RSC Advances



REVIEW

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2022, 12, 31102

Synthesis and bio-properties of 4-piperidone containing compounds as curcumin mimics

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The broad spectrum of curcumin's beneficial properties has encouraged medicinal researchers to investigate its therapeutic efficacy against diverse diseases. The clinical potential of curcumin is, however limited due to its poor pharmacodynamic/pharmacokinetic properties (such as low solubility, pH instability, poor absorption in circulation, rapid elimination from the body and photochemical degradation). 3,5-Bis(ylidene)-4-piperidone scaffolds are considered a curcumin mimic that exhibit diverse bio-properties. The current review provides a brief overview of these mimics and highlights biological activities relevant to drug development.

Received 2nd September 2022 Accepted 23rd October 2022

DOI: 10.1039/d2ra05518j

rsc.li/rsc-advances

1. Introduction

Natural products remain one of the main resources for a variety of diverse human needs. Many natural compounds are used directly as drugs or have inspired the development of potent biologically active agents for clinically use. Artemisinin 1 (Fig. 1) is a classic example which was initially extracted from *Artemisia annua* and subsequently approved as an anti-malarial (*Plasmodium falciparum*) drug. The discoverer of artemisinin was granted the Nobel Prize in 2015 in recognition of contribution

"Department of Pesticide Chemistry, National Research Centre, Dokki, Giza 12622, Egypt. E-mail: girgisas10@yahoo.com; as.girgis@nrc.sci.eg to medicine.¹ Curcumin 2 is another famous example of a natural compound with clinical potential. Curcumin was extracted from roots/rhizomes of *Curcuma longa* and has earned a high reputation among medicinal chemists due to its usage in Ayurvedic medicine, as a food additive and as a dye in many Asian countries. Historically, it was first described over 4000 years ago in ancient India where its usage was associated with religious practice.²

The broad spectrum of curcumin's beneficial properties has encouraged medicinal researchers into its therapeutic efficacy against diverse diseases. It has been reported to possess anti-inflammatory properties and its sodium salt derivative modulates iNOS and COX-2 (cyclooxygenase-2) gene expression in cultured RAW 264.7 cells.³ Antioxidant properties determined by the DPPH (1,1-diphenyl-2-picryl-hydrazyl), ABTS [2,2-azino-



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Fig. 1 Artemisinin and curcumin, examples of biologically important natural compounds

Fig. 2 Piperine, a natural compound useable in traditional medicine, responsible for the pungency of black and long pepper.

bis(3-ethylbenzthiazoline-6-sulfonic acid)], ROO¹ (TRAP) and O²¹ (NET) assays have also shown a free radical scavenging effect of curcumin.⁴ Antiproliferative properties of curcumin against many human cancer cell lines [SMMC-7721 (hepatoma), MCF-7 (breast), PC-3 (prostate), NCI-H460 (non-small cell lung) and K562 (chronic myeloid leukemia)] have been reported.⁴,⁵ In addition curcumin has been shown to display anti-malarial properties against *Plasmodium falciparum*.⁵

Although an enormous amount of research effort has been invested in bringing curcumin and its analogs towards clinical use, thus far no curcumin based product has been approved for use. Over 120 clinical trials on curcumin have failed and consumed federal funds equivalent to 150 million dollars as stated in NIH reports from the last 25 years. This is in part attributed to the promising *in vitro* results obtained in preclinical studies, but poor *in vivo* activities. Thus far curcumin has only been approved as a dietary supplement.¹

The next section highlights some of the important findings of the biological properties of curcumin reported within the last decades with focus on the promising bio-properties of curcumin and its analogs and the potential in identifying more effective hits/leads.

2. Curcumin bio-availability

It has been reported that the major clinical limitation of curcumin is due to poor pharmacological properties such as poor pharmacodynamics/pharmacokinetics, low solubility, pH instability, poor absorption in circulation, rapid elimination and photochemical degradation (giving rise to vanillin, ferulic acid, and other small phenols). 10

The bioavailability of curcumin has been intensively studied. In a phase I clinical trial, Sharma $et\ al.$ reported that the production of the prostaglandin E_2 (PGE2) was (57–62%) decreased in blood samples after 1 h oral administration in patients (with colorectal cancer). Mild diarrhea and discernible toxicity was detected at doses of 0.5–3.6 g daily administration up to four months. Curcumin and its conjugates were detected in plasma and urine and also noticed in patient feces. The entire study concluded that doses of 3.6 g curcumin are recommended for more systemic pharmacological antitumor studies. Additionally, low oral bioavailability (in both animal and human) is supported probably due to intestinal metabolism. The observed bio-properties of curcumin (anti-inflammatory and anticancer) can be attributed to its antioxidant capacity in neutral and acidic pH. $^{11-13}$

Another study investigated the bioavailability of curcumin in rats (in vivo) utilizing high performance liquid chromatography. Disappearance of curcumin was noticed from the rat's plasma within one hour of dosing (i.v. 40 mg kg⁻¹). However, it has



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Fig. 3 Agents considered for mechano-chemical grinding curcumin co-crystals

Fig. 4 Sodium salt curcumin diacetate.

been detected in plasma upon p.o. 400 mg kg⁻¹ administration suggesting that the gastrointestinal tract is more exposed to the unmetabolized curcumin compared to other tissues.¹⁴

