Application of triazoles in the structural modification of natural products

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1. Introduction

More than 200 years ago, a 21-year-old pharmacist’s apprentice named Friedrich Sertürner isolated the first pharmacologically active pure compound from a plant. This compound was morphine derived from opium produced by cut seed pods of the poppy *Papaver somniferum*. This opened an era in which the precise dosage of the purification, research and application of botanicals would not change with the source or age of the substance. After World War II, due to the discovery of penicillin, drug research expanded to large-scale screening of microorganisms to find new antibiotics. By 1990, drugs composed of natural products or analogs inspired by natural products accounted for 80%. The discovery and production of antimicrobial drugs (e.g. doxorubicin and taxol), immunosuppressants for organ transplants (e.g. rapamycin and cyclosporine), lipid control agents (e.g. lovastatin and analogs), antimalarials (e.g. artemisinin and quinine), antiparasitics (e.g. avermectin), and antibiotics (e.g. tetracycline, penicillin, and erythromycin) revolutionised medicine. It is not difficult to see from these findings that natural products play a very important role in the process of drug discovery and development. Newman’s team published a series of overviews of natural products as sources of new drugs5–6, and also reported the sources of antitumor compounds7, publishing intermediate reports describing natural products as leads to potential drugs8. All of these articles emphasise the inextricable importance of natural products and/or natural product structures in drug discovery and development.

Many marketed drugs contain heterocycles, and triazoles with a five-membered ring composed of two carbon atoms and three nitrogen atoms exist in different heterocycles. There are two types of triazole – 1,2,3-triazole and 1,2,4-triazole (Figure 1). Triazole can be readily obtained, and the framework can act as an amide, ester, carboxylic acid, and other heterocycles such as pyrazole isosteres10. By affecting the hydrogen bonding ability, polarity and lipophilicity of the molecules, the triazole moiety can improve the physicochemical properties, toxicology, pharmacokinetics and pharmacology of the compounds11,12. The synthetic moieties containing these molecular structures have been used extensively in the discovery of drugs due to their low occurrence in nature13. Meanwhile, on the basis of the literature, triazole and its derivatives have aroused enormous interest owing to their pharmaceutical and therapeutic applications, including their use as anticonvulsant14–17, antidepressant18, antitumor20–22, antiviral23–24, antimicrobial25–33, anti-acetylcholinesterase35, anti-inflammatory36, antioxidant37–40, antiparasitic41–43, and anti-diabetic drugs44. Their ability to produce various non-covalent interactions to improve solubility and binding to bimolecular targets may be the reason for this wide applicability45. Furthermore, a number of drugs that contain 1,2,3-triazole scaffolds, including TSAO46 (anti-HIV agent), Cefazolin47 (an antibiotic), CAI48 (anti-cancer agent), and Tazobactum49 (anti-bacterial agent), are currently used in clinical applications (Figure 2). The favourable properties of the enhanced biological activities of the triazole ring include hydrogen bonding capability under various conditions, a strong dipole moment, high chemical stability (they are typically inert for oxidising and reducing agents), and rigidity.

Therefore, given that natural products have become the source of most active ingredients in medicines, and the interest focussed on triazole compounds is increasing recently, we will review the latest research progress in regard to triazole-containing natural products in pharmaceutical chemistry.
2. Biological activities

2.1. Anticancer activity

2.1.1. Anticancer activity of lung cancer

Matrine (chemical formula: C15H24N2O, molecular weight: 248.36, (7aS,13aR,13bR,13cS)-dodecahydro-1H,5H,10H-dipyrido[2,1-f:3',2',1'-ij][1, 6]naphthyridin-10-one) is a quinolizidine alkaloid that is an important active compound found in the root of *Sophora flavescens* Ait (also known as Kushen). Zhao et al.\(^5\)\(^0\) found that the conjugation of matrine, 1H-1,2,3-triazole, and chalcones could form novel anticancer agents that exerted synergistic effects where the double bond of the \(\alpha,\beta\)-unsaturated moiety plays a dominant role. Adding 2'-OH into the A ring or substituting the B ring of chalcone with EWGs may increase the anticancer activity of matrine–triazole–chalcone conjugates. Among the conjugates, compound 1 (Figure 3) was 8.0-fold more potent (IC\(_{50}\) = 5.01 ± 0.59 \(\mu\)M) than was 5-fluorouracil (IC\(_{50}\) = 40.38 ± 4.61 \(\mu\)M) and possesses comparable potency to that of paclitaxel (IC\(_{50}\) = 2.82 ± 0.31 \(\mu\)M) against A549 cells. Additionally, compound 1 possesses a relatively broad anticancer spectrum and exhibits less cytotoxicity (IC\(_{50}\) = 39.21 ± 4.31 \(\mu\)M) than that of 5-fluorouracil (IC\(_{50}\) = 22.36 ± 2.09 \(\mu\)M) and paclitaxel (IC\(_{50}\) = 20.01 ± 2.38 \(\mu\)M) against NIH3T3 cells. Flow cytometry tests demonstrated that compound 1 could induce apoptosis in A549 cells in a concentration-dependent manner, and efficiently suppressed human tumour growth in a mouse xenograft model without causing obvious toxicities.

Asiatic acid (chemical formula: C\(_{30}\)H\(_{48}\)O\(_{5}\), molecular weight: 488.70, (2a,3b,4a)-2,3,23-trihydroxyurs-12-en-28-oic acid) is a pentacyclic triterpenoid isolated from the tropical medicinal plant *Centella asiatica* (Apiaceae). Huang et al.\(^5\)\(^1\) synthesised this compound and discovered that asiatic acid-based 1,2,3-triazoles derivatives could act as antitumor agents by blocking nuclear factor kappa B (NF-\(\kappa\)B) activation and cell migration. Among the tested compounds, the hydroxyl group on R2 is more beneficial than the acetyl group. Polar groups were investigated at R1 using halogen group substitution, where the 2-fluorine derivative with hydroxyl moiety substitution at R2 (Figure 3), which showed the highest activity among the investigated compounds, yielded in a significantly lower IC\(_{50}\) of 0.14 \(\mu\)M. A molecular docking study was performed to identify key interactions between compound 2 and NF-\(\kappa\)B, where the 1,2,3-triazoles moiety and the hydroxyl groups of the asiatic acid skeleton were important for improving inhibitory activity. Subsequently, surface plasmon resonance analysis verified the high affinity between compound 2 and NF-\(\kappa\)B protein with an equilibrium dissociation constant (KD) value of 0.36 \(\mu\)M. Further studies revealed that compound 2 markedly inhibited NF-\(\kappa\)B DNA binding, nuclear translocation, and I\(\kappa\)B\(\alpha\) phosphorylation. Moreover, in vitro antitumor activity screening revealed that compound 2 (IC\(_{50}\) = 2.67 ± 0.06 \(\mu\)M) exhibited the best anticancer activity against A549 cells, and this was achieved, at least in part, through the inhibition of NF-\(\kappa\)B activity. Treatment of A549 cells with compound 2 resulted in the induction of apoptosis and inhibition of cell migration *in vitro*.

![Figure 1](image1.png)  
**Figure 1.** Chemical structures of 1,2,3-triazole and 1,2,4-triazole motifs.

![Figure 2](image2.png)  
**Figure 2.** Some of the drugs available in the market containing 1,2,3-triazole ring.

![Figure 3](image3.png)  
**Figure 3.** The chemical structures of anti-lung cancer compounds 1–5.
Xanthone (chemical formula: C_{13}H_{12}O_{2}, molecular weight: 224.26, 9H-xanthen-9-one) is a bioactive substance that can be isolated from plants and from microorganisms. The basic skeleton of this compound is a biphenyl pyranone possessing a planar three-ring structure. Wu et al. synthesized several xanthone derivatives and evaluated their cytotoxicity. The IC_{50} analysis indicated that the inhibitory activity of the 18 target compounds was higher than that of the original xanthone intermediate. The extensive structure of various substituted groups resulted in a significantly increased anticancer effect versus compound with the benzyl group. In detail, the compounds with substituted electron-donating groups possessed higher activity versus those with electron-withdrawing groups. Of note, the compounds with weak electron-withdrawing groups demonstrated the highest activity in this series. Among compounds with substituted halogen, compound 3 (Figure 3) with replaced para bromide was the most active agent against A549 cancer cells (IC_{50} = 32.4 ± 2.2 μM). Western blotting analyses revealed that compound 3 significantly increased the expression of caspase 3, Bax, and c-Jun N-terminal kinase and also positively regulated p53 in cancer cells.

Flavanone (chemical formula: C_{15}H_{12}O_{2}, molecular weight: 224.26, 2,3-dihydro-2-phenyl-4H-1-benzopyran-4-one) is a flavonoid that exists as a polyphenol found in the plant kingdom. On average, the intake of flavonoids is approximately 50–150 mg per day from vegetables, fruits, and other food sources. Flavanone is an important natural secondary metabolite. A series of new flavanone-triazole hybrids were synthesised by Gutam et al. The entire synthesised group of compounds exhibited a subjective but diverse cytotoxic effect against the HCT-15, HeLa, and NCI-H522 cell lines. Among these compounds, compound 4 (Figure 3) exhibited the highest cytotoxicity against NCI-H522 cells (IC_{50} = 5.4 μM) and possessed an improved safety profile.

Camptothecin (chemical formula: C_{20}H_{17}N_{2}O_{4}, molecular weight: 348.35, (4S)-4,4-dihydro-4-hydroxy-1H-pyrano[3′,4′:4,5]indolizino[1,2-b]quinoline-3,14(4H,12H)-dione) is a topoisomerase I inhibitor that was first isolated from Camptotheca acuminata by Wall and Wani in 1966. Xu et al. reported the modifications and SAR of homocamptothecin at position C10 and used these findings to develop potent topoisomerase I inhibitors for anticancer drug discovery. Compounds 5a, 5b, and 5c (Figure 3), possessing cyclopropyl, isopropyl, and cyclohexyl groups, respectively, exhibited very high inhibitory activities that were 4–6 times more effective than camptothecin against A549 cells. The IC_{50} values of these compounds were 30, 30, and 50 nM, respectively, and compound 5a exhibited stronger topo I-dependent cytotoxic activity than did camptothecin at concentrations of 100 and 10 μM. Furthermore, compound 6 could cause cell cycle arrest in the G2 and S phases at a concentration of 0.1 μM. Interestingly, most of the alkyl and cycloalkyl groups promoted the antiproliferative activities of 1,2,3-triazole homocamptothecin derivatives against A549 cells, while the benzyl groups did not. On the contrary, the benzyl group was favourable to MDA-MB-435 cells. Surprisingly, all the glycosyl compounds showed moderate antiproliferative activities against MDA-MB-435 and HCT116, which might be due to their relatively bulky spatial positions. These findings provide valuable insights for further development of more effective homocamptothecins as antitumor agents.

2.1.2. Anticancer activity of breast cancer

Coumarin (chemical formula: C_{14}H_{16}O_{2}, molecular weight: 196.20, 9H-coumaran-9-one) was discovered in 1820 as a derivative of the tonka bean. Coumarin exists in many plants in the form of glycosides. Natural and synthetic coumarin derivatives have attracted great attention among medicinal chemists due to their wide range of biological activities. The anticancer activity of the newly synthesised triazole-linked N-glycosides of coumarins and quinolones was determined by Kumari et al. Compound 6 (Figure 4) displayed low micromolar (IC_{50} = 10.97 μM) and selective toxicity against MCF-7, a breast cancer cell line. Further study revealed that the anticancer activity of the active compound was due to the formation of reactive oxygen species (ROS) without significant DNA damage. Apart from causing DNA lesions, ROS production in the cell can also cause oxidative modifications of proteins leading to their altered functions in the cell or leads to lipid peroxidation which can generate toxic products in the cell. Since in study the active compound showed breast cancer cell line (MCF-7) specific cell death without significant nuclear DNA damage, it might be possible that other cellular macromolecules like proteins or lipids essential for the survival of targeted cell lines could be the target of this ROS generation.

Dehydroabietic acid (chemical formula: C_{20}H_{30}O_{2}, molecular weight: 300.44, (1R,4α5,10αS)-1,2,3,4,4a,9,10,10α-octahydro-1a,4a,8a,10a-tetramethyl-a-phenanthrene-carboxylic acid) is a natural resin acid that can be easily obtained from disproportionated rosin. A series of novel C-14, 1,2,3-triazole-tethered dehydroabietic acid derivatives possess antiproliferative activity. These new analogs remained effective against adriamycin-resistant MCF-7 cells at low concentrations in a dose-dependent manner. The results suggested that the introduction of 1,2,3-triazole moiety was beneficial to cytotoxicity. Further, the effect of various substituents on the C-4 position of the 1,2,3-triazole moiety was also examined and the structure activity relationship (SAR) studies revealed that the introduction of aromatic substituents was crucial for the potent cytotoxicity. Generally, the introduction of electron-rich aromatic ring system could significantly increase the cytotoxicity. Whereas, the introduction of electron-poor aromatic ring system would hamper the cytotoxicity. For example, pyridyl- and nitrobenzyl-substituted analogues only showed weak or loss of cytotoxicities. In comparison, the saturated alkyl-, ester- and tri-methyl chlorosilane- on the C-4 position of the 1,2,3-triazole moiety were only associated with moderate increase in the growth inhibitory effect. In particular, derivative 7 (Figure 4) possessing a 3-[(tert-butoxycarbonylamino)phenyl]-substituted thiazole moiety exhibited the highest potency with IC_{50} values ranging from 0.7 μM to 1.2 μM, and these values are more potent than those of the commercial anticancer drug 5-Fu (average IC_{50} value 16.1 μM). Moreover, compound 7 not only possessed broad-spectrum anti-proliferative activities against a series of tumour cells derived from different organs, but also it exhibited very weak cytotoxicity on normal cells, implying the existence of a therapeutic window for the use of compound 7.

Chalcones (chemical formula: C_{13}H_{12}O_{2}, molecular weight: 280.26, (E)-1,3-diphenyl-2-propen-1-one) are naturally occurring flavonoids that possess 1,3-diphenyl-2-propen-1-one as their framework. Chalcone is an α,β-unsaturated ketone that represents a central core for a variety of important bioactive molecules. A series of 1H-1,2,3 triazole-grafted tetrahydro-β-carboline-chalcone/ferrocenylchalcone conjugates was synthesised by Sharma et al. Analysis of SAR revealed that aryl chalcone based conjugates showed better anti-proliferative activities on both the cell lines tested. Among TH/IC-chalcone conjugates, the nature of substituent on phenyl ring of chalcone predominantly played an important role in enhancing the cytotoxicity on breast cancer cell lines whereas length of alkyl chain hardly affected the activities. Compounds with electron donating tri-methoxy substituents on...
phenyl ring displayed appreciable cytotoxicities on breast cancer cells as compared to compounds with mono-methoxy substituent, which were inactive on both breast cancer cell lines. Among trimethoxylated conjugates, Compound 8a (Figure 4) possessed an electron-donating trimethoxy substituent on the phenyl ring of chalcone and pentyl as a spacer and was the most active against MDA-MB-231 cells with an IC50 value of 21.99 μM and was therefore ~3 folds potent than Tamoxifen. Interestingly, the compounds with electron withdrawing fluorosubstitution at phenyl ring were found to be the most active amongst all the synthesised conjugates. Among these conjugates, compound 8b (Figure 4), possessing an optimum combination of electron-withdrawing and lipophilic 4-fluoro substituents on the phenyl ring of chalcone and a propyl chain as the spacer, proved to be the most potent with an IC50 value of 10.33 μM against MCF-7. Gurrapu et al.61 also synthesised novel 1,2,3-triazole chalcone hybrids as potential anti-cancer agents. All of these compounds were effective; however, meta methyl substituent attached to the triazole ring meta, para dimethoxy substituted attached to the chalcone ring of compounds 9a (Figure 4), meta chloro substitutes attached to the triazole ring and meta, para dimethoxy substituted attached to the chalcone ring of compound 9b (Figure 4), and meta chloro substituent attached to the triazole ring and meta hydroxy substituted attached to the chalcone ring of compound 9c (Figure 4) were nearly equipotent and exhibited increased efficiency against cancer cell lines. In particular, 9b exhibited the best cytotoxic activity against MCF-7 and other cell lines, displaying an IC50 of 1.27 μM and 0.02 μM at 24 and 48 h, respectively. The other compounds exhibited intermediate to moderate cytotoxic activities against the tumour cells in comparison to the cytotoxicity of cisplatin. Finally, SAR data revealed that compounds which have chloro and methoxy substituent at different position have shown promising activity when compared with other derivatives and remaining compounds showed moderate cytotoxic activity.

