocked vemurafenib (yellow), and **22c** (orange) at the binding site of B-RAF V600E (PDB code: 3OG7).



caffeine, one π -donor hydrogen bond between Thr529 and triazole ring, and one π -donor hydrogen bond between Asp594 and phenyl ring of benzyl group. The three-dimensional (3D) docked conformation of **22c** including hydrogen bonds and its two-dimensional (2D) docked conformation in the active site of enzyme are displayed in Fig. 9 and 10, respectively. The side chains of Trp531, Phe583, and Phe595 were involved in the π - π^* interactions with the caffeinyl moiety and phenyl ring of **22c**. The Ile463, Leu505, Leu414, Val471, Ala481, Lys483, Trp531, Cys532, and Phe583 create a hydrophobic pocket to stabilize **22c** in the active site of enzyme. The van der Waals interactions were also observed between **22c** and the amino acids of active site including Phe468, Val482, Ile513, Val528, Ser535, Asn581, Lys591, Ile592, Gly593, and Gly596. The calculated ΔG values for

Fig. 11 MEP of **22c**.Fig. 9 3D docked conformation and hydrogen bonds (green) of **22c** with B-RAF V600E kinase enzyme.Fig. 10 2D docked conformation of **22c** at the active site of B-RAF V600E kinase enzyme.

vemurafenib and **22c** were -160.41 and -151.02 kcal mol $^{-1}$, respectively. The total hydrogen bond energies for vemurafenib and **22c** were -9.50 and -3.24 (kcal mol $^{-1}$), respectively. These results indicated that interactions such as π - π^* , the hydrophobic and the van der Waals interactions are more important than hydrogen bonding interactions for stabilization of **22c** in the active site of B-RAF V600E kinase enzyme.

The molecular electrostatic potential (MEP) is a useful parameter to define the susceptible regions of a molecule to electrophilic or nucleophilic interaction with the surface of enzymes and receptors.⁶⁴ In this regard, MEP of **22c** was obtained using the DFT method at the B3LYP/6-31+G** level of theory (Fig. 11). As shown in Fig. 10, the atoms with the positive electrostatic potentials are depicted in blue and the regions carrying the negative electrostatic potentials are displayed in red. The negative electrostatic potentials are located around the triazole ring and some heteroatoms of caffeine moiety. Thus, these regions have high electron density and could interact with the electrophilic groups of B-RAF V600E kinase enzyme as confirmed by docking analysis.

3 Conclusion

In conclusion, it was explained the design and synthesis of the new 8-caffeinyl-triazolylmethoxy hybrid conjugates. These compounds were assessed for the *in vitro* anticancer and the cytotoxicity against two breast cancer cell lines MDA-MB-468 (ATCC HTB-22), MCF-7 (ATCC HTB-22), melanoma cell line A-375 (ATCC CRL-1619) and normal cell line HEK-293 (ATCC CRL-11268). The results have determined that **22c** displayed the highest and lowest toxicity against A375 and HEK-293 compared to methotrexate (MTX) as a standard drug, respectively. Compounds **22f** and **22i** have demonstrated the most potent activities against MCF-7 and MDA-MB-468 but their activities were weaker than MTX. The *in silico* studies were performed using the SwissADME online software, the OSIRIS Property Explorer and the preADMET online server to determine the physicochemical properties and pharmacokinetic profile. The obtained data indicated that most designated compounds were matched with established rules such as Lipinski's rule of five (RO5) and *etc*. The molecular docking study was also conducted



by the Molegro Virtual Docker (MVD 6.0) software to predict the binding mode and the interaction of **22c** as the most active anti-melanoma compound with B-RAF V600E kinase enzyme. The docking results determined that **22c** exhibited a strong binding affinity to the active site of the enzyme.

4 Experimental

4.1. Materials and cell lines

Pasteur Institute of Iran supplied the used cancer cell lines, including two breast cancer cell lines MDA-MB-468 (ATCC HTB-22), MCF-7 (ATCC HTB-22), melanoma cell line A-375 (ATCC CRL-1619) and normal cell line HEK-293 (ATCC CRL-11268). Phosphate-buffered saline (PBS) tablets, tetrazolium salt, 3-(4,5-dimethyl-thiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) were purchased from Sigma-Aldrich (USA). Penicillin streptomycin, trypsin, dimethyl sulfoxide (DMSO), and Dulbecco's Modified Eagle's Media (DMEM) cell culture medium were obtained from Shellmax (China). Fetal bovine serum (FBS) was supplied from Gibco (USA).