Low solubility is one of the major limiting parameters for curcumin's therapeutic use. Different approaches have been considered to overcome this problem and improve bioavailability. Piperine (Fig. 2) has been considered for concomitant administration with curcumin, due to the inhibitory properties in both hepatic and intestinal glucuronidation. Increased serum curcumin was noted upon oral administration (2 g kg $^{-1}$) with piperine (20 mg kg $^{-1}$) in rats, with a significant reduction in the half time of clearance. Similar observations were noticed in human volunteers with slight differences. Time to reach curcumin maximum serum concentration was shorter (earlier) in humans than rats, presumably due to physiological differences. Also, the elimination (clearance) time was shorter in humans than rats. 15

Another approach considered the mechano-chemical grinding of curcumin with diverse agents (nicotinamide, ferulic acid, hydroquinone, *p*-hydroxybenzoic acid and L-tartaric acid) (Fig. 3) in different stoichiometry ratios in an attempt to optimize the physico-chemical properties accessible for solid state oral dosage application. Significant enhancement in solubility and dissolution rates were noticed of the binary eutectics co-crystalline solids.¹⁶

Another study mentioned the possibility of water solubility enhancement through nano-vehicles with curcumin encapsulated in liposomes, exosomes, dendrimers and micelles. ¹⁷ Curcumin encapsulation in liposomes modified with DDAB (didecyldimethylammonium bromide) was studied using cervical cancer cell lines (HeLa, SiHa). It was observed that the

Fig. 5 Structure of curcumin indicating the major active sites.

uptake of DDAB liposomes was better than the non-modified ones but more toxic. Additionally, curcumin was released at a faster rate from cationic DDAB liposomes presumably due to the decrease in interaction of the lipid chains as a result of cationic charges.¹⁸

Sodium salt curcumin diacetate (Fig. 4) was also mentioned as an improved water soluble bio-active agent. Enhancement was noticed relative to that of curcumin itself in aqueous conditions and ability to protect lipid membranes. However, further detailed studies were recommended to support the accessibility for application.¹⁹ Other studies also adopted bioavailability enhancement of curcumin *via* combination with cyclodextrin,²⁰ conjugation with biopolymers²¹ or composite nanoparticles.²²

3. Curcumin chemical structure modification

Many efforts have been directed towards designing novel bioactive agents of enhanced potency and better bioavailability to overcome the drawbacks of curcumin. Manipulation of curcumin chemical structure is usually focused on the aryl rings, carbonyl groups, active methylene or the carbon linker (Fig. 5). Most of the approaches of curcumin chemical structure alterations can be summarized in one of the following approaches.⁹

- Modification of the main curcumin skeletal entities.
- Conjugation with other moieties.
- Curcumin mimics.

4. Curcumin derivatives with potential biological properties

Curcumin connected to amino acid sodium salts 3 were reported as water-soluble agents. Scheme 1 depicted the synthetic pathway via alkylation of the appropriate aldehyde with chloroacetic acid in the presence of NaOH followed by reaction with glycine ethyl ester hydrochloride. Reaction with 2,4-pentandione followed by hydrolysis with methanolic NaOH afforded the targeted agents 3. The synthesized water-soluble agents displayed enhanced antiproliferative properties (MTT assay) against HeLa (cervical cancer) cells (IC $_{50}=0.5~\mu M$ for both the synthesized agents) relative to curcumin (IC $_{50}=4.33~\mu M$) with induction of p53 activity, p21 expression and mediated apoptosis. The p53 is the tumor suppressor protein capable for induction of genes controlling cell cycle and apoptosis. It usually arrests the cell cycle at G2/M phase/transition affecting cyclin dependent kinase (Cdc2) necessary for mitosis. 23

(i) 2-chloroacetic acid, NaOH, H₂O, reflux, 6h; (ii) TBTU, glycine ethyl ester hydrochloride, TEA, CH₃CN, rt, 3h; (iii) 2,4-petanedione, boric anhydride, tri-n-butyl borate, n-butyl amine, ethyl acetate, 0.4 N HCl, rt, 6h; (iv) NaOH, MeOH, H₂O, 15 minutes

(R=H,OCH₃)

Scheme 1 Synthetic pathway of water-soluble curcumin connected to amino acid sodium salts 3.

A series of curcumin–amino acid conjugates 4 were synthesized through reaction of curcumin 2 with the corresponding protected amino acid (benzyloxycarbonyl "Cbz" and fluorenylmethyloxycarbonyl "Fmoc") in the presence of EDAC [1-ethyl-3-(3-dimethylaminopropyl)carbodiimide] and DMAP (4-dimethylaminopyridine) at -5 to 0 °C. The unprotected curcumin-amino acid conjugates 5 were obtained from the Boc (ter-butyloxycarbonyl) protected analogs (Scheme 2). Some of the synthesized conjugates revealed anti-inflammatory properties (acute carrageenan-induced paw edema in rats) with potency higher than curcumin itself and the standard

references used (indomethacin and ibuprofen, clinically used non-steroidal anti-inflammatory drugs). These agents also displayed minor or no ulcerations or lesions on the gastric mucosa of the animals tested, supporting the enhanced bio-properties of the synthesized agents. Additionally, enhanced peripheral (acetic acid-induced abdominal writhing methodology) and central (hot plate technique) analgesic properties were also revealed by some of the synthesized conjugates comparable to curcumin, indomethacin and ibuprofen. The anti-inflammatory properties observed were correlated with the nitric oxide production by lipopolysaccharide-stimulated peritoneal

R = H, Me, $CH(CH_3)_2$, $CH_2CH(CH_3)_2$, $(CH_2)_2SCH_3$, CH_2Ph , $CH_2(3-indolyl)$, $(CH_2)_2CONH_2$; Pg = Cbz, Fmoc

Scheme 2 Synthetic route towards curcumin-amino acid conjugates 4 and 5.