Costunolide (chemical formula: C15H20O2, molecular weight: 232.32, 2-Hydroxy-1,4-dihydronaphthalene-1,4-dione) is a natural bioactive compound isolated from plants of the genus Lawsonia. Kumar et al.63 successfully synthesised 20 analogs of costunolide and dehydrocostuslactone against the tested cell lines. Compound 10 (Figure 4) was demonstrated as best analog with a GI50 of < 0.12 μM against the MDA MB-231 cell line, a value that is better than that of the parent compound costunolide (GI50 = 0.56 μM). These preliminary studies laid a solid foundation for further lead optimisation of this class of compounds by a systematic chemical modification including the synthesis of water-soluble compounds to improve their overall pharmaceutical properties.

Lawsone (chemical formula: C15H20O2, molecular weight: 232.32, 2-Hydroxy-1,4-dihydronaphthalene-1,4-dione) is a natural bioactive compound isolated from plants of the genus Lawsonia. Ottoni et al.64 synthesised two series of glycosidic derivatives of Lawsone, and these corresponded to classical glycosides and glycosyl triazoles. All compounds displayed acceptable activity against the SKBR-3 cell line with IC50 values below 10 μM. The greater activity of peracetylated glycosides and glycosyl triazoles as compared to lawsone is probably due to the more favourable lipophilic–hydrophilic balance that has been achieved with the peracetylated glycosyl derivatives which could be absorbed by tumour cells more easily. The most promising derivative was the glycosyl triazole derived from peracetylated d-glucose (11) (Figure 4), exhibiting improved cytotoxicity against SKBR-3 cells (IC50 = 0.78 μM) and superior selectivity towards the tumour cell line (SI > 20). All compounds described in this work were more active than was Lawsone, thus indicating the importance of the carbohydrate and glycosyl triazole moiety for activity.

2.1.3. Anticancer activity of gastric cancer

Celastrol (Chemical formula: C29H38O4, molecular weight: 450.61, (9β,13α,14β,20z)-3-Hydroxy-9,13-dimethyl-2-oxo-24,25,26-trinoroleana-1(10),3,5,7-tetraen-29-oic acid) is a quinone methide triterpene that is an active ingredient first extracted from the roots of the Chinese medicinal plant “Thunder of God Vine” (Celastraceae, Tripterygium). Three series of novel celastrol derivatives were designed and synthesised by Zhang et al.65. The introduction of 1,2,3-triazole linked to benzyl fluoride exerted a minor influence on the anticancer activity of celastrol. However, the celastrol derivative 12 (Figure 5), a 1,2,3-triazole linked to benzyl chloride,
exerted an effective anti-proliferative effect on AGS cells (IC\textsubscript{50} = 0.97 \mu M).

Xanthotoxin (chemical formula: C\textsubscript{12}H\textsubscript{8}O\textsubscript{4}, molecular weight: 216.19, 9-methoxy-7H-furo[3,2-g][1]benzopyran-7-one) is a furanocoumarin and an active compound of the traditional Egyptian medicinal plant \textit{Ammi majus L}. Xanthotoxin-triazole derivatives possess antiproliferative properties\textsuperscript{66}. The SAR analysis revealed that the phenyl-substituted derivatives showed better activity compared with the benzyl-substituted derivatives and highlighted the importance of the substituted benzene of this molecular modification for the antiproliferative activity in xanthotoxin-linked 1,2,3-triazoles. Among the phenyl-substituted derivatives, the order of potency was p-CF\textsubscript{3} > p-Cl > p-F > p-CH\textsubscript{3} > p-OCH\textsubscript{3}. Additionally, for the phenyl-substituted xanthotoxin-linked 1,2,4-triazoles, the order of potency was p-F > p-CF\textsubscript{3}. Thus, compounds with an electron-withdrawing group (–CF\textsubscript{3}, –Cl, –F) at the 4-position of the benzyl moiety displayed better antitumor activity. Among the synthesised compounds, 9-[[1-(4-(trifluoromethyl)phenyl)-1H-1,2,3-triazol-4-yl)methoxy]-7H-furo[3,2-g]chromen-7-one (13) (Figure 5) exhibited the strongest antiproliferative activity against AGS cells (IC\textsubscript{50} = 7.5 \mu M) and possessed improved activity compared to those of the lead compound (xanthotoxin, IC\textsubscript{50} > 100 \mu M) and the reference drug (5-fluorouracil, IC\textsubscript{50} = 29.6 \mu M). The IC\textsubscript{50} value of compound 13 in L02 cells was 13.3-fold higher than that in the AGS cells. Therefore, the compound exhibited improved therapeutic activity and specificity compared to that of the positive control 5-fluorouracil. Cell cycle analysis revealed that compound 13 inhibited cell growth via the induction of S/G2 phase arrest in AGS cells.

2.1.4. Anticancer activity of ovarian cancer

Apigenin (chemical formula: C\textsubscript{15}H\textsubscript{10}O\textsubscript{5}, molecular weight: 270.24, 5,7-dihydroxy-2-(4-hydroxyphenyl)-4H-1-benzopyran-4-one) is a naturally occurring flavonoid compound that exists in a variety of plants in the form of phyto-yellow pigment, and this compound is primarily derived from the Umbelliferae plant \textit{Apium graveolens L}. Qi et al.\textsuperscript{67} found that novel triazole analogs of apigenin-7-methyl ether exhibit potent antitumor activity against ovarian carcinoma cells via the induction of mitochondria-mediated apoptosis. Of all the derivatives, the derivative 14 (Figure 6) exhibited significant and dose-dependent anticancer activity against the SKOV3 ovarian cancer cell line. The IC\textsubscript{50} of compound 14 was 10 \mu M against the SKOV3 cancer cell line. Compound 14 induced apoptosis in SKOV3 cancer cells through the accretion of reactive oxygen species and a reduction in mitochondrial membrane potential. This molecule also modulated the expression of B-cell lymphoma 2 (Bcl-2) and Bcl-2-associated X protein.

2.1.5. Anticancer activity of liver cancer

As noted earlier, derivatives of coumarin-triazole 15 exhibit anti-breast cancer activity\textsuperscript{58}. Kraljevic et al.\textsuperscript{68} synthesised a new 4-substituted 1,2,3-triazole-coumarin hybrid possessing anti-liver cancer activity. The incorporation of benzofused heterocycles led to improvements in activities. The 7-methylcoumarin-1,2,3-triazole-2-methylbenzimidazole hybrid 15 (Figure 6) can be highlighted as exerting the highest cytotoxicity against hepatocellular carcinoma HepG2 cells with an IC\textsubscript{50} value of 0.9 \mu M and high selectivity (SI = 50). This compound induced cell death that occurred primarily due to early apoptosis. The growth-suppressive properties of compound 15 in HepG2 cells could be associated with its ability to inhibit 5-lipoxygenase (5-LO) and acid ceramidase activities, as these inhibitions may, in turn, lead to the accumulation of arachidonic acid and ceramide, respectively.

Dehydroabietic acid-triazole derivatives not only exhibit anti-breast cancer activity\textsuperscript{59} but also possess anti-liver cancer activity\textsuperscript{69}. A series of dehydroabietic acid-coupled 1,2,3-triazole derivatives was synthesised by Li et al.\textsuperscript{69} The synthesised compounds were screened for cytotoxic activity against a panel of four human cancer cell lines and the human HL-7702 normal cell line using a 3-(4,5-dimethylthiazolyl-2)-2,5-diphenyltetrazoliumbromide (MTT) assay. Of these compounds, a number exhibited better antitumor activity against the tested cancer cell lines compared to that of the positive control cisplatin, and they also exhibited low cytotoxicity against the human normal liver cell line HL-7702, indicating that the introduction of 1,2,3-triazole moiety on the DHAA skeleton increased anti-tumour activity. In particular, compounds 16a and 16b (Figure 6) exhibited good antitumor activity against HepG2 with IC\textsubscript{50} values of 5.90 ± 0.41 and 6.25 ± 0.37 \mu M, better than those of positive control cisplatin.

2.1.6. Anticancer activity of pancreatic cancer

Coumarin-triazole derivatives not only possess anti-breast cancer activity\textsuperscript{58} but also exhibit anti-liver cancer activity\textsuperscript{68} and anti-pancreatic cancer activity\textsuperscript{70}. Farley et al.\textsuperscript{70} synthesised a series of functionalised coumarins and evaluated their capacity to inhibit the resistance to starvation in pancreatic cancer cells. Evaluation of two trifluoromethylphenyl compounds against three cancer cell lines showed that position of the trifluoromethyl substituent on the phenyl ring of these compounds (meta- vs para-) was correlated to selectivity for activity against MIA PaCa-2 cell line. This relatively small change in structure had a substantial effect on activity. The \textit{meta}-trifluoromethylphenyl derivative 17 (Figure 6) exhibited preferential cytotoxicity against PANC-1, Capan-1, and MIA PaCa-2 cells with PC\textsubscript{50} concentrations of 29, 8.5, and 18 \mu M, respectively. Apoptosis was determined as the mechanism of cell death (PANC-1, compound 17), based on a modified ethidium bromide and acridine orange (EB/AO) staining assay.

2.1.7. Anticancer activity of Colon cancer

Isosteviol (chemical formula: C\textsubscript{20}H\textsubscript{30}O\textsubscript{3}, molecular weight: 318.45, 4z,8\beta,13β)-13-Methyl-16-oxo-17-norkauran-18-oic acid) is a tricyclic diterpenoid possessing an ent-beyerane skeleton that exhibits multifarious bioactivities and can be readily obtained as a metabolite of stevioside isolated from the leaves of the natural stevia plant\textsuperscript{71,72}. Liu et al.\textsuperscript{73} designed and synthesised a series of novel 1,2,3-triazole-linked isosteviol derivatives using the
The cytotoxicities of these compounds against HCT-116 and JECO-1 cells were screened in vitro. From the observed cytotoxic activity data, it has been noticed that all the derivatives of isosteviol showed better cytotoxic activities than their corresponding precursor. More importantly, the inhibitory activities of most compounds were markedly improved as the 1,2,3-triazole subunit was introduced onto the skeleton of isosteviol, which indicated the 1,2,3-triazole fragment exactly played a significant role in inhibiting cancer cell proliferation. Going even further, substituted groups and positions on aromatic ring had a significant effect on cytotoxic activities. Ortho-position on aromatic ring has an important effect on cytotoxic activity. Compounds with aldehyde group on aromatic ring exhibited better inhibitory activities than the unsubstituted compounds. Oxidation of aldehyde group caused weaker activities to the cancer cell lines. To be noteworthy, the inhibitory activities of isosteviol simultaneously fusing hydroxyl and 1,2,3-triazole subunits were better than that of compounds with only 1,2,3-triazole subunit, which illustrates that introduction of hydroxyl group can result in higher inhibitory activity against HCT-116 cells. In particular, compound 18 (Figure 7) exhibited the most potent inhibitory activity against HCT-116 cells with an IC₅₀ value of 2.987 ± 0.098 μM, and this was better than that (3.906 ± 0.261 μM) of the positive control cisplatin. On the basis of these bioactivity data, hologram quantitative structure activity relationship was performed, and a statistically reliable model with good predictive power (r² = 0.848, q² = 0.544 and R²pred = 0.982) was achieved. The contribution maps derived from the optimal model explained the individual atomic contributions to the activity for each molecule.

Combretastatin A-4 (chemical formula: C₁₈H₂₀O₅, molecular weight: 316.35, 2-methoxy-5-[(1H-1,2,3-triazol-1-yl)phenol]) is one of the most well-known anticancer agents and a statistically reliable model with good predictive activity against HCT-116 cells. To be noteworthiness, the inhibitory activities of isosteviol simultaneously fusing hydroxyl and 1,2,3-triazole subunits were better than that of compounds with only 1,2,3-triazole subunit, which illustrates that introduction of hydroxyl group can result in higher inhibitory activity against HCT-116 cells. In particular, compound 18 (Figure 7) exhibited the most potent inhibitory activity against HCT-116 cells with an IC₅₀ value of 2.987 ± 0.098 μM, and this was better than that (3.906 ± 0.261 μM) of the positive control cisplatin. On the basis of these bioactivity data, hologram quantitative structure activity relationship was performed, and a statistically reliable model with good predictive power (r² = 0.848, q² = 0.544 and R²pred = 0.982) was achieved. The contribution maps derived from the optimal model explained the individual atomic contributions to the activity for each molecule.

Combretastatin A-4 (chemical formula: C₁₈H₂₀O₅, molecular weight: 316.35, 2-methoxy-5-[(1H-1,2,3-triazol-1-yl)phenol]) is among the most well-known anticancer agents and a statistically reliable model with good predictive power (r² = 0.848, q² = 0.544 and R²pred = 0.982) was achieved. The contribution maps derived from the optimal model explained the individual atomic contributions to the activity for each molecule.

Among the 1,2,3-triazole subunits and its substitutions were critical for keeping antiproliferative effect and the antiproliferative activities were almost lost when the thiol group was free or replaced by methylthio or ethylthio. Secondly, when changing the thiol group of the 3-position of triazole ring with benzylthio, the cytotoxic activities against HepG2, PC-3 and HCT116 cells were significantly increased by the chain elongation. Meanwhile, the introduction of electron withdrawing groups such as fluoro atom on the benzyl group, also caused a slight enhancement of the antiproliferative activity. These results suggest that electronic effect of substituents on benzyl group plays a crucial role on antitumor activities. Besides, linker-length of alkylsulfanyl moiety has also profound effects on the antiproliferative activities. Introduction of phenyl acetyltio substitutes on the 3-position of triazole ring leads to dramatical enhancement of antiproliferative activities against HepG2 cell lines, but naphthyl, cyclopropyl and ethoxyl groups result in dramatic decrease of the activities. It is worth noting that compound 19a (Figure 7), with N-4 chlorophenyl acetamidethio substitute, showed more potent in vitro cytotoxic activities against PC-3 with IC₅₀ values of 6.29 μM, which represented threefold improvement in activity compared to combretastatin A-4. Moreover, further flow-activated cell sorting analysis revealed that compound 19a displayed a significant effect on G2/M cell-cycle arrest in a dose dependent manner in PC-3 cells. Within the series of N,N-dimethyl analogues, the effects of substituents on the antiproliferative activities were strongly correlated with the 4-methoxy analogues. Meanwhile, analogue 19b (Figure 7) was an exception, which displayed fivefold improvement compared to combretastatin A-4 in inhibiting HCT116 cell proliferation with IC₅₀ values of 1.15 μM. More interestingly, analog 19b also displayed the most potent anti-tubulin activity with a percentage of 49% at 10 μM.

Betulinic acid (chemical formula: C₃₀H₅₆O₅, molecular weight: 456.7, (3R)-3-hydroxy-20(29)-en-28-oic acid) is a bioactive pentacyclic lupane-type triterpenoid that can be directly isolated from *Platanus orientalis* stem bark and from many other plants such as the birch tree *Betula spp.* (Betulaceae), *Ziziphus spp.* (Rhamnaceae), *Syzygium spp.* (Myrtaceae), *Diospyros spp.* (Ebenaceae), and *Paeonia spp.* (Paeoniaceae). A new library of compounds possessing a 1,2,3-triazole moiety attached to C-3 of betulinic acid was synthesised, and the anti-cancer activities of these compounds were evaluated (in vitro) against different cancer cell lines (i.e. breast, colon, liver, and leukemia) by Chakraborty et al. The structure activity relationship studies indicate that the 1,2,3-triazole moiety favours the activity when substituted at C-4 (of the heterocycle) with a hydroxymethyl group, but disfavours it when the substituent is an aromatic or heteroaromatic moiety. Compound 20 (Figure 7) was found to be the most potent inhibitor of the cell line HT-29 with an IC₅₀ value of 14.9 μM. This activity profile was improved compared to that of the parent compound (betulinic acid). Its role as an inducer of apoptosis was investigated in this cell line using an Annexin-V/PI binding assay and by following its capability for ROS generation, depolarisation of mitochondrial transmembrane potential, activation of caspases, PARP cleavage, nuclear degradation, and expression of pro- and anti-apoptotic proteins. This compound exhibited much higher cytotoxicity than did the standard drug 5-fluorouracil; however, it exhibited negligible cytotoxicity towards normal PBMCs. Elevated levels of ROS generation, activation of caspase 3 and caspase 9, DNA fragmentation, higher expression of Bax and Bad, lower expression of Bcl2 and Bcl-x, and increased levels of Bax/Bcl-x ratio identified compound 20 as a promising inducer of
apoptosis that follows a mitochondria-dependent pathway. Biological studies indicate that compound 20 acts as a minor groove binder to the DNA.