4.2. Determination of the anticancer activity

The anticancer activity of 10 new compounds and Methotrexate (MTX) were investigated using MTT assay.⁶² The various cell lines were cultured in DMEM complete medium (containing FBS 10% and penicillin-streptomycin 1%) and incubated at 37 °C air (95%) and CO₂ (5%). Cells (MDA-MB-468, MCF-7, A-375 and HEK-293) were separated by trypsin; after that; they were seeded into 96-well cell culture plates (separately) at a cell density of 5000 cells per well and incubated for 24 h. The culture media was discarded, cells were exposed to serial dilutions of MTX or the tested compounds in the 12.5 to 300 μM range (100 μL per well). After incubation (24 h, 37 °C), the plates' content was discarded, and wells were replaced with 100 μL PBS to wash. Then, 100 μL MTT solution (0.5 mg. mL⁻¹) was added to each well and incubated for another 4 h. In continuation, 100 μL per well DMSO was added to dissolve formed formazan crystals. In each plate, six wells were considered as the control group, which was filled with 200 μL per well of DMEM. Finally, the absorbance (A) was measured at 570 nm on a plate reader (ELISA Plate Reader). The cell viability at each concentration was calculated by eqn (1). IC₅₀ values (μM) were determined from the dose response curves as the concentration of the compound caused a 50% decrease in MTT reduction compared to the control (DMSO) using Prism 7 (GraphPad).

$$\text{Cell viability(\%)} = \frac{\text{Mean } A \text{ sample}}{\text{Mean } A \text{ control}} \times 100 \quad (1)$$

4.3. Chemistry general

All chemicals were purchased from Merck and/or other chemical vendors and straightforwardly applied without further purifications. The 8-bromocaffeine (8-BC) was synthesized *via* our previously explained procedure in the literature.^{27,34} The Reactions were monitored by TLC using SILG/UV 254 silica-gel plates. The purifications by a short column chromatography

were done on the silica gel 60 (0.063–0.200 mm, 70–230 mesh; ASTM). The melting points are measured on the Electrothermal IA 9000 in open capillary tubes and they are uncorrected. The PerkinElmer 240 B micro-analyzer, Shimadzu GC/MS-QP 1000-EX apparatus (*m/z*; rel.%), and the Shimadzu FT-IR-8300 spectrophotometer were used to obtain the elemental analyses, GC/MS, and IR spectra, respectively. ¹H and ¹³C-NMR spectrum was recorded on the Brüker Avance-DPX-300 spectrometer operating at 300/75 MHz, respectively. Chemical shifts are given in δ relative to tetramethylsilane (TMS) as an internal standard, coupling constants *J* are given in Hz. The abbreviations used for ¹H-NMR signals are: s = singlet, d = doublet, t = triplet, q = quartet, m = multiplet, br = broad and *etc.*

4.4. General procedure for synthesis of alkyl azides

Alkyl azides were made according to the established method reported in the literature⁴²

4.5. General procedure for synthesis of (1-alkyl-1H-1,2,3-triazol-4-yl)methanol (23)

In a double-necked round bottom flask (100 mL), it was added a mixture of propargyl alcohol (0.56 g, 10 mmol), desire alkyl azide (12 mmol) and CuI (0.05 g, 0.26 mmol) in THF/H₂O (1 : 1 V/V, 30 mL). The reaction vessel was stirred at reflux until TLC monitoring indicates no further progress of the reaction (6–12 h). Then, the catalyst was separated (sintered glass) and the filtrate was evaporated *in vacuo*. Afterward, the residue was diluted in CHCl₃ (200 mL) and washed with H₂O (3 × 100 mL). Finally, the obtained crude product was pure enough that it was used for the subsequent reaction without requiring further purification.

4.6. General procedure for synthesis of 8-caffeinyl-triazolymethoxy hybrid conjugates (22a–22j)

In a double-necked round bottom flask (100 mL), it was added a mixture of 8-BC (2.73 g, 10 mmol), desire **23** (12 mmol), and KOH (0.56 g, 10 mmol) in DMSO (15 mL). The reaction vessel was stirred at 100 °C until TLC monitoring indicates no further progress of the reaction (24 h). Then, the reaction was diluted in distilled water and the organic phase was extracted by CHCl₃ (3 × 100 mL). Finally, the obtained crude product was purified by a simple recrystallization or a short column chromatography using the solvents described below.