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7, R = H; 8, R = OMe

Scheme 3 Curcumin analogs 7 and 8.

macrophages. Most of the synthesized conjugates showed high antibacterial properties against S. aureus, S. pyogenes (Gram-

positive) and *S. typhi*, *P. aeruginosa* (Gram-negative) bacteria with potency higher than norfloxacin and ciprofloxacin (standard reference antibiotic useable drugs) ²⁴

dard reference, antibiotic useable drugs).²⁴

Dimethoxycurcumin **6** (Fig. 6) revealed antiproliferative properties with efficacy comparable to that of curcumin **2** capable to arrest the cell cycle at S-phase. Dimethoxycurcumin is about three times more metabolically stable relative to curcumin in mice. The mode of action was mentioned through oxidative stress and mitochondrial dysfunction.²⁵

Curcumin analogs 7 and 8 synthesized through basecatalyzed (NaOH, EtOH) condensation of the appropriate aldehyde with acetone were found to be potent inhibitors (MTT

Scheme 4 Synthetic route towards the curcuminoid-difluroborons 9

assay) on a variety of human pancreatic cancers (PANC-1, BXPC-3, MIA-PACA-2, ASPC-1, HPAC and HPDE) and apoptosis inducers relative to curcumin²⁶ (Scheme 3).

Curcuminoid-difluoroborons **9** were synthesized through the reaction of acetyl acetone–BF $_2$ complex with the appropriate aldehyde under nitrogen atmosphere in ethyl acetate in presence of n-butylamine (Scheme 4). Some of the synthesized agents [Ar = 2,3,4-(H $_3$ CO) $_3$ C $_6$ H $_2$; 3,4-5-(H $_3$ CO) $_3$ C $_6$ H $_3$; 3-(H $_3$ CO)-4-(H $_3$ COO)C $_6$ H $_3$] showed notable antiproliferative properties through NCI (National Cancer Institute) screening program (NCI-60) against a variety of human cancer cell lines (SRB technique). 27

Multi-component Biginelli reaction of curcumin with furochromone carbaldehyde **10** and amines, hydrazines, hydroxylamine hydrochloride, urea or thiourea afforded the corresponding furochromone-containing heterocycles **11–14** (Scheme 5). All the synthesized agents showed mild to moderate antiproliferative properties (MTT assay) against MCF7 (breast) and HepG2 (hepatocellular) cancer cell lines relative to doxorubicin and 5-fluorouracil (antitumor standard reference drugs).²⁸

Knoevenagel condensation of curcumin with the appropriate aldehyde in presence of catalytic amounts of piperidine in DMF afforded the corresponding condensate analogs **15**. Trifluoroacetic acid in CH₂Cl₂ was used for Boc group removal (Scheme 6). Higher antiproliferation properties were observed by the synthesized agents against MCF7 (breast) cancer cell line (SRB assay) than curcumin itself. It has also been noted that the synthesized condensates affect microtubules and

 $R = 4-(OH)-3-(H_3CO)C_6H_3; \ R' = 4-MeC_6H_4, \ 2-naphthyl, \ 2-thiazolyl, \ 5-methyl-3-isoxazolyl; \\ R'' = H, \ 4-(NO_2)C_6H_4, \ 2,4-(NO_2)_2C_6H_3; \ X = O, \ S$

Scheme 5 Synthetic route towards furochromone-containing heterocycles 11–14.

MeO
$$_{\text{HO}}$$
 OMe $_{\text{CF}_3\text{CO}_2\text{H}, \text{ CH}_2\text{CI}_2}$ OMe $_{\text{NH}_2\text{CF}_3\text{COO}}$ OMe $_{\text{NH}_2\text{CF}_3\text{COO}}$ R = Ph, 4-(OH)-3-(H₃CO)C₆H₃, NBoc

Scheme 6 Synthetic route towards curcumin condensates 15

Fig. 6 Dimethoxycurcumin 6

Fig. 7 (Z)-3-Hydroxyl-1-(2-hydroxyphenyl)-3-phenylprop-2-ene-1-one.

polymerization of purified tubulin in addition to the ability to induce p53 mediated apoptosis in tested cancer cells.^{29,30}

The antiproliferation properties of (Z)-3-hydroxy-1-(2-hydroxyphenyl)-3-phenylprop-2-ene-1-one **16** (mimicking curcumin) (Fig. 7) was observed by MTT assay against human colon adenocarcinoma (HT29, SW620) cell lines with potency higher than curcumin. Induction of apoptosis indicated by DNA fragmentation and arresting cell cycle at G0/G1 phase was also observed.³¹

Bis(arylidene)monocarbonyl compounds 17–19 were synthesized as modified curcuminoid analogs comprising one ketonic group conjugated with two olefinic groups (Fig. 8) through acid or base catalyzed Claisen Schmidt condensation of

$$\begin{split} R &= 2\text{-}(H_3\text{CO})C_6H_4,\ 4\text{-}(H_3\text{CO})C_6H_4,\ 2\text{-}3\text{-}(H_3\text{CO})_2C_6H_3,\ 2\text{-}5\text{-}(H_3\text{CO})_2C_6H_3,\ 4\text{-}(OH)\text{-}3\text{-}(H_3\text{CO})C_6H_3,\ 2\text{-}CIC_6H_4,\ 4\text{-}CIC_6H_4,\ 4\text{-}FC_6H_4,\ 4\text{-}BrC_6H_4,\ 3\text{-}4\text{-}(H_3\text{CO})_2C_6H_3,\ 4\text{-}4\text{-}BrC_6H_4,\ 4\text{-}BrC_6H_4,\ 4\text{-}BrC_6H_4,\$$