2.1.8. Anti-leukaemia activity
Andrographolide (chemical formula: C_{30}H_{56}O_{5}, molecular weight: 350.45, (3E,4S)-3-[2-[(1R,4αS,5R,6R,8aS)-decahydro-6-hydroxy-5-(hydroxymethyl)-5,8a-dimethyl-1-naphthalenyl]ethylidene]dihydro-4-hydroxy-2(3H)-furanone) is one of the labdane diterpenoids that are the principal active constituents of Andrographis paniculata. A series of new andrographolide-1,2,3-triazole derivatives were synthesised from the natural bioactive labdane-type diterpenoid andrographolide. All of the derivatives were screened against the human cancer cell lines MCF7, MDA-MB-231, COLO205, HepG2, K562, Hela, and HEK293 to evaluate their cytotoxic activity. All of these compounds exhibited anti-cancer activity selectively against the K562 cell line with IC_{50} values ranging from 8.00 to 17.11 μM and were inactive against the rest of the cell lines. Andrographolide-1,2,3-triazole can be considered as parent moiety, and the substitution at the first position of triazole was varied. It is very clear that the first position with a substituted benzene ring found to be active over unsubstituted or aliphatic substitution, and electron-donating groups are preferred over withdrawing groups. The increase in the carbon chain length linking the substituted benzene with the triazole group reduces the activity. Compounds 21a and 21b (Figure 8) exhibited better cytotoxicity against K562 cell lines compared to that of the other compounds in the series with 62.9% and 51.8% inhibition at 50–μM concentration IC_{50} values of 8 and 9.7 μM, respectively.

Betalenic acid-triazole derivatives not only possess anti-colon cancer activity but also exhibit an anti-leukaemia effect. A novel triazole derivative of betulinic acid induces extrinsic and intrinsic apoptosis in human leukaemia HL-60 cells. The structure activity relationship of structural modifications can be summarised as follows: the more electron-negative p-fluoro exhibited much better cytotoxicity in all the cancer cell lines compared to p-bromo derivative which showed only 61% and 54% of growth inhibition against MiaPaCa-2 and A549 cancer cell lines. The probable reason for the same may be the high lipophilicity, by molecule can penetrate easily to the cells and exerted its activity. Generally, compounds containing electron withdrawing functional groups (–F, –Cl) exhibited more potent cytotoxic effects against the cancer cells compared to the electron donor functional groups (–CH_{3}, –OCH_{3}). Among all of the tested compounds, compound 22 (Figure 8) displayed promising antiproliferative activity in all four cell lines (HL-60, MiaPaCa2, PC-3, and A549), with IC_{50} values of 7, 5, 7, and 7 μM, respectively. The cytotoxic profile of compound 22 was similar to that of betulinic acid. Based on this, Khan et al. chose this compound for use in further cell death mechanistic studies in HL-60 cells. The effect of compound 22 on DNA damage and apoptosis was investigated using cell cycle analysis. It was observed that treatments with up to 10 μM of compound 22 resulted in mild induction of apoptosis (7%). Furthermore, the cell cycle G1 phase was significantly blocked by compound 22 at a concentration of 20 μM, indicating that it caused a delay in the cell cycle. Compound 22 induced loss of mitochondrial membrane potential in a concentration-dependent manner. Compound 22 at 5 μM induced a 7% loss of mitochondrial membrane potential, and this increased to 18% and 27% at 10 and 20 μM concentrations, respectively. Compound 22 decreased the expression of mitochondria-associated anti-apoptotic protein Bcl-2 in a concentration-dependent manner. HL-60 cells treated with compound 22 exhibited significant loss of mitochondrial membrane potential. Compound 22 significantly decreased the level of the mitochondrial antiapoptotic protein Bcl-2 and increased the expression of the pro-apoptotic protein Bax with a concurrent decrease in the Bcl-2/Bax ratio. Compound 22 treatment results in the activation of caspases and in PARP-1 cleavage. Compound 22 inhibited both Procaspase-9 and Procaspase-8, indicating that it induced apoptosis via both the extrinsic and intrinsic pathways. Therefore, we can conclude that it induces apoptosis via both intrinsic and extrinsic activation pathways in HL-60 cells. The 18C-30-triazole-substituted betulin and betulinic acid derivatives were synthesised by Shi et al. These compounds were tested for their cytotoxic activity against the leukaemia cell line HL-60 using an MTT assay. New C-30 triazole-substituted betulinic acid derivatives exhibited improved cytotoxic activity compared to that of betulin derivatives (> 25 μM). The majority of the new triazole-substituted betulinic acid derivatives displayed improved bioactivity compared to that of betulinic acid (11.5 μM). The rank order of potency based on the C-30 triazole substituent is 4-fluorophenyl > n-hexyl > phenyl = 2-thienyl > n-butyl > cyclopropyl. Overall, the compound 23 (Figure 8) [4-(4-fluorophenyl)-1H-1,2,3-triazol-1-yl] betulinic acid possessed the best IC_{50} value (1.3 μM) against the leukaemia cell line HL-60 (eight- to ninefold higher potency than that of betulinic acid). Therefore, larger C-30 side chains with aromatic substitutions were favoured for the cytotoxic activity.

2.1.9. Miscellaneous
 Diosgenin (chemical formula: C_{27}H_{44}O_{3}, molecular weight: 414.63, (3β,25S)-Spirost-5-en-3-ol) is rich in the tubers of D. deltoidei, where it exist as a steroidal sapogenin. Diosgenin is structurally similar to cholesterol and to other steroids, and this compound is in high demand in the pharmaceutical industry. Both diosgenin and its analogs exhibited interesting anti-proliferative effects against four human cancer cell lines (HBL-100 [breast], A549 [lung], HT-29 [colon], and HCT-116 [colon]) according to the results of MTT assays. Among the synthesised analogs, compound 24a (Figure 9) that possesses a simple phenyl R moiety attached via triazole to the parent molecule was identified as the most
potent analog against the A549 cancer cell line. This analog possessed an IC50 of 5.54 μM, which was improved compared to that of the positive control (BEZ-235). Compounds 24b (Figure 9) and 24c (Figure 9) that possess o-nitrophenyl and o-cyanophenyl R moieties, respectively, displayed impressive anti-proliferative activity against all the tested human cancer cell lines and exhibited IC50 values ranging from 5.77 to 9.44 μM. These observations highlight the beneficial impact of electron withdrawing ortho substituents attached to R moiety towards the anti-proliferative activity.

Berberine (chemical formula: C20H18NO4, molecular weight: 336.37, 5,6-dihydro-9,10-dimethoxybenzo[g]-1,3-benzodioxolo[5,6-a]quinolizinium) is a naturally occurring isoquinoline alkaloid that can be isolated from various Berberis plants.85,86 Jin et al.87 designed and synthesised a series of new triazolyl berberine derivatives. These synthesised compounds and their anticancer activities were evaluated against a panel of four human cancer cell lines that included MCF-7 (breast), MCF-7/ADR (breast), SW-1990 (pancreatic), and SMMC-7721 (liver) and against the non-cancer cell line HUVEC (human umbilical vein endothelial cell). The results revealed that the majority of the compounds exhibited improved anticancer activities against MCF-7 and SMMC-7721 compared to that of berberine. SAR analysis indicated the following: (i) hetero-aryl substituents at the R position showed poor anticancer activity against four cancer cell lines; (ii) when methylene-naphthalene was substituted for methylene-quinoline, the cytotoxicity was significantly reduced. The results of the in vitro anticancer activity tests indicated that compounds 25a and 25b (Figure 9) exhibited the most potent inhibitory activities against the SMMC-7721 and SW-1990 cell lines with IC50 values of 14.861 ± 2.4 μM and 16.798 ± 3.4 μM, respectively.

Bergenin (chemical formula: C14H16O9, molecular weight: 328.27, 2R,3R,5,6-dihydroxy-2-(hydroxymethyl)-9-methoxypyrano[3,2-c][2]benzopyran-1-one) represents a dihydroisocoumarin derivative. Kumar et al.88 designed and synthesised a series of novel bergenin-trizole hybrid derivatives and evaluated their anti-cancer activities against DU-145, A549, HCT-116, Hep G2, and HeLa cell lines. It is evident that majority of synthetic derivatives displayed better cytotoxic activity than parent compound on A549, HepG2 and HeLa cell lines. Preliminary structure activity relationship indicates that presence of a substituent such as a chloride atom or a cyano, floro and CF3 group on aromatic triazole partner enhanced activity with IC50 values ranging from 1.33 to 9.9 μM on A459 cell as well as HeLa cell lines within the studied series. Among them, compound 26 (Figure 9) demonstrated potent activity against A549 and HeLa cell lines with IC50 values of 1.86 μM and 1.33 μM, respectively, and was equipotent to doxorubicin. Furthermore, mechanistic studies revealed that compound 26 triggered cell cycle arrest at the G2/M phase and induced cell apoptosis in a dose- and time-dependent manner. Taken together, these results indicate that compound 26 effectively inhibited tubulin polymerisation and disrupted the intracellular tubulin-microtubule balance, ultimately resulting in prolonged G2/M cell cycle arrest. Docking studies also indicated a strong hydrophobic interaction with tubulin, thus leading to stable binding and subsequent apoptosis of cancer cells. A series of triazole derivatives of bergenin were synthesised by Yang et al.89 The introduction of 1,2,3-triazole into the bergenin skeleton, exhibit enhancing effect in terms of anti-tumour activities than bergenin. The electron properties and steric effects of the substituents of the benzene ring affect the antiproliferative activity. Either an electron-donating substituent such as 4-methyl, 4-ethyl and 4-tert-butyl group or an electron-withdrawing substituent such as 4-methoxy, 4-fluoro, 4-nitro and 4-nitro group of the benzene ring of the side chain are all more effective than the unsubstituted benzene derivative. Particularly, the introduction of larger steric hindrance substituent on the benzene ring, such as the tert-butyl group, caused compound 27 (Figure 9) to exhibit significantly improved antiproliferative activity against three cancer cell lines (EC9706, MGC803, and B16) compared to that of bergenin. The IC50 dropped to 6.2 μmol/L, 12.0 μmol/L, and 17.6 μmol/L, respectively, and this compound had a more effective antiproliferative activity than the positive drug 5-fluorouridine against the cancer cell line EC9706.

Coumarin-triazole derivatives not only exhibit anti-breast cancer activity,88 anti-liver cancer activity,68, and anti-pancreatic cancer activity70, but also they possess activity against multiple types of cancer cells. A series of coumarin-based target-specific probes for cancer theranostic applications that played a dual role in the field of both diagnosis and therapy was screened for anticancer activity against breast cancer (MCF7) and human epitheloid cervix carcinoma (HeLa) cell lines90. All hybrids with IC50 values ranging from 9.83 to 26.21 μM exhibited high potency against MCF7 and HeLa cell lines, and this potency was comparable to that of cisplatin (IC50: 18 and 10 μM) but was less than that of doxorubicin (IC50: 5.2 and 3.83 μM). The SAR revealed that the electron-withdrawing groups-Cl and-Br on the 1,2,3-triazole motif boost the anticancer profile in both cell lines, while the electron-donating groups-OH and -OMe and the alkyl groups were unfavourable in regard to activity. The most active was hybrid 28 (Figure 9) (IC50: 17.5 and 9.83 μM), and this compound was slightly more potent than cisplatin and also exhibited the lowest cytotoxicity in normal human foetal lung fibroblast (MRC-5) cells (IC50: 185.22 μM). Moreover, hybrid 28 exhibited strong cellular uptake in the MCF-7 cell line. Thus, hybrid 28 may be useful for cancer theranostics due to its high potency, low cytotoxicity, and strong cellular uptake. The coumarin derivatives also exhibited improved antiproliferative activities against several cancer cell lines.91 It is necessary to point out that the cytotoxic activities of the derivatives with
different halogen substitutions on the benzene ring were in the following order, $p$-Cl > o-Cl > m-Cl, $p$-CH$_3$ > o-CH$_3$ > m-CH$_3$ and 2,4-(CH$_3$)$_2$ > 2,6-(CH$_3$)$_2$. Based on an overall comparison, the compounds derived from structures with electron-withdrawing substituents on the 1,2,4-triazole ring exhibited potent activity, and those with electron-donating substituents on the 1,2,4-triazole ring exhibited moderate activity, against the six cancer cell lines. Compound 29 (Figure 9) exhibited potent activity against AGS, MGC-803, and HCT-116 cell lines, where the IC$_{50}$ values were 2.63 ± 0.17, 3.05 ± 0.29, and 11.57 ± 0.53 μM, respectively. This compound also exerted strong activity against the HeLa cell line, with an IC$_{50}$ value of 13.62 ± 0.86 μM. A more detailed mechanistic
study demonstrated that compound 29 could inhibit the proliferation of AGS cancer cells by inducing apoptosis and arresting cells in the G2/M phase. In an attempt to develop potent anti-tubulin agents against various cancers, a library of 28 novel triazole-tethered isatin-coumarin hybrids was synthesised through the use of a click chemistry approach. Cytotoxicity results revealed an interesting structure activity relationship for the designed hybrids: (i) type of substituent on isatin and the length of carbon-bridge connecting isatin moiety with triazole ring considerably influences cytotoxic potential of hybrids; (ii) presence of unsubstituted isatin remarkably enhances the cytotoxic potential; (iii) enhanced cytotoxicity for hybrids having isatin with electron withdrawing substituent in comparison to electron donating substituent; (iii) toxicity for hybrids having isatin with electron withdrawing substituent remarkably enhances the cytotoxic potential; (ii) enhanced cytotoxicity increases with increase in electronegativity of substituent on isatin; Thus, the overall preference order of R is as follows: H > F > Cl > Br > I > NO2 > OCH3; (iv) cytotoxicity decreases significantly with the increase in chain length of carbon-bridge. Thus, the overall preference order of n (chain length) is as follows: 1 > 2 > 3 > 4. The most active was hybrid 30 (IC50: 0.73, 3.45, and 3.04 μM against THP-1, COLO-205, and HCT-116 cancer cell lines), and this compound also displayed the most potent anti-tubulin activity with an IC50 value of 1.06 μM. The in vitro tubulin polymerisation assay clearly indicated that these hybrids exert their anti-cancer activity through tubulin inhibition. Singh et al. synthesised triazole ring-binding molecule hybrids of C5-curcumino- noid and coumarin. Cytotoxicity results revealed an interesting structure activity relationship for these designed hybrids: (i) methoxy substituted phenyl ring remarkably enhances the cytotoxic potential; (ii) placement of a heteroaryl ring such as furan and thiophene in place of the unsubstituted phenyl ring improved the activity profile; (iii) an enhanced effect was observed with the increased number of methoxy substituents on phenyl ring such as trimethoxy phenyl > dimethoxy phenyl > monomethoxy phenyl; (iv) placement of naphthyl ring behaved as a surrogate for dimethoxy substituted phenyl ring; (v) cytotoxicity of hybrids with monomethoxy substituted phenyl ring was found similar to the heteroaryl ring substituted hybrids. Thus, the overall preference order of ring is established as follows: trimethoxy phenyl > dimethoxy phenyl = naphthyl > monomethoxy phenyl = furan = thiophene > phenyl. The most active hybrid 31 with trimethoxy phenyl ring exhibited significant cytotoxicity with IC50 values ranging from 0.82 to 4.68 μM against THP-1, HCT-116, and COLO-205 cell lines, respectively. Compound 31 also displayed the most potent anti-tubulin activity, with an IC50 value of 1.55 μM.

Melanomagnolide B (chemical formula: C31H32O6, molecular weight: 496.52, 1b,10a,10b-Octahydrop-5-(hydroxymethyl)-1a-methyl-8-methyleneoxireno9,10b-Octahydro-5-(hydroxymethyl)-1a-methyl-8-methyleneoxireno derivatives can potent NF-κB inhibition and cell cycle arrest in multiple cancer cells lines. Compound 32 possessed nanomolar activity (EC50 = 0.26 μM) against SH-SY5Y cells, Compound 33 exerted moderate inhibitory activity against MCF-7, U87, MGC-803, and EC109 cells (IC50 = 32.25, 9.57, 5.95, and 20.77 μM, respectively). Compound 34 arrested the cell cycle at the G2/M phase, induced apoptosis accompanied by a decrease in mitochondrial membrane potential, and potently inhibited LSD1 (IC50 = 3.18 μM). Docking studies indicated that compound 34 formed interactions with the surrounding amino acid residues, and the steroid nucleus occupied the tubular hydrophobic cavity of the active site.