4.6.1 1,3,7-Trimethyl-8-((1-(3-phenylpropyl)-1H-1,2,3-triazol-4-yl)methoxy)-1H-purine-2,6(3H,7H)-dione (22a). Column chromatography on silica gel with *n*-hexane/EtOAc (1:10) afforded the pure product as white solid (2.98 g, 73%); m. p. 164–166 °C; IR (KBr): 1150, 1476, 1591, 1690, 1720, 2974, 3100 cm⁻¹. ¹H NMR (300 MHz, CDCl₃) δ_{ppm} = 2.27–2.32 (m, 2H, PhCH₂CH₂), 2.65–2.67 (m, 2H, PhCH₂), 3.10 (s, 3H, NCH₃), 3.22 (s, 3H, NCH₃), 3.45 (s, 3H, NCH₃), 4.41 (br. s, 2H, NCH₂), 5.84 (s, 2H, OCH₂), 7.16–7.30 (m, 5H, aryl), 7.71 (s, 1H, C(5)-H of triazole). ¹³C NMR (75 MHz, CDCl₃) δ_{ppm} = 18.84, 28.18, 28.97, 33.28, 34.79, 53.49, 73.45, 105.79, 121.51, 127.03, 127.81, 128.85, 137.65, 142.50, 149.78, 151.30, 152.57, 155.85. MS (EI):



m/z (%) = 409 (20.8) [M^+]. Anal. Calc. for $C_{20}H_{23}N_7O_3$: C, 58.67; H, 5.66; N, 23.95; found: C, 58.81; H, 5.84; N, 24.06.

4.6.2 1,3,7-Trimethyl-8-((1-phenethyl-1H-1,2,3-triazol-4-yl)methoxy)-1H-purine-2,6(3H,7H)-dione (22b). Column chromatography on silica gel with *n*-hexane/EtOAc (1 : 10) afforded the pure product as creamy solid (2.80 g, 71%); m. p. 169–171 °C; IR (KBr): 1163, 1482, 1594, 1686, 1715, 2956, 3078 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 2.56 (t, J = 7.2 Hz, 2H, $PhCH_2$), 3.35 (s, 3H, NCH_3), 3.52 (s, 3H, NCH_3), 3.62 (s, 3H, NCH_3), 4.44 (t, J = 6.9 Hz, 2H, NCH_2), 6.23 (s, 2H, OCH_2), 7.17–7.32 (m, 5H, aryl), 8.39 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 19.19, 28.46, 31.91, 35.21, 58.05, 73.53, 104.79, 121.66, 126.99, 128.60, 129.54, 140.42, 143.20, 150.13, 151.24, 152.63, 156.57. MS (EI): m/z (%) = 395 (22.5) [M^+]. Anal. Calc. for $C_{19}H_{21}N_7O_3$: C, 57.71; H, 5.35; N, 24.80; found: C, 57.87; H, 5.50; N, 24.98.

4.6.3 8-((1-Benzyl-1H-1,2,3-triazol-4-yl)methoxy)-1,3,7-trimethyl-1H-purine-2,6(3H,7H)-dione (22c). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as creamy solid (2.93 g, 77%); m. p. >265 °C (dec.); IR (KBr): 1145, 1476, 1598, 1680, 1718, 2969, 3050 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 3.38 (s, 3H, NCH_3), 3.50 (s, 3H, NCH_3), 3.72 (s, 3H, NCH_3), 5.39 (s, 2H, NCH_2), 5.71 (s, 2H, OCH_2), 7.27–7.30 (m, 5H, aryl), 7.86 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 18.27, 29.49, 32.60, 57.69, 73.47, 105.09, 120.21, 126.43, 128.46, 129.61, 137.42, 142.65, 150.67, 151.61, 153.42, 155.29. MS (EI): m/z (%) = 381 (15.9) [M^+]. Anal. Calc. for $C_{18}H_{19}N_7O_3$: C, 56.69; H, 5.02; N, 25.71; found: C, 56.86; H, 5.19; N, 25.90.