Fig. 8 Bis(arylidene)monocarbonyl compounds 17–19 as modified curcuminoid analogs.

the appropriate aldehyde with acetone, cyclopentanone or cyclohexanone. Many of the synthesized agents showed promising antiproliferative properties against MCF7 (estrogen-dependent breast), MDA-MB-231 (estrogen-independent breast), K562 (chronic myelogenous leukemia) and HeLa (cervical) human cancer cell lines (MTT assay) higher than that of curcumin. Some of them revealed higher potency than that of doxorubicin. (1E,4E)-1,5-Bis(2,5-dimethoxyphenyl)-penta-1,4-dien-3-one [R = 2,5-(H₃CO)₂C₆H₃] is the most potent analog synthesized against the breast cancer cell lines tested (MCF7, MDA-MB-231).³²

A series of curcumin mimics conjugated with chromen-4-one analogs **20** were synthesized through reaction of (1E,4E)-1-aryl-5-(hydroxyphenyl)penta-1,4-dien-3-ones with 3-(bromoalkyloxy)-5,7-dimethoxy-2-(3,4,5-trimethoxyphenyl)-4H-chromen-4-one in DMF containing K_2CO_3 (Scheme 7). Some of the synthesized agents exhibited promising antiproliferative properties of which, 7-dimethoxy-3-(3-(2-((1E,4E)-3-oxo-5-(pyridin-2-yl)penta-1,4-dien-1-yl)phenoxy)propoxy)-2-(3,4,5-trimethoxyphenyl)-4H-chromen-4-one, showed the most promise against gastric cancer cell lines (SGC-7901, MGC-803) relative to doxorubicin (standard reference) with inhibitory properties against TrxR (thioredoxin reductase) observed.³³

5. 3,5-Bis(ylidene)-4-piperidone, bioactive curcumin mimics

5.1. Antitumor active agents

3,5-Bis((*E*)-2-fluorobenzylidene)piperidin-4-one **21** (Fig. 9) was reported to possess antiproliferation properties against MDA-MB231 (breast) and PC3 (pancreatic) cancer cell lines with potency higher than that of curcumin. It was presumed that its mode of action due to the inhibition of intracellular proangiogenic transcription factor (HIF).³⁴

3,5-Bis(4-hydroxyarylidene)-4-piperidones 22 with alkylaminomethyl substituent (Fig. 10) revealed potent antiproliferative properties against Molt 4/C8, CEM (T-lymphocyte) and L1210 (murine leukemia) cell lines (MTT assay) with higher efficacies

n = 3, 4; $R = 3,4-(MeO)_2C_6H_3$, $3-NO_2C_6H_4$, 4-methyl-5-thiazolyl, 2-pyridinyl, $4-ClC_6H_4$, 2-furanyl, 2-thienyl

Scheme 7 Synthetic route towards curcumin mimics conjugated chromen-4-one analogs 20

Fig. 9 3,5-Bis((E)-2-fluorobenzylidene)piperidin-4-one 21.

 $\mathsf{R} = \mathsf{NMe}_2,\, \mathsf{NEt}_2,\, \mathsf{1-pyrrolidinyl},\, \mathsf{1-piperidinyl},\, \mathsf{4-morpholinyl},\, \mathsf{1-(4-methylpiperazinyl)}$

Fig. 10 3,5-Bis(hydroxyarylidene)-4-piperidones 22 with alkylaminomethyl substituents.

than melphalan. Some of synthesized agents characterized as inducer of apoptosis in addition to ability for DNA fragmentation. Cleavage of poly ADP-ribose polymerase is the mode of action assumed for the bio-active agents.³⁵

20

N-Acryloyl-3,5-bis(ylidene)-4-piperidones **23** were synthesized through acid catalyzed condensation (AcOH/HCl gas) of 4-piperidone hydrate hydrochloride with the appropriate aldehyde followed by reaction with acryloyl chloride in CH₂Cl₂ containing TEA (triethylamine) at 0 °C. However, 3,5-bis(ylidene)-1-[*N*-(aryl)maleamoyl]-4-piperidones **24** were obtained through reaction of maleamic acid (generated from the reaction of maleic anhydride and aryl amine in CH₂Cl₂) and the corresponding 3,5-bis(ylidene)-4-piperidones in THF (tetrahydrofuran) containing ethyl chloroformate and TEA (Scheme 8).

 $R = Ph, \ 4-NO_2C_6H_4, \ 4-EtO_2CC_6H_4; \\ R' = 2-FC_6H_4, \ 2-ClC_6H_4, \ 2-MeC_6H_4, \ 2-MeOC_6H_4, \ 2-NO_2C_6H_4, \ 2-6-Cl_2C_6H_3, \ 2,4,6-(H_3C)_3C_6H_2, \ 2-(H_3C)-6-(C_2H_5)C_6H_3, \ 2-(H_3C)-6-(H_3C)_3C_6H_2, \ 2-(H_3C)-6-(H_3C)-6-(H_3C)_3C_6H_2, \ 2-(H_3C)-6-(H_3C$

Scheme 8 Synthetic route towards N-acryloyl-23 and 1-[N-(aryl)maleamoyl]-3,5-bis(ylidene)-4-piperidones 24.