Derivatives of isosteviol-triazole not only possess anti-colon cancer activity, but also they exhibit anti-multiple cancer cell activity. Compounds with different phenyl 1,2,3-triazole chloroacetamide showed considerably higher antiproliferative activity against the HCT-116 and HepG2 cell lines. Perhaps the triazole acts as a hydrogen bond acceptor and binds to some key enzymes involved in cancer cell metabolism, inhibiting their expression. In particular, compound 35 possesses a methyl group introduced at the para position of the benzene ring exhibited the strongest antiproliferative activity among all of the target compounds (IC50 values of 5.38 ± 0.26 μM, 15.91 ± 0.41 μM, and 8.92 ± 0.44 μM against HCT-116, BEL-7402, and HepG2 cell lines, respectively). Compound 35 was 4.6-fold (against HCT-116 cells), 1.3-fold (against BEL-7402 cells), and 2.6-fold (against HepG2
cells) more active than the positive control drug 5-fluorouracil. The compound was 18.6-fold (against HCT-116 cells), 6.3-fold (against BEL-7402 cells), and 11.2-fold (against HepG2 cells) more active than the lead compound isovitexin. Compound 35 also inhibited colony formation in HCT-116 cells in a concentration-dependent manner. Cell cycle analysis revealed that compound 35 inhibited cell growth via the induction of S phase arrest in HCT-116 cells. The possible mechanism of action may be correlated with downregulation of cyclin A and cyclin E1 expression and with the upregulation of cyclin B1 expression.

Oridonin (chemical formula: C_{20}H_{28}O_{6}, molecular weight: 364.39, \((12\times6,7\times4,14\times7)-20\text{-Epoxy-1,6,7,14-tetrahydroxykaure-16-en-15-one}\)) was initially isolated from various Isodon species that are commonly used as a home remedy herb in China and Japan. Derivatives of oridonin and triazole possess anti-tumour activity\(^{39}\). All oridonin derivatives containing different phenyl 1,2,3-triazoles exhibited stronger anti-proliferative activities against all three selected cancer cell lines than did oridonin and 5-Fu. Compound 36a (Figure 9), with 4-methoxophenyl 1,2,3-triazole, was the most potent compound in the series against the HCT116 cell line, with an IC\(_{50}\) value of 1.94 \(\mu\)M. This compound was approximately threefold more potent than oridonin against the tested cancer cell lines. Compound 36b (Figure 9), with 4-methylphenyl 1,2,3-triazole, was the most potent compound in this series, with an IC\(_{50}\) value of 3.01 \(\mu\)M in MCF-7 cell lines. This compound was approximately sixfold more potent than oridonin against the tested cancer cell lines. Preliminary results suggested that the phenyl 1,2,3-triazole groups would improve the anti-proliferative activities of oridonin, and highlighting the importance of the linker.

Ferulic acid (chemical formula: C_{10}H_{12}O_{4}, molecular weight: 194.18, \(3-(4\text{-Hydroxy-3-methoxyphenyl})-2\text{-propanoic acid}\)) is an abundant phenolic phytochemical found in plant cell walls. Aneja and et al.\(^{100}\) reported potent and selective ferulic acid-based small molecule inhibitors of carbonic anhydrase IX that possess significant inhibitory potential against various oncogenic parameters. The potency of compounds bearing \(-\text{Cl}, -\text{NO}_2\) and \(-\text{COOH}\) functionality was significantly enhanced in comparison to its natural precursor, ferulic acid with IC\(_{50}\) in the range of 0.024–3.78 \(\mu\)M. Fluorine substituent did not exhibit any significant effect on the activity against these carbonic anhydrase isoforms. However, the effect of electron-donating substituents on inhibition of carbonic anhydrase isoforms was found to be moderate only in the range of 1.96–6.67 \(\mu\)M. Interestingly, it was found that compound 37 (Figure 9) selectively inhibited carbonic anhydrase IX in the nanomolar range (IC\(_{50}\) = 24 \(n\)M). In silico analysis revealed the binding of compound 37 to the catalytically important amino acid residues of carbonic anhydrase IX. Further, cell-based studies indicated that compound 37 inhibits the activity of carbonic anhydrase IX, decreases epithelial to mesenchymal transition, induces apoptosis, and inhibits cell migration and colonisation potential in cancer cells. Taken together, these results emphasise the potential for use of compound 37 as a prospective pharmacological lead molecule in carbonic anhydrase IX-targeted anticancer therapeutics.

Derivatives of chalcones-triazole not only exert anti-breast cancer activity\(^{50,61}\), but also they exhibit activity against multiple types of cancer cells\(^{101,102}\). A new series of 1,2,3-triazole-chalcone hybrids was synthesised by Ashour and et al.\(^{101}\). The results suggest that the electronic and steric properties of the substituents play an important role in the binding affinity of chalcones to their cellular target(s). As a general statement, the presence of meta \(\text{OCH}_3\) group on the right side and \(\text{para}\ \text{Cl}\) atom on the left side of the hybrid was proved to be essential for general anticancer activity of the tested series. The para chloro compound 38 (Figure 9) that possesses a 3–4-dimethoxyphenyl chalcone moiety was the most potent derivative and inhibited the growth of RPMI-8226 and SR leukaemia cell lines by 99.73% and 94.95% at 10 \(\mu\)M, respectively. Furthermore, this compound inhibited the growth of M14 melanoma, K-562 leukaemia, and MCF7 breast cancer cell lines by more than 80% at the same test concentration. Compound 38 exhibited IC\(_{50}\) values of less than 1 \(\mu\)M against six types of tumour cells and possessed a high selectivity index that reached 104-fold in MCF7. Compound 38 possessed superior activity compared to that of methotrexate and gefitinib against the most sensitive leukaemia cell lines, and it exhibited higher or comparable activity against the other sensitive cell lines. Flow cytometry analysis of RPMI-8226 cells revealed that compound 38 caused cell cycle arrest at the G2/M phase and induced apoptosis in a dose-dependent manner. Mechanistic evaluation indicated that apoptosis induction triggered the mitochondrial apoptotic pathway by inducing ROS accumulation and increasing the Bax/ Bcl-2 ratio and the activation of caspases 3, 7, and 9. The current study clearly identified the potential of compound 38 as a promising lead for the future development of active anticancer agents, and the results of this study may offer new insights for treating multiple myeloma based on the data generated using RPMI-8226 cells. A series of chalcone linked-1,2,3-triazoles was synthesised by Yadav and et al.\(^{102}\). All of the synthesised products were subjected to MTT cytotoxicity assays against a panel of four human cancer cell lines (MCF-7, MIA-Pa-Ca-2, A549, and HepG2) to assess their anticancer potential. Structure activity relationship results demonstrated the broad spectrum anticancer potential of compounds bearing methoxy substituents on ring A. In case of triazole derivative, the introduction of bromo or fluoro groups on benzene ring C led to decrease in anticancer activity compared to the parent compound 39 especially against MCF-7 and MIA-Pa-Ca-2 cell lines. Compound 39 exhibited more than 20-fold and 25-fold cytotoxicity towards the MCF-7 and MIA-Pa-Ca-2 cell lines as compared to normal cell lines F2. All the synthesised triazoles were found to be less toxic towards the normal cells compared to the reference. Compound 39 (Figure 9) was found to be most active against all of the tested cancer cell lines, with IC\(_{50}\) values in the range of 4–11 \(\mu\)M. This compound also exhibited improved or comparable activity compared to that of the reference drug against all the tested cell lines. Cell cycle analysis revealed that compound 39 induces apoptosis and G2/S arrest in MIA-Pa-Ca-2 cells. Compound 39 triggers mitochondrial potential loss in pancreatic cancer MIA-Pa-Ca-2 cells Furthermore, compound 39 triggers caspase-3 and PARP-1 cleavage, and this cleavage increases in a dose-dependent manner.

Podophyllotoxin (chemical formula: C_{29}H_{32}O_{8}, molecular weight: 414.41, \((5\text{-R,SaR,8aR,9R})-5,8,8a,9\text{-Tetrahydro-9-hydroxy}\text{-5-(3,4,5-trimethoxyphenoxy)furo}[3'4':6,7]naphtho[2,3-d]-1,3-dioxol-6(\text{SaH})\text{-one}\)) is a natural lignin and a natural product that is isolated from the roots of Podophyllum hexandrum growing in the wild. A series of new 4\(\beta\)-amidotriazole-linked podophyllotoxin derivatives was designed and synthesised by Reddy and et al.\(^{103}\). The majority of derivatives exhibited promising anti-proliferative activity with IC\(_{50}\) values ranging from 1 to 10 \(\mu\)M in six human cancer cell lines that included cervical (HeLa), breast (MCF-7), prostate (DU-145), lung (A549), liver (HepG2), and colon (HT-29) cancer lines. The structure activity relationship indicates that the aryl triazolic-amide derivatives shown higher cytotoxicity in comparison to their corresponding benzyl triazolic amide derivatives, and the strong electron withdrawing groups like chloro, trifluoromethyl at position-4 in aryl triazolic-amide derivatives exhibited superior
occurring oleane-type pentacyclic triterpene. A series of novel compounds were found to be more active than etoposide. Moreover, compound 40b exhibited remarkable cytotoxicity with IC50 values in the range of 0.70–4.11 µM, and this was the most promising compound in the series. For example, cervical (HeLa), breast (MCF-7), prostate (DU-145), lung (A549), liver (HepG2), and colon (HT-29) cancer cell lines were affected by compound 40b with IC50 values of 0.78, 0.97, 0.70, 1.20, 0.78, and 4.11 µM, respectively. Similarly, compound 40a exhibited IC50 values of 6.49, 1.10, 0.99, 1.61, 2.79, and 11.4 µM, while compound 40c exhibited IC50 values of 1.21, 1.35, 0.89, 1.96, 2.11, and 4.40 µM against these human cancer cell lines (HeLa, MCF-7, DU-145, A549, HepG2, and HT-29), respectively. The IC50 of etoposide when used as a standard drug ranged from 1.62 to 2.84 µM. Topoisomerase-mediated DNA relaxation assay results revealed that the derivatives could efficiently inhibit the activity of topoisomerase-II. Flow cytometry analysis of DU-145 cells revealed that these compounds cause arrest at the G2/M phase of the cell cycle. Further apoptotic studies were also performed on these DU-145 cells, and the results indicated that this class of compounds could effectively induce apoptosis.

Eugenol (Chemical formula: C10H12O2, molecular weight: 164.2, 2-Methoxy-4-(2-propen-1-yl)phenol) is the primary active component of clove oil (75%–90%)104. Taia et al.105 reported the synthesis of a new series of heterocyclic eugenol derivatives through the mixed condensation reaction of 1,3-dipolar azide and the oxide of p-chlorophenylazide on 4-allyl-2-methoxy-1-(prop-2-yn-1-ylxylo) benzene. The monoadducts and bis-heterocyclic derivatives were active against the fibrosarcoma cell line and against lung and breast carcinoma cell lines. The hybrid compound 41 (Figure 9) showed the highest anticancer activity against all tumour cell lines, with IC50 values between 15.31 and 23.51 µM.

Hederagenin (chemical formula: C30H48O4, molecular weight: 472.7, (3β,4α)-3,23-Dihydroxyolean-12-en-28-oic acid) is a naturally occurring oleane-type pentacyclic triterpene. A series of novel aryl-1H-1,2,3-triazol-4-yl methylester and amide derivatives of the natural product hederagenin was synthesised by Rodriguez-Hernandez and et al.106. The cytotoxic activities of all compounds were screened against a panel of six human cancer cell lines using the well-established photometric sulforhodamine B assay. The majority of the compounds displayed higher levels of antitumor activity than did the parent hederagenin. These results indicate that the presence of a bulky group bonded to carbonyl-28 of the triterpene skeleton modulates their cytotoxic activity. 1,2,3-Triazolyl compounds having an ester group are generally more cytotoxic than compounds carrying an amide moiety. The ester derivatives 42a, 42b, and 42c (Figure 9) that possess an m-bromo, m-chloro, and m-nitro substituent, respectively, were the most active compounds against all human cell lines tested, and they exhibited EC50 values ranging from between 3.2 and 4.0 µM for 42a, between 3.1 µM and 4.0 µM for 42b, and between 3.2 and 4.1 µM for 42c. These results revealed that these compounds (42a, 42b, and 42c) are at least eightfold more active than is the parent hederagenin; however, they were not selective between malignant and non-malignant cells. The ortho-fluorobenzyl-1,2,3-triazolyl ester 42d (Figure 9) was the most active compound against this cell line (EC50 = 1.6 µM). This compound (42d) also exhibited some selectivity in cytotoxicity (SI = 5.4; SI is defined as the quotient of EC50 values according to EC50 [NIH 3T3]/EC50 [tumour cell line]), discriminating between the cancer cell line HT29 and the non-malignant mouse fibroblast line NIH 3T3.

Oleanolic acid (chemical formula: C30H48O6, molecular weight: 456.71, (3β)-3-Hydroxyolean-12-en-28-oic acid) is a natural pentacyclic triterpenoid compound that is synthesised in many plants through the cyclisation of squalene107. A series of novel oleanolic acid-coupled 1,2,3-triazole derivatives was designed and synthesised by Wei and et al.108. The synthesised compounds were screened for anticancer activity against a panel of five human cancer cell lines using an MTT assay. A number of these compounds exhibited improved anticancer activity against the tested cancer cell lines compared to that of the positive controls 5-fluorouracil and oleanolic acid. Compounds with p-substitutions at an

Figure 10. The chemical structures of anti-inflammatory compounds 44–49.
aromatic ring are more active than corresponding compounds without substitutions or substitutions at an ortho- or meta-position. Compounds with electron withdrawing groups at an aromatic ring are generally more active than compounds without substitutions or substitutions with an electron donating group at the identical position. Compound 43 (Figure 9) possessed strong inhibitory activity against A375-S2 and HT1080 cells, with IC_{50} values of 4.97 and 3.51 μM. A series of pharmacology experiments revealed that compound 43 significantly induced HT1080 cell apoptosis. This compound can serve as a promising lead candidate for further study.

2.2. Anti-inflammatory activity

Maslinic acid (chemical formula: C_{30}H_{48}O_{4}, molecular weight: 472.7, (2α,3β/2,3-Dihydroxyolean-12-en-28-oic acid) is a natural pentacyclic triterpene. In the European diet, olive oil and table olives represent a significant source of maslinic acid. The introduction of triazole moieties into maslinic acid considerably improves its anti-inflammatory effect. The tested compounds from the series of tri-1,4-disubstituted triazoles were found to be the most potent among the synthesised compounds (% IL-1β production = 23–47; 30–100 μM). This finding showed the importance of the number and may be also the position of the triazole moieties to improve the anti-inflammatory activity of maslinic acid. Compounds 44a (m-Me) (Figure 10) and 44b (Figure 10) possessing a naphthyl group on the triazole ring exhibited relatively high activities (% IL-1β production = 23 ± 3 and 34 ± 3, respectively; 30 μM) compared to those of the remaining analogs and of maslinic acid (% IL-1β production = 109 ± 3; 30 μM).

Ursolic acid (chemical formula: C_{30}H_{48}O_{5}, molecular weight: 456.7, (3β/3-Hydroxypurslavone-12-en-28-oic acid) is a well-known pentacyclic triterpene that is one of the major active components of many traditional Chinese medicines. Most ursolic acid derivatives containing oxadiazole, triazole, and piperazine moieties exhibited pronounced anti-inflammatory effects at 100 mg/kg. Compound 45 (Figure 10) showed the most potent inhibitory activity against ear inflammation of all the synthesised compounds (69.76%), and this activity was higher than that of UA (57.67%), ibuprofen (25.17%) and indomethacin (26.83%) at 100 μg/kg (i.p.) and was one- and twofold more potent than were the standard drugs. The cytotoxicity of the compounds was assessed using the MTT assay, and no compounds exhibited any appreciable cytotoxic activity (IC_{50} >100 μmol/L), which was in contrast to ursolic acid. These results indicate that incorporation of a triazole moiety to ursolic acid can improve the anti-inflammatory activity, and the order of activity for the different carbon chain lengths was C_{3} > C_{4} > C_{2} > C_{5}.