4.6.4 1,3,7-Trimethyl-8-((1-(4-methylbenzyl)-1H-1,2,3-triazol-4-yl)methoxy)-3,7-dihydro-1H-purine-2,6-dione (22d). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as white solid (3.28 g, 83%); m. p. 213–215 °C; IR (KBr): 1153, 1485, 1596, 1689, 1720, 2983, 3069 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 2.26 (s, 3H, $PhCH_3$), 3.19 (s, 3H, NCH_3), 3.41 (s, 3H, NCH_3), 3.64 (s, 3H, NCH_3), 5.27 (s, 2H, NCH_2), 5.51 (s, 2H, OCH_2), 7.14–7.21 (m, 4H, aryl), 8.15 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 19.33, 25.10, 29.53, 31.81, 59.54, 73.40, 104.12, 121.26, 128.63, 129.80, 134.62, 136.15, 142.50, 150.37, 151.26, 152.93, 155.62. MS (EI): m/z (%) = 395 (24.3) [M^+]. Anal. Calc. for $C_{19}H_{21}N_7O_3$: C, 57.71; H, 5.35; N, 24.80; found: C, 57.58; H, 5.20; N, 24.62.

4.6.5 8-((1-Benzhydryl-1H-1,2,3-triazol-4-yl)methoxy)-1,3,7-trimethyl-1H-purine-2,6(3H,7H)-dione (22e). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as bright brown solid (3.65 g, 80%); m. p. 213–215 °C; IR (KBr): 1145, 1479, 1590, 1693, 1720, 2951, 3050 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 3.20 (complex, 6H, $2NCH_3$), 3.64 (s, 3H, NCH_3), 5.31 (s, 2H, OCH_2), 6.28 (s, 1H, NH), 7.21–7.38 (m, 10H, aryl), 8.18 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 18.98, 29.66, 32.67, 70.25, 74.42, 104.81, 121.29, 127.07, 128.24, 129.86, 138.65, 143.55, 150.23, 151.62, 153.29, 155.13. MS (EI): m/z (%) = 457 (28.4) [M^+]. Anal. Calc. for $C_{24}H_{23}N_7O_3$: C, 63.01; H, 5.07; N, 21.43; found: C, 62.86; H, 4.93; N, 21.25.

4.6.6 1,3,7-Trimethyl-8-((1-pentyl-1H-1,2,3-triazol-4-yl)methoxy)-1H-purine-2,6(3H,7H)-dione (22f). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as pink solid (2.16 g, 60%); m. p. >273 °C (dec.); IR (KBr): 1156, 1459, 1597, 1690, 1718, 2971, 3028 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} =

0.85 (t, J = 6.9 Hz, 3H, CH_2CH_3), 1.26–1.30 (m, 4H, $(CH_2)_2CH_3$), 1.43–1.53 (m, 2H, NCH_2CH_2), 3.34 (s, 3H, NCH_3), 3.48 (s, 3H, NCH_3), 3.70 (s, 3H, NCH_3), 3.93 (t, J = 7.2 Hz, 2H, NCH_2), 5.53 (s, 2H, OCH_2), 7.76 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 13.84, 18.37, 21.80, 27.69, 28.83, 29.70, 32.42, 53.34, 73.28, 105.09, 120.94, 142.73, 150.47, 151.61, 152.76, 155.26. MS (EI): m/z (%) = 361 (21.9) [M^+]. Anal. Calc. for $C_{16}H_{23}N_7O_3$: C, 53.17; H, 6.41; N, 27.13; found: C, 53.35; H, 6.56; N, 26.98.

4.6.7 8-((1-Hexyl-1H-1,2,3-triazol-4-yl)methoxy)-1,3,7-trimethyl-1H-purine-2,6(3H,7H)-dione (22g). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as white solid (2.36 g, 63%); m. p. 135–137 °C; IR (KBr): 1149, 1470, 1590, 1692, 1720, 2983, 3050 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 0.81 (br s, 3H, CH_2CH_3), 6.57 (br s, 6H, $(CH_2)_3CH_3$), 1.83 (br s, 2H, NCH_2CH_2), 3.30 (s, 3H, NCH_3), 3.52 (s, 3H, NCH_3), 3.93 (s, 3H, NCH_3), 4.24 (t, J = 7.2 Hz, 2H, NCH_2), 5.27 (s, 2H, OCH_2), 7.65 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 13.54, 18.93, 23.14, 27.57, 28.33, 29.91, 31.56, 32.73, 52.61, 73.44, 104.04, 121.38, 142.66, 151.08, 152.03, 153.16, 154.84. MS (EI): m/z (%) = 375 (20.7) [M^+]. Anal. Calc. for $C_{17}H_{25}N_7O_3$: C, 54.39; H, 6.71; N, 26.12; found: C, 54.57; H, 6.90; N, 26.28.