R, R' = 2-F, 3-NO₂, 4-CN, 3,5-OMe, 3,4,5-OMe, R" = F, NHAc

Scheme 9 Synthetic route towards N-arylsufonyl-3,5-bis(arylidene)-4-piperidones 25

R = 3,5-OMe, 3,4,5-OMe, 2,3,4-OMe, 2-F, 2-F-5-NO₂, 2-Br-5-F, 2-CF₃, 4-CF₃, 4-CN; R' = F, CF_3 ; R'' = H, R' = H,

Fig. 11 Dissymmetric pyridine-containing 3,5-(arylidene)-4-piperidones 26–29

Most of the synthesized agents exhibited cytostatic properties against human Molt4/C8, CEM (T-lymphocytes) and L1210 (leukemic) cell lines with higher potency than curcumin and melphalan (standard reference, used in chemotherapeutical combination for chronic leukemia and wide range of malignancies). Safety profile was achieved in vitro (WI-38, human fibroblasts cells) and in vivo (mice) testing. Some of the synthesized agents showed potent inhibitory properties of topoisomerase IIa. This enzyme facilitates DNA replication by preventing the buildup of supercoils during replication fork progression. Inhibitors of this enzyme result in the generation of multiple DNA strand breaks, arrest cell division and eventually lead to apoptosis. As such numerous topoisomerase inhibitors are currently used in clinics as chemotherapy for numerous malignancies. The synthesized agents also revealed antioxidant properties.36

A series of *N*-arylsulfonyl-3,5-bis(arylidene)-4-piperidones **25** were synthesized through base-catalyzed arylsulfonation

(catalytic amount of pyridine in CH₂Cl₂ at room temperature) of the corresponding *N*-unsubstituted piperidones (Scheme 9). The synthesized agents showed anti-inflammatory properties supported by the inhibition of IL-6 and TNF-α in RAW264.7 cells induced by lipopolysaccharide (LPS of Gram-negative bacteria). Promising antiproliferation properties were also mentioned by some of the synthesized agents against liver (HepG2, SMMC-7721, QGY-7703) cancer cell lines (MTT assay) with induction of apoptosis. Association of chronic inflammation with cancer progression especially, hepatic cancer is the rational for investigation of anti-inflammatory and antiproliferation properties of the synthesized agents.³⁷

Other sets of dissymmetric pyridine-containing 3,5-bis(arylidene)-4-piperidones $\bf 26-29$ were also reported (Fig. 11) with anti-inflammatory and anti-hepatoma properties similar to compounds $\bf 25.^{38}$

Pyrido[4,3-d]pyrimidines **30** were synthesized through reaction of the corresponding fluoro-containing *N*-arylsulfonyl-3,5-

R = 2-F, 3-F, 4-F; R' = H, Me, NO₃, CF₃, F, CI, Br, CN, NHAc

Scheme 10 Synthetic route towards pyrido[4,3-d]pyrimidines 30.

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 $R = Ph, 4-CIC_6H_4, 4-FC_6H_4, 4-BrC_6H_4, 4-H_3CC_6H_4, 4-H_3COC_6H_4, 3,4-(H_3CO)_2C_6H_3, 2-thienyl; \\ R' = Me, Et, 1-Pr$

Scheme 11 Synthetic route towards 1-(alkylsulfonyl)-3,5-bis(ylidene)-4-piperidinones 31.

$$\begin{split} R = Ph, \, 4\text{-}CIC_6H_4, \, 4\text{-}FC_6H_4, \, 4\text{-}H_3COC_6H_4, \, 4\text{-}H_3COC_6H_4, \, 2\text{-}thienyl;} \\ R' = Ph, \, 4\text{-}CIC_6H_4, \, 4\text{-}H_3COC_6H_4, \, C_2H_5, \, H_2CC_6H_5 \end{split}$$

Scheme 12 Synthetic route towards 3,5-bis(ylidene)-*N*-substituted-4-piperidinone-1-carboxamides **32**.

bis(arylidene)-4-piperidones with guanidine hydrochloride in ethanolic KOH (Scheme 10). Antiproliferative properties were observed by the targeted agents against a variety of hepatocellular carcinoma cells (HepG2, SMMC-7721) using MTT assay. Inhibition of the nuclear translocation of NF- κ B induced by TNF- α or LPS supports the anti-inflammatory properties of these compounds, considering that NF- κ B is the signal pathway connecting the chronic inflammation and hepatocellular carcinoma.³⁹

Potent antiproliferative agents against HCT116 (colon) and A431 (skin/squamous) cancer cell lines were exhibited by 1-(alkylsulfonyl)-3,5-bis(ylidene)-4-piperidinones **31** (Scheme 11) relative to 5-fluorouracil (approved drug for colon, breast and skin cancers). Some of the synthesized agents also showed high potency against MCF7 (breast) and A549 (lung) cancer cell lines (relative to 5-fluorouracil and doxorubicin) with minimal cytotoxicity against RPE1 (non-cancer, retinal pigment epithelial) cell line. The synthesized agents exerted their mode of action *via* inhibitory properties of topoisomerase IIα which is the enzyme responsible for breaking double strand DNA helix during DNA replication, transcription and repairing.⁴⁰

A set of 4-piperidone-1-carboxamides 32 were synthesized *via* reaction of isocyanate with the appropriate *N*-unsubstituted 3,5-diylidene-4-piperidine in DMF in the presence of TEA (Scheme 12). Most of the synthesized agents revealed high potency against HCT116 (colon), MCF7 (breast) and A431 (skin/squamous) cancer cell lines with higher efficacy than that of 5-fluorouracil and safe behavior against non-cancer (RPE1) cell line. The synthesized agents revealed topoisomerase II-α inhibitory properties supporting their mode of action.⁴¹

 $R = Ph, 4-FC_6H_4, 4-CIC_6H_4, 4-H_3CC_6H_4, 4-H_3COC_6H_4, 4-NO_2C_6H_4, 3,4-(H_3CO)_2C_6H_3$

Scheme 13 Synthetic route towards 1-[3-(2-methoxyethyloxy)propionyl]-4-piperidones 33 and their thio analogs 34-36.