Lappaconitine (chemical formula: C_{32}H_{44}N_{2}O_{8}, molecular weight: 584.7, (1a,14a,16b)-20-Ethyl-1,14,16-trimethoxyaconitane-4,8,9-triol 4-(2-acylamino)benzoate) is extracted from the roots of Aconitum sinomontanum Naka. A series of novel lappaconitine derivatives possessing various substituents at the 20-N position was designed and synthesised by Pang and et al. In the initial screening of lappaconitine derivatives against NO production, the target compounds exhibited excellent inhibitory ability relative to that of lappaconitine. In particular, compound 46 (Figure 10) exhibited the most potent inhibition, with an IC_{50} of 12.91 μmol/L. The elementary structure activity relationship of NO inhibitory activity indicated that replacement of the benzene ring with an electron donating group (4-(morpholine-4-yl) < 4-(4-phenylpiperazin-1-yl) < 4-(4-benzylpiperazin-1-yl) < 3,4,5-triOCH_{3} < 3,4-diOCH_{3} < 4-OCH_{3} < 4-(piperidin-1-yl)) could improve anti-inflammatory efficacy, and replacement of the benzene ring with an electron withdrawing group (4-CF_{3} < 4-F) could reduce anti-inflammatory efficacy. Furthermore, compound 46 exerts its anti-inflammatory activity by inhibiting NO, PGE2, and TNF-α generation via the suppression of NF-κB and MAPK signalling pathways. Notably, compound 46 could exert a significant therapeutic effect on LPS-induced acute lung injury (ALI) in vivo.

Chalcone-triazole derivatives not only possess anticancer activity, but also they exhibit anti-inflammatory activity. Boshra et al. synthesised new 2-hydroxylchalcone-triazole hybrids that possessed anti-inflammatory activity. The majority of the synthesised compounds exhibited anti-inflammatory activity that was equivalent to or even higher than that of celecoxib. The results showed that introducing electron donating group such as 3,4-dimethoxy or electron withdrawing group such as 4-Br to the phenyltriazole derivative improved the potency. Particularly, introducing lipophilic moiety as Cl or Br increases inhibitory potency as well as selectivity against COX-2. In order to investigate the effect of phenyltriazole moiety on activity, phenyl group was replaced by isatin to give compounds. Interestingly, some compounds of this series showed higher potency and selectivity for COX-2 than the corresponding phenyltriazole derivative. This improvement in potency of this series may be attributed to the synergistic effect of isatin moiety. An in vitro COX-1/COX-2 inhibition study revealed that among the synthesised compounds, compound 47 (Figure 10) exhibited the highest inhibitory activity against COX-2, with an IC_{50} value of 0.037 μM and a selectivity index of 359.46. Most of the compounds possessed significant in vitro 15-LOX inhibitory activity that was higher than that of zileuton. Therefore, compound 37 is a potent dual inhibitor of COX-2 and 15-LOX.

Coumarin-triazole derivatives not only possess antitumor activity, but also they exhibit anti-inflammatory activity. Lippeva et al. synthesised conjugates of coumarin possessing lupan triterpenoids and 1,2,3-triazole and tested their anti-inflammatory activity. Among them, compound 48 (Figure 10), a conjugate of lupine triterpenoid with furocoumarin oreoselone, possessed marked anti-inflammatory activity. This compound significantly reduced paw edema caused by the injection of histamine (edema index: 24.5%) to a level that was comparable to that of the nonsteroidal anti-inflammatory drug indomethacin. Among the hybrids of betulonic acid with the coumarin preurtheunicin, compound containing an alkyl triazole linker with a short three-unit alkyl chain exhibited a weak anti-inflammatory effect. This compound statistically significantly reduced the histamine-induced edoema; its effect was two times weaker than the effect of conjugate (48). The elongation of the alkyl chain of the linker led to a loss of anti-inflammatory activity.

Piperine (chemical formula: C_{17}H_{19}NO_{3}, molecular weight: 285.37, 5-(1,3-Benzodioxol-5-yl)-1-(1-piperidinyl)-2,4-pentadien-1-one) is a major alkaloid constituent of piper species, including Piper nigrum Linn and Piper longum Linn. This compound is commonly used in various traditional medicine systems. Nineteen novel piperine-based triazoles were synthesised using a click chemistry approach and were tested for in vivo anti-inflammatory activity. The most active compounds were evaluated for in vitro TNF-α expression. Compounds 49a and 49b (Figure 10) were found to exert significant in vivo inhibition of inflammation at levels of 80.40% and 76.71%, respectively, after 5 h in comparison to that caused by piperine (54.72%) and the standard drug indomethacin (77.02%) without causing any damage to the stomach. Compounds 49a and 49b suppressed TNF-α levels by 73.73% and 70.64%, respectively, and reduced the protein expression of
COX-2, NF-κB, and TNF-α to a greater degree than that caused by indomethacin. Moreover, compound 49a was found to exert a significant analgesic activity of 54.09%, and this was comparable to that caused by indomethacin (57.43%). The structure-activity relationship of the synthesised compounds has been analysed as follows. Presence of electron withdrawing groups on the aryl moiety decreased the in vivo anti-inflammatory activity as compared to the presence of electron donating groups. Better in vivo anti-inflammatory activity was observed for compounds having halogen at ortho position of the aromatic ring A as compared to halogen attached at para position. Increasing order of activity found was F < Cl < Br.

2.3. Antimicrobial activity

2.3.1. Antibacterial activity

Carvacrol (chemical formula: C_{10}H_{14}O, molecular weight: 150.22, 2-Methyl-5-(1-methylethyl)phenol) is a monoterpane-phenol that is found in the essential oil of many aromatic plants of the Lamiaceae family, including thyme and oregano. Aneja et al. designed and synthesised 1,2,3-triazole analogs of natural bioactive precursors. These compounds exhibited moderate to potent antibacterial activity against both Gram-positive and Gram-negative bacteria. The analogues bearing electron-donating substituents such as p-methyl and p-methoxy exhibited better...
inhibition of all bacterial strains in comparison to halogensubstituted (p-fluoro and p-chloro substitution) analogues, where the activity was considerably lost against all strains. Among all of these triazole analogs, compound 50 (Figure 11) (derived from carvacrol) that possesses carboxylic acid functionality emerged as a potent antibacterial agent against Streptococcus pneumoniae (IC50: 62.53 μg/mL), Enterococcus faecalis (IC50: 36.66 μg/mL), and Escherichia coli (IC50: 15.28 μg/mL). Furthermore, compound 50 also demonstrated moderate efficacy against multidrug-resistant Escherichia coli strains. Compound 50 in combination with ciprofloxacin displayed a synergistic effect on the multidrug-resistant Escherichia coli MRA11 and MRC17 strains. Growth kinetics studies performed on Streptococcus pneumoniae and Escherichia coli treated with compound 50 revealed an extended lag phase. TEM analysis indicated that compound 50 caused significant cell wall damage and membrane disruption in bacterial cells (Streptococcus pneumoniae and Escherichia coli), ultimately leading to cell death. Moreover, this compound was also found to be a potent anti-biofilm agent against Streptococcus pneumoniae and Escherichia coli strains and exhibited non-cytotoxic effects on a human embryonic kidney (HEK293) cell line up to a concentration of 100 μg/mL. Additionally, this compound did not cause an alteration in haemocyte density, indicating the lack of an immune response, and it was also non-toxic to the larvae of Galleria mellonella up to a concentration of 2.5 mg/mL.

Dehydroabietane-tetra-triazole derivatives not only possess anti-cancer activity59,69, but also they exhibit antibacterial activity118. Hou et al.118 obtained a diverse natural product-like synthetic abietane diterpenoid library containing 86 compounds. The structure-activity relationships showed that the introduction of 1,2,3-triazole ring at the C-14 position of dehydroabietic acid could retain the potent antibacterial activities. Generally, the introduction of electron-poor aromatic ring system on the C-4 position of the 1,2,3-triazole moiety would significantly increase the antibacterial activities. In particular, compounds 51a, 51b and 51c (Figure 11) showed promising low MICs against the tested bacteria. While, with the saturated alkyl-, ester- and trimethylsilyl- substituted groups on the C-4 position of the 1,2,3-triazole moiety there were no significant structure activity relationships. Among them, compound 51c exhibited the most potent activities against both gram-positive bacteria (Bacillus subtilis and Staphylococcus aureus) and gram-negative bacteria (Escherichia coli and Pseudomonas fluorescens) with comparable MICs (1.6 μg/mL to 3.1 μg/mL) to those observed in response to the positive control norfloxacin. More importantly, compound 51c exhibited very low cytotoxicity (CC50: 31.7 μg/mL and 41.8 μg/mL) against normal human foreskin fibroblast (HFF) and liver (HL-7702) cells.

Spiramycin (chemical formula: C18H38N2O16, molecular weight: 843.05, Spiramycin) is a natural antibiotic produced by Streptomyces ambofaciens in the form of a mixture of three compounds known as spiramycins I–III19,20. Huisgen cyclo-addition allowed for the acquisition of novel triazole-bridged antibiotics possessing the reconstructed C(5) arm of spiramycin21. Further comparison of the physicochemical parameters of these compounds possessing a relatively hydrophilic C(5) arm revealed good solubility for all of these derivatives. Antibacterial activity studies demonstrated that the most active among novel triazole conjugates is that containing the terminal N-acetylsaccharide moiety (compound 52). Compound 52 (Figure 11) possesses a lipophilicity that is similar to that of spiramycin, and this contributes to its generally comparable activity to that of spiramycin (is 2-4 less active than spiramycin [μM]). All of these results clearly show that 16-membered macroclide derivatives, even those not containing an aldehyde group but having additionally functionalised saccharide, can be active against Gram-positive bacteria at a level comparable to that of spiramycin.

Usnic acid (chemical formula: C18H18O7, molecular weight: 344.32, (9bR)-2,6-Diacetyl-7,9-dihydroxy-8,9-bimethyl-1,3(2H,9bHf)-dibenzo[3,4]furandione) is a secondary metabolite derived from lichen sources such as Usnia, Ramalina, and Cladonia22,123. Usnic acid enamine-coupled 1,2,3-triazole can be used as an antibacterial and antitubercular agent24. Among the synthesised derivatives, the most active analog (53a) (Figure 11) inhibited Mycobacterium tuberculosis (Mtbb) at an MIC value of 2.5 μM, and this was slightly better than that of the standard reference isoniazid (2.9 μM). Compounds 53b (3,4-difluorobenzoyl) (Figure 11) and 53c (2-acylnaphthyl) (Figure 10) proved to be effective and exhibited MIC values of 5.4 and 5.3 μM, respectively. In contrast, the uniqueness of the synthesised triazoles was observed for Bacillus subtilis, where compound 53d [3-(trifluoromethyl) phenacyl] (Figure 11) exhibited the greatest antibacterial efficacy (zone of inhibition of 15 mm) while also possessing moderate anti-tuberculosis activity. In addition to its anti-tuberculosis activity, compound 53e (N-acyl-morphololinyl) (Figure 11) also exhibited a 13 mm diameter inhibition zone against Bacillus subtilis. The active anti-tuberculosis compounds 53b (3,4-difluorobenzoyl) and 53c (2-acylnaphthyl) exert bacteriostatic effects on Bacillus subtilis, with inhibition zones of 12 and 11 mm, respectively. A structure–activity relationship assessment suggests that the presence of the fluorine atoms in 53b might be a contributing factor for the increased potency of the molecule. Naphthalene substituted triazole 53d displayed activity similar to that of 53b, indicating that the meta- and para-positions should be occupied for antitubercular activity. The presence of a p-halogen is important and favourable, but the presence of an m-halogen might be a deciding factor for increased potency. The presence of fluorne is more favourable than chlorine. The presence of an electron-donating methoxy group is unfavourable for antitubercular activity. The present study shows that a versatile bridging unit like triazole, when coupled to usnic acid, might be beneficial in imparting an antitubercular property.

Chalcone-triazole derivatives not only possess anticancer activity60,61,101,102 and anti-inflammatory activity114, but also they exhibit antibacterial activity125–127. The newly synthesised bis-1,2,3-triazole based chalcones were tested for their antibacterial activity125. The compounds demonstrated significant inhibition of tested gram positive and gram negative strains compared to the standard drug Gentamicin sulphate. The compounds 54a, 54b, 54c, 54d, 54e, and 54f (Figure 11) demonstrated high antibacterial activity at concentrations of 75 and 100 μg/mL. The accumulated data indicated that presence of the strong electron donating group (OMe) at ortho and para positions of phenyl ring of chalcone could enhance the antibacterial activity of the other products. In contrast, triazolyl-pterostilbene derivatives exhibited potent antibacterial activity, particularly against methicillin-resistant Staphylococcus aureus (MRSA)126. Among these compounds, compound 55 (Figure 11) exhibited the most potent anti-MRSA activity with a minimum inhibitory concentration (MIC) value of 1.2–2.4 μg/mL and a minimum bactericidal concentration (MBC) value of 19.5–39 μg/mL. Anti-MRSA mechanism studies indicated that active compounds may inhibit MRSA by acting on DNA polymerase instead of the bacterial cell wall and cell membrane. The structure-activity relationship indicated that the carboxylic acid must be the essential moiety in the antibacterial activity. And the spacer between triazole and carboxylic acid plays an important role in the inhibition of MRSA. Phenyl groups with the characteristics of planarity and resonance would be the most favourable.
A series of new dehydroacetic acid chalcone-1,2,3-triazole hybrids was designed, synthesised, and characterised for use as potential antimicrobial agents by Lal et al.\textsuperscript{127} Preliminary studies revealed that a number of the compounds exhibited bioactivities that were comparable to or even superior to those of the reference drugs. All of the triazole derivatives containing a substituted benzene ring displayed improved activity compared to that of dehydroacetic acid and dehydroacetic acid-chalcone alkynes, thereby highlighting the role of the 1,2,3-triazole moiety on the antimicrobial potential of the target compounds. Compounds containing bromo and methoxy groups on benzene ring exhibited better activity against most of the studied microorganisms. Compound 56 (Figure 11) was found to be the most active against \textit{Escherichia coli}, with an MIC value of 0.0030 µM/mL. Additionally, molecular modelling studies indicated that compound 56 binds effectively to the active sites of DNA gyrase via hydrogen bonding, hydrophobic interactions, and electrostatic interactions.

Coumarin-triazole derivatives not only possess anticancer activity\textsuperscript{58,68,70,90} and anti-inflammatory activity\textsuperscript{115}, but also they exhibit antibacterial activity\textsuperscript{128,129}. A new class of triazole-linked coumarinyl 2-mercaptobenzimidazole hybrids was synthesised and screened for their anti-tuberculosis potential against \textit{MTB H37Rv} by Anand et al.\textsuperscript{128} All conjugates exhibited promising anti-mycobacterial activity against \textit{MTB H37Rv} with MIC values ranging from 3.8 to 59.6 mM. In particular, the most active bis-substituted hybrids (57a and 57b) (Figure 11) (MIC: 3.8 µM) were found to be 2.4- to 6.6-fold more potent than were the references Pyrazinamide, streptomycin, and ciprofloxacin (MIC: 25.3, 10.7, and 2.4- to 6.6-fold more potent than were the references screened for their antibacterial activity. Compounds containing bromo and methoxy groups on benzene ring exhibited better activity against most of the studied microorganisms. Compound 56 (Figure 11) was found to be the most active against \textit{Escherichia coli}, with an MIC value of 0.0030 µM/mL. Additionally, molecular modelling studies indicated that compound 56 binds effectively to the active sites of DNA gyrase via hydrogen bonding, hydrophobic interactions, and electrostatic interactions.