4.6.8 8-((1-Heptyl-1H-1,2,3-triazol-4-yl)methoxy)-1,3,7-trimethyl-1H-purine-2,6(3H,7H)-dione (22h). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as white solid (2.29 g, 59%); m. p. 122–124 °C; IR (KBr): 1130, 1459, 1597, 1695, 1718, 2964, 3057 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 0.82–0.84 (m, 3H, CH_2CH_3), 1.23–1.30 (complex, 8H, $(CH_2)_4CH_3$), 1.89 (br. s, 2H, NCH_2CH_2), 3.34 (s, 3H, NCH_3), 3.51 (s, 3H, NCH_3), 3.63 (s, 3H, NCH_3), 4.32 (t, J = 7.2 Hz, 2H, NCH_2), 5.57 (s, 2H, OCH_2), 7.70 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 14.47, 19.52, 22.74, 27.91, 29.08, 29.98, 30.70, 32.32, 33.50, 53.52, 73.82, 105.25, 121.36, 143.22, 150.21, 152.11, 153.74, 155.36. MS (EI): m/z (%) = 389 (17.8) [M^+]. Anal. Calc. for $C_{18}H_{27}N_7O_3$: C, 55.51; H, 6.99; N, 25.18; found: C, 55.36; H, 6.81; N, 25.02.

4.6.9 1,3,7-Trimethyl-8-((1-octyl-1H-1,2,3-triazol-4-yl)methoxy)-1H-purine-2,6(3H,7H)-dione (22i). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as creamy solid (2.62 g, 65%); m. p. 117–119 °C; IR (KBr): 1145, 1467, 1593, 1691, 1720, 2975, 3071 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 0.82 (t, J = 6.9 Hz, 3H, CH_2CH_3), 1.22–1.32 (complex, 10H, $(CH_2)_5CH_3$), 1.87–1.92 (m, 2H, NCH_2CH_2), 3.35 (s, 3H, NCH_3), 3.52 (s, 3H, NCH_3), 3.64 (s, 3H, NCH_3), 4.33 (t, J = 7.2 Hz, 2H, NCH_2), 5.58 (s, 2H, OCH_2), 7.70 (s, 1H, C(5)-H of triazole). ^{13}C NMR (75 MHz, $CDCl_3$) δ_{ppm} = 14.43, 18.87, 22.19, 26.85, 28.48, 29.66, 30.62, 31.80, 32.22, 53.55, 73.22, 105.54, 121.72, 142.80, 150.72, 152.35, 153.31, 155.16. MS (EI): m/z (%) = 403 (22.8) [M^+]. Anal. Calc. for $C_{19}H_{29}N_7O_3$: C, 56.56; H, 7.24; N, 24.30; found: C, 56.74; H, 7.40; N, 24.49.

4.6.10 8-((1-Decyl-1H-1,2,3-triazol-4-yl)methoxy)-1,3,7-trimethyl-1H-purine-2,6(3H,7H)-dione (22j). Recrystallization with CH_2Cl_2 /EtOAc afforded the pure product as white solid (3.02 g, 70%); m. p. 115–117 °C; IR (KBr): 1137, 1456, 1590, 1694, 1718, 2958, 3050 cm^{-1} . 1H NMR (300 MHz, $CDCl_3$) δ_{ppm} = 0.84 (t, J = 6.9 Hz, 3H, CH_2CH_3), 1.24–1.31 (complex, 14H, $(CH_2)_7CH_3$), 1.89–1.93 (m, 2H, NCH_2CH_2), 3.37 (s, 3H, NCH_3), 3.53 (s, 3H,



NCH₃), 3.66 (s, 3H, NCH₃), 4.34 (t, $J = 7.2$ Hz, 2H, NCH₂), 5.59 (s, 2H, OCH₂), 7.70 (s, 1H, C(5)-H of triazole). ¹³C NMR (75 MHz, CDCl₃) $\delta_{\text{ppm}} = 13.48, 18.35, 21.88, 26.53, 27.92, 28.65, 29.55, 30.28, 32.35, 33.75, 52.67, 72.94, 106.09, 121.71, 142.25, 150.15, 152.06, 153.23, 155.30$. MS (EI): m/z (%) = 431 (24.7) [M⁺]. Anal. Calc. for C₂₁H₃₃N₇O₃: C, 58.45; H, 7.71; N, 22.72; found: C, 58.31; H, 7.86; N, 22.58.

Conflicts of interest

The authors have declared no conflict of interest.

Acknowledgements

The authors wish to thank Shiraz University of Technology research council for partial support of this work.

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