R =H, CI; R' = Ph, $4-FC_6H_4$, $4-CIC_6H_4$, $4-B_3CC_6H_4$, $4-H_3CO_6H_4$, $4-H_3CO_6H_4$, $4-(H_3CO)_2C_6H_3$, 2-thienyloon = 1

Scheme 14 Synthetic route towards piperidone-salicylate conjugates 37.

R = Ph, $4-FC_6H_4$, $4-CIC_6H_4$, $4-BrC_6H_4$, $4-H_3CC_6H_4$, $3,4-(H_3CO)_2CIC_6H_3$; n = 1,2

Scheme 15 Synthetic route towards 3,5-di[(E)-arylidene]-1-[3-(4-methylpiperazin-1-yl)alkyl]piperidin-4-ones 38.

A set of 1-[3-(2-methoxyethyloxy)propionyl]-4-piperidones 33 and their thio-analogs 34 were synthesized through reaction of the acyl chloride with the appropriate 3,5-bis(ylidene)-4piperidone in CH₂Cl₂ containing triethylamine. The sulfinyl-35 and sulfonyl-36 derivatives were obtained through oxidation of compounds 34 with peracetic acid and 3-chloroperoxybenzoic acid in CH₂Cl₂, respectively (Scheme 13). Antiproliferative properties were observed by the synthesized agents against human Molt 4/C8, CEM (T-lymphocyte) and L1210 (murine leukemia) cell lines with safer behavior towards nonmalignant cells. Some of the synthesized agents demonstrated PARP1 [poly(ADP-ribose)polymerase 1] cleavage, a characteristic hallmark of apoptosis. PARP1 is capable of repairing DNA single-stranded breaks, thus compounds that induce PARP1 cleavage can prevent DNA replication and are useful in cancer chemotherapy.42

The piperidone-salicylate conjugates 37 were synthesized through dehydrohalogentation of acetylsalicylic acid chloride

with the appropriate piperidone in DMF containing TEA as basic catalyst (Scheme 14). Potent antiproliferation properties were noticed by the synthesized conjugates against A431 (squamous skin), HCT116 (colon) and MCF7 (breast) cancer cell lines (MTT assay) with comparable efficacies to that of 5-fluorouracil and sunitinib (standard references). Multi-targeted inhibitory properties were observed against VEGFR-2 (vascular endothelial growth factor receptor-2) and EGFR (epidermal growth factor receptor) in both MCF7 (breast) and HCT116 cancer cells. Enhanced COX-1 and COX-2 (cyclooxygenase-1 and -2) inhibitory properties were also revealed by the synthesized agents than that of aspirin supporting their anti-inflammatory properties. Selective inhibition was noticed towards COX-2 compared to COX-1. Additionally, some of the synthesized agents revealed antiviral properties against SARS-CoV-2 (respiratory syndrome coronavirus 2) which is the responsible infectious microorganism of COVID-19 (corona virus disease 2019) pandemic.43

$$\begin{array}{c} O \\ \hline \\ N_{a_2CO_3} \\ H.HCl.H_2O \end{array} \begin{array}{c} O \\ \hline \\ N_{a_2CO_3} \\ \hline \\ MeCN, H_2O \end{array} \begin{array}{c} O \\ \hline \\ N_3(CH_2)_nP(OEt)_2 \\ \hline \\ Cu(l), {}^tBuOH/H_2O \end{array} \begin{array}{c} O \\ \hline \\ N_1 \\ \hline \\ N_2 \\ \hline \\ OEt \end{array} \begin{array}{c} O \\ \hline \\ R \\ \hline \\ N_1 \\ \hline \\ N_2 \\ \hline \\ N_2 \\ \hline \\ N_2 \\ \hline \\ OEt \end{array}$$

 $R = 4-O_2NC_6H_4$, $4-NCC_6H_4$, $4-FC_6H_4$, $4-H_3COC_6H_4$; n = 3, 4

Scheme 16 Synthetic route towards 3,5-bisarylidene-4-piperidones connected with 1,2,3-triazolyl heterocycle bearing phosphonate group 39.

 $\begin{array}{c} O \\ N \\ N \\ H \\ H_2O \end{array} \begin{array}{c} R \\ O \\ P(OEt)_3, Mg(CIO_4)_2, \\ Et_3N \\ P \\ O \\ OEt \\ \end{array} \begin{array}{c} O \\ R \\ O \\ OEt \\ OEt \\ \end{array} \begin{array}{c} R' \\ O \\ CIOO_4/Et_3N \\ R' \\ O \\ OEt \\ O \\ OEt \\ \end{array}$

R = H, 4-F, 4-MeO, 2,4-(MeO)₂,2,3,4-(MeO)₃; R' = H, 4-F, -4-MeO, 4-NO₂, $4-Me_2N$, 4-CN, 2,3,4-(MeO)₃

Scheme 17 Synthetic route towards 3,5-bis(arylidene)-4-piperidones attached to diethyl[(aryl)methyl]phosphonate moiety 40.