Coumarin-triazole derivatives not only possess anticancer activity\textsuperscript{58,68,70,90} and anti-inflammatory activity\textsuperscript{115}, but also they exhibit antibacterial activity\textsuperscript{128,129}. A new class of triazole-linked coumarinyl 2-mercaptobenzimidazole hybrids was synthesised and screened for their anti-tuberculosis potential against \textit{MTB H37Rv} by Anand et al.\textsuperscript{128} All conjugates exhibited promising anti-mycobacterial activity against \textit{MTB H37Rv} with MIC values ranging from 3.8 to 59.6 mM. In particular, the most active bis-substituted hybrids (57a and 57b) (Figure 11) (MIC: 3.8 µM) were found to be 2.4- to 6.6-fold more potent than were the references Pyrazinamide, streptomycin, and ciprofloxacin (MIC: 25.3, 10.7, and 9.4 µM, respectively). Thus, hybrids 57a and 57b may become useful as new potential anti-tuberculosis agents in the future. Lipeeva et al.\textsuperscript{129} synthesised a new series of coumarinotriazol compounds. The synthesised coumarinotriazole-type derivatives were screened for their \textit{in vitro} antimicrobial activity. Compounds 58a and 58b (Figure 11) possessing a 4-(carboxyphenyl) triazolyl substituent at the 6 or 7 position of the coumarin ring exhibited excellent antibacterial activity against \textit{Staphylococcus aureus} strains, with MIC values of 0.16–3.75 µg/mL and 0.21–6.28 µg/mL, respectively. The coumarin-2,3-dihydrofurocoumarin hybrid compound 58c (Figure 11) was found to be selective against \textit{Bacillus subtilis} and \textit{Escherichia coli}, with MIC values of 0.02–0.15 µg/mL. A molecular docking study was performed for the most active compounds against the MurB protein, and the molecular docking results were consistent with the \textit{in vitro} antibacterial activity findings.

**Figure 12.** The chemical structures of antifungal compounds 59–60.

**Figure 13.** The chemical structures of anti-multiple timicrobial compounds 61–62.

### 2.3.2. Antifungal activity

Gossypol (chemical formula: C\textsubscript{30}H\textsubscript{30}O\textsubscript{8}, molecular weight: 518.55, 1,1’6,6’,7,7’-Hexahydroxy-3,3’-dimethyl-5,5’-bis[1-methylethyl][2,2’-binaphthalene]-8,8’-dicarboxaldehyde) is a yellow pigment that is present in various parts of cotton plants where it acts as a plant defense system against pathogenic fungi and insects\textsuperscript{130}. This compound is a natural bisesquiterpene. Pyta et al.\textsuperscript{131} designed and synthesised novel gossypol triazole conjugates that were functionnalised with aliphatic chains and benzoyloxy groups. Biological evaluation of the new gossypol-triazole conjugates revealed that the potency of the 59a and 59b derivatives (Figure 12) possessing triazole-benzoyloxy moieties was comparable to that of miconazole against \textit{Fusarium oxysporum} (MICs = 16 µg/mL). The results of HPLC evaluation of ergosterol content in different fungal strains upon treatment with gossypol and its derivatives indicate that the mechanism of antifungal activity of gossypol and its triazole-containing derivatives may be involve the inhibition of ergosterol biosynthesis, as this process is crucial for controlling the permeability and fluidity of fungal plasma membranes in a manner similar to that of cholesterol in animals.

Carnosol (chemical formula: C\textsubscript{20}H\textsubscript{26}O\textsubscript{4}, molecular weight: 518.55, 4aR,9S,10aS)-1,3,4,9,10a-Hexahydro-5,6-dihydroxy-1,1-dimethyl-7-(1-methylethyl)-2H-9,4a-epoxymethano]phenanthren-12-one) is one of the major phenolic diterpenes derived from the leaves of \textit{Rosmarinus officinalis}. L. Pertino et al.\textsuperscript{132} used click chemistry to synthesise a series of twenty-four novel abietane diterpene derivatives with good to reasonable yields. The antifungal activity of the compounds was determined as the percentage of inhibition of \textit{Candida albicans} ATCC 10231 and \textit{Cryptococcus neoformans} ATCC 32264 in the range of 250–3.9 µg/mL. From these data, the MIC\textsubscript{100} and MIC\textsubscript{50} were determined for all of the synthesised compounds. The most active carnosol derivative was the p-bromobenzyl derivative 60, which reduced the growth of \textit{Cryptococcus neoformans} by about 91.3% at 250 µg/mL while compound, with a p-nitrobenzyl unit decreased fungal growth by about 71% at the same concentration. The results indicate some selectivity for the different fungi and that the placement of the lactone (either C-20, C-11 or C-20, C-7) is important for the effect.
Candida utilis were highly active against Candida albicans. Most of the compounds exhibited very good antifungal activity. Pseudomonas aeruginosa, Bacillus subtilis, and Bacillus cereus, with MIC values of 8 μg/mL and 16 μg/mL, respectively, possessed chloro and methoxy substitution in coumarin and 1-aza coumarin (Figure 13) were found to be very active against Candida albicans, Candida krusei, and Candida tropicalis, with MIC values of 1 μg/mL, respectively. Compounds possessing chloro and methoxy substitution in coumarin 62 (Figure 13) were found to be very active against Staphylococcus aureus, Candida albicans, Candida utilis, and Candida krusei, with an MIC of 1 μg/mL that was similar to that of the standard. This compound also exhibited good activity against Bacillus subtilis, Bacillus cereus, and Bacillus bronchiseptica, with MIC values of 8 μg/mL and 16 μg/mL, respectively. The majority of the synthesised hybrids also exhibited good antifungal activity. Compound 62 possessed higher potency compared to that of fluconazole (MIC = 0.0102 μM/mL) against A. niger and C. albicans, with an MIC value of 0.0032 μM/mL. Structure activity relationship: i) Most of the fluorinated triazoles exhibited good results than the non-fluorinated compound. ii) The synthesised triazole analogues with a substituted benzene showed better activity than naphthaldehyde-chalcone alkynes, exhibiting the significance of 1,2,3-triazole. These outcomes revealed the additive effect of biological activity when two pharmacophoric moieties, i.e. chalcone and 1,2,3-triazole are conjugated. iii) Compounds with electron withdrawing substituents on benzene displayed superior activity than having electron releasing groups. iv) It was observed that compounds with nitro and methoxy substituents on benzene ring demonstrated better activity against majority of the microorganisms under test. v) Molecule 62 containing p-nitro group exhibited good antifungal activity and was more effective than Fluconazole. vi) Activity results also revealed that most of the triazole hybrids exhibited superior antifungal potency compared to antibacterial activity.

### 2.3.3. Miscellaneous

Coumarin-triazole derivatives not only possess anticancer activity, anti-inflammatory activity, and antibacterial activity, but also they exhibit antifungal activity. A series of new coumarin-, quinolinone-, and benzyl-linked 1,2,3-triazole derivatives were synthesised and screened for their antibacterial and antifungal activities by Savanur and et al. Most of the compounds exhibited good activity against the Gram-positive bacteria Staphylococcus aureus and Bacillus subtilis and the Gram-negative bacteria Pseudomonas aeruginosa. Almost all of the compounds exhibited very good antifungal activity. Most of them were highly active against Candida albicans, Candida tropicalis, Candida utilis, and Candida krusei and exhibited moderate activity against A. fumigatus, A. niger, R. oryzae, and R. bataticola. Compounds possessing chloro and methoxy substitution in coumarin (61a) (Figure 13) were found to be very active against Staphylococcus aureus, Pseudomonas aeruginosa, Candida albicans, Candida utilis, and Candida krusei, with an MIC of 1 μg/mL that was similar to that of the standard. This compound also exhibited good activity against Bacillus subtilis, Bacillus cereus, and Bacillus bronchiseptica, with MIC values of 8 μg/mL and 16 μg/mL, respectively. The majority of the synthesised hybrids also exhibited good antifungal activity. Compound 62 possessed higher potency compared to that of fluconazole (MIC = 0.0102 μM/mL) against A. niger and C. albicans, with an MIC value of 0.0032 μM/mL. Structure activity relationship: i) Most of the fluorinated triazoles exhibited good results than the non-fluorinated compound. ii) The synthesised triazole analogues with a substituted benzene showed better activity than naphthaldehyde-chalcone alkynes, exhibiting the significance of 1,2,3-triazole. These outcomes revealed the additive effect of biological activity when two pharmacophoric moieties, i.e. chalcone and 1,2,3-triazole are conjugated. iii) Compounds with electron withdrawing substituents on benzene displayed superior activity than having electron releasing groups. iv) It was observed that compounds with nitro and methoxy substituents on benzene ring demonstrated better activity against majority of the microorganisms under test. v) Molecule 62 containing p-nitro group exhibited good antifungal activity and was more effective than Fluconazole. vi) Activity results also revealed that most of the triazole hybrids exhibited superior antifungal potency compared to antibacterial activity.

### 2.4. Antiparasitic activity

#### 2.4.1. Anti-leishmanial activity

Eugenol-triazole derivatives not only possess anti-cancer activity, but also they exhibit anti-leishmanial activity. Eugenol derivatives exert leishmanicidal activities with varying degrees of effectiveness. The most active compound, 4-[(3-(4-allyl-2-methoxyphenoxypropyl)-1-(4-methylbenzyl)-1H-1,2,3-triazole (63) (Figure 14) (IC50 = 7.4 ± 0.8 μM/L), also targeted Leishmania parasites inside peritoneal macrophages (IC50 = 1.6 μM/L) without interfering with cell viability. The cytotoxicity of compound 63 against macrophage cells was indicated by an IC50 of 211.9 μM/L and a selective index of 132.5. Under similar conditions, compound 52 was more effective than were glucantime and pentamidine, two drugs currently used in the clinic. Theoretical calculations indicated that this compound also exerts the majority of its physicochemical and pharmacokinetic activities within the ranges expected for orally available drugs. It is believed that eugenol bearing 1,2,3-triazole functionalities may represent a scaffold to be explored towards the development of new agents to treat leishmaniasis.
Hederagenin-triazole derivatives not only possess anti-cancer activity\(^{106}\), but also they exhibit anti-leishmanial activity\(^{136,137}\). A series of hederagenin derivatives synthesised by Rodríguez-Hernández et al.\(^{136}\) exerted highly potent anti-leishmanial effects\(^{136}\). Some derivatives possessed activity at the micromolar level and exhibited low toxicity against BGM and HepG2 cells. Moreover, the ability of the hederagenin derivatives \(64a\) (11 \(\mu\)M) and \(64b\) (2 \(\mu\)M) (Figure 13) to prevent the proliferation of intracellular amastigote forms of \textit{Leishmania infantum} and their higher selectivity index and low toxicity compared to commercial positive control drugs (potassium antimonyl tartrate trihydrate) (IC\(_{50}\) = 80 \(\mu\)M, SI = 0.1) make these compounds promising candidates for the treatment of leishmaniasis. Although a considerable number of hederagenin derivatives, carrying or not a triazol unit, bonded with an aromatic system with several electron donating and withdrawing groups at different positions, have been prepared, no structural-activity relation could be clearly established. Rodríguez-Hernández et al.\(^{137}\) also found that derivatives \(65a\), \(65b\), and \(65c\) (Figure 13) were highly effective in preventing the proliferation of intracellular amastigote forms of \textit{Leishmania infantum} (IC\(_{50}\) = 25.9, 5.6, and 7.4 \(\mu\)M, respectively). All of these compounds exhibited a higher selectivity index and a low toxicity compared to two strains of kidney (BGM) and liver (HepG2) cells. Compound \(65b\) possessed a higher selectivity (1780 times) in comparison to that of the commercial drugs and is approximately 8 times more selective than is the most active compound previously reported as a commercial antimony drug. Moreover, \(65a\), \(65b\), and \(65c\) (Figure 13) exhibited antimalarial activity\(^{128,129}\) and antifungal activity\(^{135}\), but also they exhibit antimalarial activity\(^{140}\). The results show that most of the new compounds of coumarin-triazole series exhibited very promising activity ranges, depending on the substitution on position-1 of triazole ring, like alkyl chain having polar group and phenyl ring having different functional group. The presence of electron releasing group and electron withdrawing group, their position (ortho, meta and para) in phenyl ring and their steric hindrance, all the factors might affect the activity, but varying behaviour of compounds with similar nature of functional groups is still elusive. Among these compounds, compound \(67\) (7-[1-(2, 4-dimethoxyphenyl)-1H-[1–3] triazol-4-ylmethyl]-4-methyl-chromen-2-one) (Figure 15) having 2, 4-dimethoxy phenyl ring was the most active, with an IC\(_{50}\) value of 0.763 ± 0.0124 \(\mu\)g/mL, presence of two electron releasing methoxy group at ortho and para position of phenyl ring enhanced the antimalarial activity quite significantly. Convincing results were obtained by analysing their inhibitory effect on the supercoiling activity of the enzyme gyrase through assays that employed the use of \textit{Escherichia coli} DNA gyrase and

**2.4.2. Antimalarial activity**

Quinine (chemical formula: C\(_{20}\)H\(_{24}\)N\(_2\)O\(_2\), molecular weight: 324.42, (8z,9R)-6’-Methoxycinchonan-9-ol) is the most abundant Cinchona alkaloid and was the only known antimalarial drug for over 300 years\(^{138}\). Faidallah et al.\(^{139}\) synthesised a variety of 1,2,3-triazole-quinine conjugates. The synthesised compounds were bioassayed against the blood stage of \textit{Plasmodium falciparum} strain 3D7 according to the \textit{in vitro} standard procedure. All the amino acid and dipeptide analogs were found to be less potent than quinine. Some of the aryl and heteroaryl analogs showed more promising submicromolar potency. Upon inserting a second methylene between quinine and triazole moieties, the potency dramatically improved. A number of the synthesised analogs (IC\(_{50}\) = 43, 37, 41, 40, 30, and 27 nM for compounds \(66a\), \(66b\), \(66c\), \(66d\), \(66e\), and \(66f\) [Figure 15], respectively) exhibited antimalarial properties with higher potency than that of the starting precursor quinine (standard reference used). This may be attributed to the lipophobic/hydrophilic properties of long alkyl chain containing compounds.

Coumarin-triazole derivatives not only possess anticancer activity\(^{28,68,70,90–93}\), anti-inflammatory activity\(^{115}\), antibacterial activity\(^{128,129}\), and antifungal activity\(^{135}\), but also they exhibit antimalarial activity\(^{140}\). Coumarin-triazole analogs possess antiplasmodial efficacy \textit{in vitro}\(^{140}\). The results show that most of the new compounds of coumarin-triazole series exhibited very promising activity ranges, depending on the substitution on position-1 of triazole ring, like alkyl chain having polar group and phenyl ring having different functional group. The presence of electron releasing group and electron withdrawing group, their position (ortho, meta and para) in phenyl ring and their steric hindrance, all the factors might affect the activity, but varying behaviour of compounds with similar nature of functional groups is still elusive. Among these compounds, compound \(67\) (7-[1-(2, 4-dimethoxyphenyl)-1H-[1–3] triazol-4-ylmethyl]-4-methyl-chromen-2-one) (Figure 15) having 2, 4-dimethoxy phenyl ring was the most active, with an IC\(_{50}\) value of 0.763 ± 0.0124 \(\mu\)g/mL, presence of two electron releasing methoxy group at ortho and para position of phenyl ring enhanced the antimalarial activity quite significantly. Convincing results were obtained by analysing their inhibitory effect on the supercoiling activity of the enzyme gyrase through assays that employed the use of \textit{Escherichia coli} DNA gyrase and

![Figure 15](image-url) The chemical structures of antimalarial compounds 66–71.
relaxed plasmid DNA. Thus, these compounds can be used as potential agents to synthesise new antimalarial drugs that target the DNA gyrase enzyme.

Berberine-triazole derivatives not only possess anticancer activity, but also they exhibit antimalarial activity. Novel sulphonamide-based berberine–1,2,3-triazole hybrids were successfully synthesised by Batra et al. The majority of the synthesised compounds possessed significant antimalarial activity, with IC50 values in the range of 0.1–20 μg/mL. These compounds were also found to be non-cytotoxic under the tested conditions. The majority of synthesised sulphonamide-based berberine-triazole hybrids are proved to be significantly active against Plasmodium falciparum. Among the halogen-substituted derivatives, compound 68 (Figure 15) containing a p-chlorophenylamino substituent was determined to be the most active molecule, with an IC50 of 0.1 μg/mL.

Kaurenoic acid (chemical formula: C20H30O2, molecular weight: 302.455, 4α)-Kaur-16-en-18-0ic acid) is a widely-occurring diterpenoid. Santos et al. reported the synthesis of hybrid kaurenoind molecules of type 1,2,3-triazole–1,4 disubstituted with the aim of improving the antimalarial activity of kaurenoic and xylpic acids. A total of nine novel hybrid kaurenoind–1,2,3-triazole derivatives were obtained via the CuAAC reaction, although the products were much less active than were the starting diterpene acids. Derivative 69 (Figure 15) possessed the highest selective index (SI = 22.2). Chloroquine exhibited a CC50 value of 543.6 ± 71.4 μM, and the SI was 1308. Derivative 69 possessed CC50 values (> 2102.3 μM) that were greater than those of chloroquine. Specifically, the cytotoxicity of these substances as tested in HepG2 cells was significantly lower than that of chloroquine. However, the SI value of chloroquine was higher than the values found for derivative 69 due to its higher activity. Consequently, its IC50 is much lower. In this case, the presence of nitrogen groups, and possibly of protonated forms, would certainly influence in receptors interaction and, consequently, in the antimalarial effect.