 $R = H, CI, OMe; R' = Ph, 4-CIC_6H_4, 2,4-CI_2C_6H_3, 4-FC_6H_4, 4-H_3CCC_6H_4, 4-H_3COC_6H_4\\ 2-thienyl, 3-pyridinyl; R'' = Me, Et$

Scheme 18 Synthetic route towards dispiro[3*H*-indole-3,2'-pyrrolidine-3',3"-piperidines] 41.

A set of antiproliferative active agents 3,5-di[(*E*)-arylidene]-1-[3-(4-methylpiperazin-1-yl)alkyl]piperidin-4-ones **38** were prepared through acylation of the appropriate 3,5-bis(ylidene)-4-piperidones followed by dehydrohalogenation *via* reaction with *N*-methylpiperazine (Scheme 15). Potent antiproliferative properties were exhibited against HCT116 (colon) and MCF7 (breast) cancer cell lines relative to sunitinib and 5-fluorouracil (standard references). Dual inhibitory properties were observed against human topoisomerase I and IIα. However, with higher efficacy against topoisomerase IIα than I. Promising anti-SARS-CoV-2 properties were also revealed relative to favipiravir (standard reference) during the VERO-E6 standard technique. Lack of cytotoxicity against normal RPE1 and VERO-E6 cells supports the possibility that these compounds may be potential drug candidates.⁴⁴

A series of 3,5-bis(arylidene)-4-piperidones connected with 1,2,3-triazolyl heterocycle bearing phosphonate group **39** were obtained via Aldol condensation (Et₂O·BF₃) (Scheme 16). Promising antitumor properties were observed against HCT116 (colon) and MCF7 (breast) cancer cell lines (relative to

doxorubicin, reference standard) with limited toxicity against normal (HEF) cells.⁴⁵

40

A series of 3,5-bis(arylidene)-4-piperidones attached to diethyl[(aryl)methyl]phosphonate moiety **40** were synthesized *via* reaction of α -amino(aryl)methyl phosphonates (obtained through Kabachnik–Fields reaction of triethyl phosphite, appropriate aldehyde and 4-piperidone hydrochloride monohydrate) with aromatic aldehyde in the presence of LiClO₄/Et₃N (Lewis acid) (Scheme 17). Noticeable antiproliferative properties were observed against a variety of human tumor cell lines (RD "rhabdomyosarcoma", PC3 "pancreatic", HCT116 "colon", and MCF7 "breast") cell lines relative to doxorubicin and daunorubicin (reference standards).⁴⁶

A series of dispiro[3*H*-indole-3,2'-pyrrolidine-3',3"-piperidines] **41** were synthesized through dipolar cycloaddition of the appropriate 3,5-bis(ylidene)-4-piperidones with azomethine ylide (which obtained *in situ* through reaction of sarcosine with isatins) (Scheme 18). The stereochemical structure of **41** was established through single crystal X-ray studies. Many of the synthesized agents showed promising antiproliferation properties against HeLa (cervical), MCF7, T-47D (breast), HepG2

Fig. 12 Dispiro-analog bearing 1-[(4-morpholinyl)methylene] function 42.

(liver) and HCT116 (colon) carcinoma cell lines (SRB technique) relative to that of doxorubicin and cisplatin (standard references). QSAR (CODESSA-Pro, CODESSA III software) and 3D-

pharmacophoric (Discovery Studio 2.5) studies discussed the biological properties observed, optimized molecular models and exhibited the descriptors/chemical features necessary for bio-observations.⁴⁷⁻⁴⁹

Dispiro-analog bearing a 1-[(4-morpholinyl)methylene] group 42 was also synthesized through azomethine dipolar cycloaddition reaction with the appropriate piperidone (Fig. 12). Quantum chemical calculations [DFT/B3LYP, 6-31G(d,p)] determined the stereochemical structure. Promising antitumor properties were observed against diverse human cancer cell lines (National Cancer Institute screening program). 50

5.2. Cholinesterase inhibitors

Alzheimer's disease is a neurodegenerative disease comprising the main cause of dementia particular in elder people. One of

 $R = Ph, 4-CIC_6H_4, 4-H_3CC_6H_4, 2-pyridinyl; R' = H, Cl, CH_3, NO_2$

Scheme 20 Synthetic route towards spiropyrrolidines 47.

the most common methods for treatment is the elevation of acetylcholine levels in the brain. Acetylcholinesterase (AChE) and butyrylcholinesterase (BChE) are two enzymes that exist in the central nervous system capable of hydrolyzing acetylcholine (neurotransmitter at cholinergic synapses). For this reason compounds with inhibitory properties against these enzymes are valuable for controlling progress of this disease. A series of 3,5-bis(ylidene)-4-piperidones 43 (Fig. 13) showed promising inhibitory properties against AChE and BChE (Ellman's method) compared to Tacrine and Donepezil (standard references). 51,52

Pyrido[4,3-d]pyrimidines 44 obtained through reaction of 3,5-bis(ylidene)-4-piperidones with thiourea in presence of NaOEt. Reaction of 44 with phenacyl bromides afforded the formation of pyrido[4,3-d]thiazolo[3,2-a]pyrimidines 45. Alternatively, 45 could be obtained directly through domino reaction of the appropriate piperidone, thiourea and phenacyl bromide in ionic liquid (1-butyl-3-methylimidazolium bromide "[bmim] Br") under microwave irradiation (Scheme 19). Promising AChE and BChE inhibitory properties were observed by some of the synthesized agents 44 and 45 (Ellman's method) relative to Galantamine. ⁵³⁻⁵⁵