Chalcone-triazole derivatives not only possess antitumor activity, but also they exhibit antimalarial activity, and antibacterial activity. Kant et al. attempted to synthesise a new class of triazole chalcone derivatives possessing antimalarial activity through the use of copper-catalyzed click chemistry. A number of these compounds exhibited moderate activities. The most potent compound was compound 70 (Figure 15) that possessed a 3-chloro-4-fluoro substituted benzene ring, as this compound exhibited an IC50 of 2.74 μg/mL in vitro against the erythrocytic stages of Plasmodium falciparum (3D7 strain). These compounds were also evaluated for cytotoxicity in vitro against the Huh-7 cell line, and they exhibited no cytotoxic activity and CC50 values that were higher than 100 μg/mL. This is a preliminary result and to reach more appropriate conclusion 2nd and 3rd generation compounds should be synthesised in order to establish meaningful structure activity relationship. Chalcone-quinoline hybrids possessing aminoethylene and aminopropylene linkers between quinoline and 1,2,3-triazole moieties also showed potential in vitro antimalarial activities against the CQR W2 strain of Plasmodium falciparum. The structure activity relationship results demonstrated that the shorter aminoethylene linker between quinoline and 1,2,3-triazole motifs was more potent than the longer aminopropylene linker between the quinoline and 1,2,3-triazole motifs. The most active hybrid (71) (Figure 15) with an IC50 of 114.1 nM against the CQR W2 strain of P. falciparum was not inferior to chloroquine (IC50: 150 nM), and it (CC50: 35.6 μM) also showed low cytotoxicity towards HeLa cells and exhibited a good selectivity index (SI: 311).

### 2.4.3. Anti-Toxoplasma gondii activity

Arctigenin (chemical formula: C37H41O6, molecular weight: 372.41, (3(R,4R)-4-[(3,4-Dimethoxyphenyl)methyl]dihydro-3-[(4-hydroxy-3-methoxyphenyl)methyl]-2(3H)-furanone) is derived from the dried ripe fruit of Arctium lappa. Four new series of arctigenin derivatives were designed, synthesised, and evaluated for their anti-Toxoplasma gondii activity in vitro and in vivo by Zhang and et al. For the different substituted phenyl or benzyl 1,2,3-triazole compounds, the introduction of electron-withdrawing groups, such as halogens, at the ortho and meta positions of the benzene ring enhanced the ability of HeLa cells to resist Toxoplasma gondii. However, the introduction of a halogen at the para position of the benzene ring was detrimental to the increase of anti-Toxoplasma gondii activity. On the other hand, the anti-Toxoplasma gondii ability was enhanced after the introduction of electron-donating groups to the para position of the benzene ring. Among the synthesised compounds, 4–(3,4-dimethoxybenzyl)–3–(4–(1-(2-fluorobenzyl)-1H,1,2,3-triazol-4-yl) methoxy)-3-methoxybenzyl)dihydrofuran-2(3H)-one (72) (Figure 16) exhibited the most potent anti-Toxoplasma gondii activity and a low cytotoxicity (IC50 in Toxoplasma gondii: 17.1 μM; IC50 in HeLa cells: > 600.0 μM; Selectivity: 35.09), thus yielding improved results compared to those of the lead compound arctigenin (IC50 in Toxoplasma gondii: 586.4 μM; IC50 in HeLa cells: 572.7 μM; Selectivity: 0.98) and the clinically applied positive-control drug spiramycin (IC50 in Toxoplasma gondii: 262.2 μM; IC50 in HeLa cells: 189.0 μM; Selectivity: 0.72). Compound 72 not only significantly reduced the number of tachyzoites in the peritoneal cavity of mice, but also it resulted in their partial malformation (p < 0.05) in vivo. Additionally, the results of a docking study of compound 72 into the Toxoplasma gondii calcium-dependent protein kinase 1 (TgCDPK1) receptor protein-binding site revealed that its mode of action was possibly as a TgCDPK1 inhibitor.

Ursolic acid-triazole derivatives not only possess anti-inflammatory activity, but also they exhibit anti-Toxoplasma gondii activity. Ursolic acid derivatives displayed some anti-Toxoplasma gondii activity and exhibited less cytotoxicity than ursolic acid.
in vitro \(^{146}\). The compounds with 1,2,4-phenyltriazole showed considerably higher anti-Toxoplasma gondii activity. It seems that the anti-Toxoplasma gondii ability was enhanced after the introduction of strong electron withdrawing group (–F, –NO\(_2\)) to the para position of the benzene ring. Compound 73 (Figure 16) exhibited the most potent anti-Toxoplasma gondii activity in vivo (Selectivity: 1.77) and was superior to ursolic acid (selectivity: 0.62) and the positive control spiramycin (Selectivity: 0.72). Additionally, determination of biochemical parameters, including the liver and spleen indexes, indicated that compound 73 effectively reduced hepatotoxicity and significantly enhanced anti-oxidative effects compared to those values in response to ursolic acid. Furthermore, a molecular docking study indicated that compound 73 possesses a strong binding affinity for Toxoplasma gondii calcium-dependent protein kinase 1.

Usnic acid-triazole derivatives not only possess antibacterial activity \(^{124}\), but also they exhibit anti-Toxoplasma gondii activity \(^{147}\). Six series of (+)-usnic acid derivatives were synthesised by Guo et al. \(^{147}\). In vitro, the majority of the derivatives tested in this study exhibited more anti-Toxoplasma gondii activity than did the parent compound (+)-usnic acid and the positive control drugs. Compounds introduced different substituted aminotriazoles into the lead (+)-usnic acid, and the sequence of the selectivity index was as follows: p-Br > p-CF\(_3\) > p-F > o-CI > o-OCH\(_3\) > p-OCH\(_3\) > p-H > p-Cl. To some extent, this suggests that for the anti-Toxoplasma gondii activity it may be beneficial to introduce strong electron withdrawing groups in the para position of the benzene ring. The selectivity index of the triazole-linked (+)-usnic acid derivatives was 0.41–1.34. Among them, derivative 74 (Figure 16) exhibited the best anti-Toxoplasma gondii activity (selectivity: 1.34), and this activity was greater than that of the positive control drugs sulfadiazine (selectivity: 1.15), pyrimethamine (selectivity: 0.89), and spiramycin (selectivity: 0.72) and the lead compound (+)-usnic acid (selectivity: 0.96).

2.4.4. Anti-Trypanosoma cruzi activity

Eugenol-triazole derivatives not only possess anticancer activity \(^{135}\) and anti-leishmanial activity \(^{135}\), but also they exhibit anti-Trypanosoma cruzi activity. \(^{148}\) De Souza et al. \(^{148}\) reported the synthesis of 1,2,3-triazoles obtained from eugenol and di-hydroeugenol and their \textit{in vitro} and \textit{in vivo} tryp路演idal activity. Compound 75 exhibited the highest activity against the epimastigote forms of \textit{Trypanosoma cruzi} (Y strain) \((IC_{50} = 42.8 \mu M)\) and were weakly toxic to cardiomyoblast cells (H9c2 cells), similarly to that presented by the control drug benznidazole. Although this derivative had a lower SI than benznidazole, it can be considered an innovative structural core for optimisation and design of new tripanocidal agents. Moreover, the triazole 75 was twice as active as the corresponding eugenol derivative \((IC_{50} = 88.4 \mu M)\) pointing the importance of the \textit{n}-propyl side chain for this activity. It is possible to note among these 1,2,3-triazoles that the phenyl group was the best substituent at the triazole core, because derivatives with hydroxymethyl, acetyl or cyclohexyl groups showed lower or no tripanocidal activity. Compounds 75 (Figure 17) could reduce greater than 50% of parasitemia after a 100 mg/kg oral treatment in mice infected with \textit{Trypanosoma cruzi}. Molecular docking studies suggested that this compound could act as a trypanocidal agent by inhibiting cruzain, an essential enzyme for \textit{Trypanosoma cruzi} metabolism that is typically inhibited by triazole compounds.

2.4.5. Miscellaneous

Quinine-triazole derivatives not only possess antimalarial activity \(^{139}\), but also they exhibit antileishmanial activity \(^{149}\). Sahu et al. \(^{149}\) synthesised a series of quinine-triazole molecular hybrids. From the explored biological and toxicological evaluation it was observed that addition of triazole moiety to quinine resulted in reduction of toxicity of the conjugated scaffold. Out of the varied substituents linked to the parent quinine-triazole scaffold it was found that the tertiary amines linked compounds were relatively more potent than both the reference standards. On increasing the chain length of the compounds the potency of the compounds also exhibited an increasing trend. This may be attributed to the increased penetrating power rendered to the pharmacophoric lead due to long alkyl chain substitution of the parent scaffold. Among them, compounds 76a, 76b, 76c, 76d, and 76e (Figure 18) possessed significant antimalarial \((Plasmodium falciparum)\) and antileishmanial activities \((Leishmania donavani)\), with \(IC_{50}\) values of 0.28, 0.28, 0.25, 0.33, 0.76 \(\mu M\) and 8.26, 4.4, 1.78, 3.95, and 4.06 \(\mu M\), respectively. Further toxicological analyses established the median lethal dose (LD50), no observed adverse effect level (NOAEL), and human equivalent dose (HED) of the most potent compounds by acute and subacute toxicity studies performed in rodent animal models. The studies revealed that these compounds 76a, 76b, 76c, and 76d did not cause any measurable toxicity at a dose of 1000 mg/kg, and based on this, the corresponding HED was calculated to be 13.84 mg/kg.

Veraguensin (chemical formula: \(C_{22}H_{28}O_{5}\), molecular weight: 372.45, \((2R,3S,4R,5S)-2,5\)-Bis\((3,4\)-dimethoxyphenyl)\)tetrahydro-3,4-dimethylfuran), grandisin (chemical formula: \(C_{24}H_{32}O_{7}\), molecular weight: 356.41, \((3,4\)-Dimethoxyphenyl)tetrahydro-3,4-dimethylfuran), and machilin G (chemical formula: \(C_{21}H_{24}O_{5}\), molecular weight: 338.45, \((3,4\)-Dimethoxyphenyl)tetrahydro-3,4-dimethyl-2-furanyl\)>-5-(3,4-Dimethoxyphenyl)tetrahydro-3,4-dimethyl-2-furanyl\)>-1,3-benzodioxole) are three of the sixteen 1,4-diaryl-1,2,3-triazole compounds derived from the natural products veraguensin, grandisin, and machilin G that were previously synthesised. \(^{150}\) Biological activity tests against \textit{L. amazonensis} promastigotes revealed that compounds 77a, 77d, and 77b (Figure 18) were the most active, with maximum inhibitory concentration \((IC_{50})\) values of 1.1, 3.71, and 7.23 \(\mu M\) that were higher than that of pentamidine \((IC_{50} = 8.9 \mu M)\). Compound 77a, an analog of machilin G, 3, was eightfold more active than pentamidine, while 77d and 77b were 2.5- and 1.2-fold more active, respectively, than was pentamidine. Compounds 77a, 77d, and 77b exhibited low cytotoxicity and possessed high selectivity indexes (SI) that are tens to hundreds of times higher than those of the recommended drugs for leishmaniasis, such as pentamidine and amphotericin B (SI 8.8 and 8.2, respectively) and for \textit{Trypanosoma cruzi}, such as benznidazole (SI 13.2). Compound 77b was highly active against \textit{Leishmania infantum}, with an \(IC_{50}\) value of 5.2 \(\mu M\), and derivative 77c (Figure 18) exhibited an \(IC_{50}\) value of 28.6 \(\mu M\) against \textit{Trypanosoma cruzi} trypanomastigotes. Regarding SAR, hybrid 1,2,3-triazole compounds 77a and 77c and positional isomers 77b and 77d, containing the methylenedioxy group present in machilin G, were the most active against the trypanosomatids, indicating that this group is
responsible for the high antileishmanial activity and moderate antitrpanosomal activity of these compounds.

2.5. Antiviral activity

Camphor (chemical formula: C₁₀H₁₆O, molecular weight: 152.23, 1,7,7-Trimethylbicyclo[2.2.1]heptan-2-one) is a member of the terpenoid family. Artushin et al. synthesized six new terpene (+)-camphor and alkaloid (+)-cystine conjugates. Among them, imino-derivatives were, in general, of higher activity. Elongation of linker attached to the camphor moiety and introduction of bulky lipophilic groups decreased virusinhibiting activity. Conjugate 78 (Figure 19) that contains a cisvynine fragment separated from the triazole ring by a C₆H₁₂– aliphatic linker exhibited the highest activity at relatively low toxicity (values of 50% cytotoxic dose CC₅₀=168 μmol, 50% inhibition dose IC₅₀=8 μmol, selectivity index SI = 20). Its selectivity index appeared to be higher than that of the reference compound rimantadine.

Baicalein (chemical formula: C₁₅H₁₀O₅, molecular weight: 270.24, 5,6,7-Trihydroxy-2-phenyl-4H-Camphor (chemical formula: C₁₀H₁₆O, molecular weight: 152.23, 2.5. Antiviral activity antitrypanosomal activity of these compounds.

Figure 18. The chemical structures of anti-multiple parasite compounds 76–77.

Figure 19. The chemical structures of antiviral compounds 78–79.

2.6. Antioxidant activity

L-ascorbic acid (chemical formula: C₆H₈O₆, molecular weight: 176.12, Vitamin C) exists in numerous types of fresh vegetables and fruits. The novel 4-substituted 1,2,3-triazole L-ascorbic acid conjugates possessing a hydroxyethylene spacer and their conformationally restricted 4,5-unsaturated analogs were synthesised as potential antioxidant agents by Harej et al. An evaluation of the antioxidant activity of these novel compounds revealed that the majority of the 4,5-unsaturated L-ascorbic acid derivatives exhibited improved antioxidant activity compared to that of their saturated counterparts. m-Hydroxyphenyl (80a), p-pentylphenyl (80b), and 2-hydroxyethyl (80c) substituted 4,5-unsaturated 1,2,3-triazole L-ascorbic acid derivatives exhibited highly efficient and rapid (within 5 min) 2,2-diphenyl-1-picrylhydrazyl (DPPH) radical scavenging activity (80a: IC₅₀ = 0.06 M/mL; 80c: IC₅₀ = 0.07 mM). The structure activity analysis through principal component analysis indicated radical scavenging activity by the participation of OH group with favourable reaction parameters: the C₃-OH group of saturated C₄-C₅(OH) derivatives and the C₂-OH group of their unsaturated C₄-C₅ analogues.

Coumarin-triazole derivatives not only possess anticanccer activity, anti-inflammatory activity, antibacterial activity, antifungal activity, and antimalarial activity, but also they exhibit antioxidant activity. Coumarin-based 1,4-disubstituted 1,2,3-triazole derivatives were synthesised by Dharavath et al. All of the synthesised compounds were screened for their inhibitory activity on neutrophils and lymphocytes in the BALF of RVS-infected mice. In summary, compound 79a inhibits RSV-infection and prevents pulmonary airway inflammation through the activation of the IFN signalling pathway. Structure activity relationship analysis revealed that all the four compounds 79a, 79b, 79c and 79d which demonstrated good activity against RSV induced infection contained substituents in the ortho-position. The substituents present in the ortho-position of compounds 79a, 79b, 79c and 79d are floro, trifluoromethyl, nitrite and bromo, respectively. It appears that these compounds exhibit inhibitory effect on RSV infection by interacting with the proteins containing active site that can accommodate only ortho substituted compounds.
2.7. Anti-Alzheimer’s activity

Sarsasapogenin (chemical formula: C27H44O3, molecular weight: 416.64, (3β,5β,25S)-Spirostan-3-ol) is an active ingredient in *Rhizoma anemarrhenae*. Wang et al. designed and synthesised a novel series of sarsasapogenin-triazolyl hybrids and evaluated their Aβ1-42 aggregation inhibitory activities. The result showed that the potent inhibitory activity exhibited by the benzyl groups series of compounds was possibly due to their steric effects and the ability of the benzyl groups forming pi-pi stacking interactions with the aromatic residues of Aβ. Among them, the most potent compound was 82a (IC50 = 5.84 µM) (Figure 21), and this compound exhibited an inhibition ratio of 84.74%, which was approximately 1.5- and 5.2-fold higher than that of curcumin (55.87%). Compound 82b (IC50 = 8.28 µM) (Figure 21) containing a triazole moiety also exhibited good Aβ1-42 aggregation inhibitory activity that was improved compared to that of the standard drug curcumin (IC50 = 14.99 µM). Moreover, 82a and 82b exhibited moderate neuroprotective effects against H2O2-induced neurotoxicity in SH-SY5Y cells. Oral treatment with 82a and 82b significantly ameliorated cognitive impairments in behavioural tests, and TUNEL staining revealed that 82a and 82b attenuated neuronal loss in the brain.

Coumarin-triazole derivatives not only possess anticancer activity, anti-inflammatory activity, antibacterial activity, antifungal activity, antimalarial activity, and antioxidant activity, but also they exhibit anti-Alzheimer’s activity. In 2016, Torres et al. screened the *in vitro* acetylcholinesterase inhibitor activity of a novel series of 1,2,3-triazole-linked coumarin hybrids. The most prominent compound was coumarin 83 (triazole and 4-methylcoumarins) (Figure 21), and this compound inhibited nearly 60% of the acetylcholinesterase activity at a concentration of 200 µmol/L. However, docking simulations suggest that this compound binds similarly to donepezil and, consequently, this compound can putatively block the secondary non-chelating functions of the enzyme, including adhesion, differentiation, and deposition of beta-amyloid in Alzheimer’s disease. Therefore, can be explored in 4-methylcoumarins/1,2,3-triazoles conjugates to yield multitarget ligands in the search of new compounds for the treatment of Alzheimer’s disease. The same year, Park et al. designed and synthesised new triazole-linked decursinol derivatives possessing potent inhibitory activities against cholinesterase (acetylcholinesterase [AChE] and butyrylcholinesterase [BuChE]). Compound 84 (IC50 = 5.89 ± 0.31 mM against BuChE) (Figure 21) possessed more effective inhibitory activity against BuChE than did galantamine (IC50 = 9.4 ± 2.5 mM). Moreover, compound 84 exhibited no inhibitory activity against AChE (IC50 value > 350 mM). Inhibitory activity and selectivity (AChE/BuChE) of triazole linked decursinol derivatives may result from not decursinol or triazole moiety but hybrid compounds. Since decursinol itself is an interesting bioactive pharmacological compound, the new biological activity of decursinol derivatives against BuChE will result in beneficial effects for treating AD patients. Also, selective inhibition of BuChE over AChE may have another beneficial effect compared with exclusive use of AChE inhibitors. Additionally, in the same year, the multi-target-directed ligand paradigm was applied to the design of carbamates able to simultaneously target the recently proposed endocannabinoid system and the classic cholinesterase system to achieve effective dual FAAH/cholinesterase inhibitors. Among the two series of synthesised compounds, some derivatives proved to be extremely potent on a single target, and compound 85 (Figure 21) was identified as an effective dual FAAH/ChE inhibitor, with well-balanced nanomolar activities (IC50 = 42.7 nM and 27.9 nM, respectively), and substitution at the phenyl ring carrying the carbamate group proved to be crucial for structure-activity relationships. Thus, compound 85 may be considered as a new promising candidate for AD treatment. In 2017, a novel series of chromenones linked to the 1,2,3-triazole ring system were synthesised and evaluated for their anti-AChE activity by Akbarzadeh et al. The presence of halogen at 2-position of aryl group connected to 1,2,3-triazole increased AChE activity. Among them, N-(1-(2-chlorobenzyl)-1H-1,2,3-triazol-5-yl)methyl)-8-methoxy-2-oxo-2H-chromene-3-carboxamide (86) (Figure 21) bearing methoxy group on the chromeno moiety and 2-chlorophenyl on the pendant 1,2,3-triazole group, exhibited good anti-acetylcholinesterase activity (IC50 = 15.42 µM). Additionally, compound 86 demonstrated a neuroprotective effect against H2O2-induced cell death in PC12 neurons; however, it exhibited no beta-secretase (BACE1) inhibitory activity. Docking and kinetic studies separately confirmed the dual binding activity of compound 86, as it targeted both the catalytic active site (CAS) and the peripheral anionic site (PAS) of AChE. In the same year, multifunctional iminocromene-2H-carboxamide derivatives containing different aminomethylene triazoles with BACE1 inhibitory, neuroprotective, and metal chelating properties targeting Alzheimer’s disease were synthesised. The majority of the synthesised compounds were demonstrated to possess moderate to potent BACE1 inhibitory activity according to a FRET assay. Substitution of phthalimide on amino methylene triazole plays a significant role in BACE1 inhibition. Compound 87 (Figure 21) that possesses a phthalimide pendant on the amino methylene triazole was the most potent derivative, with an IC50 value of 2.2 µM. Compound 87 did not exhibit any cytotoxicity up to a concentration of 100 µM. In 2018, Gharbi et al. studied the anticholinesterase and cytotoxic activities of 2-(1-(2,4,6-trichlorophenyl)-1H-1,2,3-triazol-4-yl)methyl benzopyran-4,3-c] pyrazol-4(2H)-one. The result showed the importance of the triazole ring to enhance the activity. The importance of the chlorine atom, characterised by an electro-attractor effect, compared to the methyl group which, on the contrary, has an electro-donor effect. The importance of the position of the triazole ring in the molecule and the position of this system on the nitrogen N-1 seems more favourable to this activity. Compound 88 (IC50 = 18 µM) (Figure 21) was found to possess the most potent anti-cholinesterase activity and to be less cytotoxic (IC50 = 6.0 µM against the HCT-116 cell line). Compound 88 was the starting compound used to treat the multi-targeted anticholinesterases, anti-5-lipoxygenases, and anti-tyrosinases and to assess anti-cancer activities. In the same year, thirty coumarin-lipoic acid conjugates were synthesised and screened for their new multi-target-directed ligands (MTDLs) for the treatment of Alzheimer’s disease by Jallli-Baleh et al. Among them, 3,4-dimethoxyphenyl-coumarin derivative 89 (IC50 = 16.4 µM) (Figure 21) was approximately twofold more effective than the reference drug donepezil in terms of the inhibitory activity on self-induced and AChE-induced Aβ1-42 aggregation. Moreover, compound 89
provided significant protection against Aβ1-42-induced cytotoxicity that was superior to that of donepezil. The compound 89 contains a propylene linker ($n = 3$). Elongation of linker ($n = 4$ and 5) led to homologs with no activity on cholinesterases. On the other hand, the 3,4-dichlorophenyl-coumarin derivatives with 4 C or 5 C linker showed suitable inhibitory activity against BuChE ($IC_{50}$ values of 10.3 and 7.8 μM, respectively). The 3,4-dichlorophenyl-coumarin analog with propylene spacer ($n = 3$) showed marginal activity against BuChE ($IC_{50} = 73.5$ μM). Therefore, it can be concluded that in 3,4-dichlorophenyl-coumarin derivatives, the elongation of linker increased the anti-BuChE potency. Additionally, a novel hybrid series of umbelliferone and benzyl amine scaffolds linked via triazole rings was synthesised and evaluated for use as both an acetylcholinesterase (AChE) and butyrylcholinesterase (BuChE) inhibitor. All of the synthesised compounds possessed moderate to excellent inhibitory activities. In umbelliferone series, substitution at C-2 position of the phenyl ring was shown to increase the activity in comparison with the unsubstituted derivatives. The order of activity was $\text{Cl} > \text{F} > \text{NO}_2 > \text{H}$, which shows that hydrophobic, bulky and electron withdrawing substituent at C-2 position is needed for optimal activity. According to the $IC_{50}$s in umbelliferone series, the presence of polar substituents such as methoxy at 3 position afforded compounds, exhibiting potent AChE inhibition $\text{OCH}_3 > \text{H} > \text{Cl} > \text{F} > \text{Br} = \text{CH}_3$. Of these, compound 90 (Figure 21) that possessed a 3-methoxy substituent on the benzyl moiety was the most active (AChE and BuChE, $IC_{50} = 3.4$ and $1.1 \mu$M, respectively). Neuroprotection evaluation revealed that this compound efficiently protected PC12 neurons against H$_2$O$_2$-induced cell death. In 2019, a set of novel 1,2,3-triazole-chromone carboxamide derivatives was synthesised and screened for in vitro cholinesterase inhibitory activity by Mina Saeedi. The majority of the synthesised compounds were inactive at a concentration of 100 μM, whereas $N$-(1-benzylpiperidin-4-y)-7-((1-(3,4-dimethylbenzyl)-1H-1,2,3-triazol-4-yl)methoxy)-2-oxo-2H-chromene-3-carboxamide (91) (Figure 21) displayed potent acetylcholinesterase inhibitory activity ($IC_{50} = 1.80 \mu$M), although it was inactive towards butyrylcholinesterase ($IC_{50} \geq 100 \mu$M). The compound 91 possessed benzylpiperidinyl moiety connected to amide functional group and 3,4-dimethylbenzyl connected to 1,2,3-triazole moiety. Compound 91 was evaluated for its BACE1 inhibitory activity, and the calculated $IC_{50}$ of 21.13 μM confirmed the desired inhibitory activity. This compound also exhibited a satisfactory neuroprotective effect against H$_2$O$_2$-induced cell death in PC12 neurons at 50 μM as well as metal chelating ability towards Fe$^{2+}$, Cu$^{2+}$, and Zn$^{2+}$ ions. It seems that benzylpiperidinyl moiety induced better anti-AChE activity than 3-morpholinopropyl which can be associated to similar moiety (benzylpiperidinyl group) ubiquitous in donepezil. Also, the presence of benzylpiperidinyl moiety and 3,4-disubstituted benzyl group connected to 1,2,3-triazole ring is significant for induction of AChEI activity in this series of 1,2,3-triazole-chromone carboxamides.

2.8. Enzyme inhibitors

2.8.1. Pancreatic lipase inhibitors

(E)-labda 8(17), 12-diene-15, 16-dial (Chemical formula: $C_{20}H_{30}O_2$, molecular weight: 302.4580) is a labdane that was isolated from...
the fresh rhizomes of Curcuma amada\textsuperscript{165}. Jalaja et al.\textsuperscript{166} designed and synthesised a new natural product-derived labdane incorporating appended triazoles with pancreatic lipase inhibition potential. The cytotoxicity of the compounds against the Hep G2 human liver carcinoma-derived cell line was measured according to MTT assays. Based on the percentage of cell viability, none of the compounds exhibited any signs of toxicity at any of the tested concentrations. Among the semisynthetic derivatives, the labdane triazole appendages 92a and 92b (Figure 22) were the most active candidates from the series and exhibited excellent pancreatic lipase inhibitory activity (IC\textsubscript{50}: 0.75 ± 0.02 μM and 0.77 ± 0.01 μM) that was slightly higher than that of the positive control Orlistat (IC\textsubscript{50}: 0.8 ± 0.03 μM). In a structure activity relationship point of view, we observed that the labdane-triazole hybrids incorporating benzyl azides were more active than the phenacyl azides. Most of the analogues synthesised from benzyl azides exhibited excellent inhibition property slightly better than or equal to that of Orlistat. In contrast, the triazole analogues synthesised from phenacyl azides did not show any significant inhibition potential except for the compound 6m. In precise, among the triazoles incorporated from the variously substituted benzyl azides, all the para-substituted analogues showed the lowest IC\textsubscript{50}. However, the unsubstituted, ortho- and meta-substituted benzyl azide incorporated triazole appendages showed moderate activity. Interestingly, there was no clear trend followed by the nature of the para substitution, i.e. among the halogen, electron donating and electron withdrawing groups. Overall, among the various substituted hybrids 92a and 92b with p-F and p-Cl substituted benzyl azide incorporated triazole appendages were found to be the most potent candidates of the series.

Coumarin-triazole derivatives not only possess anticancer activity\textsuperscript{58,68,70,90–93}, anti-inflammatory activity\textsuperscript{115}, antibacterial activity\textsuperscript{128,129}, antifungal activity\textsuperscript{133}, antimalarial activity\textsuperscript{140}, antioxidant activity\textsuperscript{145}, and anti-Alzheimer's activity\textsuperscript{154–164}, but also they exhibit pancrelipase inhibitory activity\textsuperscript{167}. Kahveci et al.\textsuperscript{167} designed, synthesised, and studied the anti-lipase activity of the coumarin-triazole hybrid molecule. The majority of the compounds possessed anti-lipase activities at various concentrations. Among the tested compounds, 93a and 93b (Figure 22) exhibited the best anti-lipase activity. These compounds inhibited pancreatic lipase by 99.30 ± 0.56% and 99.85 ± 1.21%, respectively, at a concentration of 10 μM. Orlistat, a pancreatic lipase inhibitor used as an anti-obesity drug, yielded an inhibitory effect of 99.88 ± 0.41% at a concentration of 300 nM (IC\textsubscript{50} = 0.41 ± 0.01 nM). The IC\textsubscript{50} values of compounds 93a and 93b were calculated as 2.64 ± 0.33 μM and 1.80 ± 0.08 μM, respectively. Based on the biological activity, these results indicate that coumarin-triazol hybrid molecules show more activity than coumarin and triazol derivatives.

2.8.2. Xanthine oxidase inhibitors

Anthraquinone (chemical formula: C\textsubscript{14}H\textsubscript{8}O\textsubscript{2}, molecular weight: 208.21, 9,10-Anthracenedione) possesses the structural core of anthracyclines. (1H-1,2,3-triazol-4-yl)methoxybenzaldehyde derivatives containing an anthraquinone moiety were synthesised and identified as novel xanthine oxidase inhibitors\textsuperscript{168}. Among them, the most promising compounds (94a [IC\textsubscript{50} = 0.6 μM] and 94b [IC\textsubscript{50} = 0.8 μM]) (Figure 22) were obtained, and these compounds exhibited greater than 10-fold higher potencies compared to that of the reference xanthine oxidase inhibitor allopurinol. The structure activity relationship analysis revealed that the benzaldehyde moiety played a more important role than did the anthraquinone moiety in its inhibitory potency. Additionally, a formyl group fixed at the 2-position of the phenyl moiety was essential for bioactivity. The Lineweaver–Burk plot showed that compound 94a acted as a mixed-type xanthine oxidase inhibitor. The basis for the significant inhibition of xanthine oxidase by compound 94a was rationalised by molecular modelling studies.

3. Conclusion

Natural products or their derivatives are increasingly used in commercial drugs, which reflects their widespread use as lead compounds for discovering new drugs with novel structures and mechanisms. Triazole has enough potential therapeutic applicability, and is still expanding, it is a multifunctional stent in medicinal chemistry. This review article is an effort to summarise medicinal chemistry investigations of natural product-triazole derivatives in the past six years, in search for new natural product-triazole derivatives which may be an excellent source of promising biological activities. It will help the scientific community rationally design and develop varied, optimised, novel and target oriented natural product-triazole based drugs for treating multifactorial diseases. The structural analysis, molecular docking, activity analysis and mechanism research may provide convenience for further exploration and development of new natural product-triazole derivatives with improved efficacy and reduced toxicity.

4. Future perspective

Due to its wide chemical diversity, natural products have become an important source of biologically active compounds. However, natural biologically active compounds may have inappropriate pharmacological properties, limiting their use, such as cytotoxicity, high lipophilicity or poor oral absorption. The inability to obtain most of these derivatives from sustainable sources is another major limitation in the use of natural products in drug development. Therefore, natural products that have undergone structural
modifications to facilitate the discovery of new drugs with novel structures and mechanisms are widely used as lead compounds. The triazole part has developed into an indispensable heterocyclic scaffold due to its extensive biological properties, especially for the production of natural product-triazole hybrids, even though the triazole part does not exist in nature. This review article mainly studies the biological activity and structure of natural product-triazole derivatives. Among the many biological activities, we comprehensively reviewed the progress in many aspects of the natural product triazole derivatives that have been specially reported in the past six years, such as anti-cancer, anti-inflammatory, anti-microbial, anti-parasitic, anti-viral, antioxidant, and anti-Alzheimer’s disease and enzyme inhibitor action. In this review, we aimed to provide a wide range of data resources on natural product triazole derivatives to medicinal chemists who are engaged in drug design and development, so as to help them conduct more fertile and organised drug discovery operation in the process of experimental studies. Simple structural analysis and molecular docking would be helpful in the structural modification of natural product derivatives with triazole scaffolds. Activity analysis and mechanism research would be very useful for enhancing pharmacokinetic properties and biological activity, and overcoming the difficulty of reducing toxicity and increasing selectivity.

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