 $R = 2-\text{Cl-}3-\text{H}_3\text{COC}_6\text{H}_3, \ 2-\text{Cl-}3,4-(\text{H}_3\text{CO})_2\text{C}_6\text{H}_2, \ 2-\text{Cl-}3,4,5-(\text{H}_3\text{CO})_3\text{C}_6\text{H}, \ 4-(\text{EtO})_2\text{CH}_2\text{C}_6\text{H}_4, \ 2-\text{NO}_2-4-(\text{CH}_3)_2\text{NC}_6\text{H}_3, \ 4-(\text{pyrrolidin-}1-\text{yl})\text{C}_6\text{H}_4; \ R' = \text{H, Me, CH}_2\text{C}_6\text{H}_5, \ R' = \text{H, Me, CH}_2\text{C}_6\text{H}_6, \ R' = \text{H, Me, CH}_2$

Fig. 13 3,5-Bis(ylidene)-4-piperidones of cholinesterase inhibitory properties 43.

 $R = Ph, 2-FC_6H_4, 4-FC_6H_4, 2-ClC_6H_4, 4-ClC_6H_4, 2,4-Cl_2C_6H_3, 4-BrC_6H_4, 2-H_3CC_6H_4, 4-H_3CC_6H_4, 2-H_3COC_6H_4, 3,5-(H_3CO)_2C_6H_3, 3-NO_2C_6H_4; R' = H, Et, 4-(morpholinyl)CH_2CH_2; R'' = Ph, 4-FC_6H_4, 4-ClC_6H_4, 4-BrC_6H_4, 4-H_3CCC_6H_4, 4-NO_2C_6H_4, 2-naphthyl$

Scheme 19 Synthetic route towards pyrido[4,3-d]pyrimidine 44 and pyrido[4,3-d]thiazolo[3,2-a]pyrimidines 45.

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 $R = Ph, 2-H_3CC_6H_4, 2-H_3COC_6H_4, 2-CIC_6H_4, 2-FC_6H_4, 3-NO_2C_6H_4, 2,4-CI_2C_6H_3, 4-H_3CC_6H_4, 4-CIC_6H_4, 4-FC_6H_4, 1-naphthyl$

Scheme 21 Synthetic route towards spiro-heterocycles 48.

Spiropyrrolidines 47 were synthesized *via* [3 + 2]-dipolar cycloaddition reaction of 3-(ylidene)-*N*-substituted-4-piperidones 46 with azomethine ylides formed from condensation of the appropriate isatin and sarcosine in presence of ionic liquid "[bmim]Br" (Scheme 20). Promising AChE and BChE inhibitory properties were observed by some of the synthesized agents (Ellman's method) compared with Galantamine (reference standard).⁵⁶

Cycloaddition reaction of azomethine ylide (formed from condensation of 5-chloroisatin and L-proline) with *N*-acryloyl-3,5-bis(ylidene)-4-piperidones **23** afforded the corresponding spiro-heterocycles **48** *via* reaction with the acryloyl linkage rather than the exocyclic ylidene olefinic linkage (Scheme 21). Stereochemical structure of **48** was supported by single crystal X-ray studies. AChE and BChE properties were exhibited by the synthesized agents in comparison with Galantamine (standard reference).⁵⁷

Additionally, azomethine cycloaddition reaction (formed from isatin and L-proline) with *N*-acryloyl-3,5-bis(ylidene)-4-piperidones **23** in refluxing MeOH in equimolar values afforded the mono-spiro-heterocycles **49** in a similar manner to that

of the aforementioned formation of **48**. However, reaction of azomethine ylide in double folds amount (two molar equivalents) to that of piperidones **23**, the bis-spiro-heterocycles **50** were obtained due to double cycloaddition reactions with both acryloyl and ylidene linkages (Scheme **22**). Both compounds **49** and **50** revealed AChE and BChE inhibitory properties (Ellman's method) and some of them showed potency comparable to that of Galantamine.⁵⁸

Meanwhile, ionic liquid "[bmim]Br" mediated cycloaddition reaction of azomethine ylide derived from isatin and sarcosine with *N*-acryloyl-3,5-bis(ylidene)-4-piperidones 23 afforded the mono-spiro-pyrrolidines 51 due to cycloaddition reaction with the exocyclic ylidene linkage. However, reaction of the azomethine ylide with 23 in 2:1 molar value equivalent afforded the bis-spiro-pyrrolidines 52. The difference in observations of this reaction to that mentioned in Scheme 22 is attributed to the different reactant azomethine ylide derived from diverse amino acid (sarcosine and L-proline). Additionally, bis-spiro-pyrrolidines 52 were obtained from mono-spiro-pyrrolidines 51 by reaction with another mol equivalent of the azomethine ylide (Scheme 23). Both mono-spiro-51 and bis-spiro-pyrrolidines 52

 $R = Ph, 2-H_3CC_6H_4, 2-H_3COC_6H_4, 2-CIC_6H_4, 2-FC_6H_4, 3-NO_2C_6H_4, 2,4-CI_2C_6H_3, \\ 4-H_3CC_6H_4, 4-CIC_6H_4, 4-FC_6H_4, 1-naphthyl$

Scheme 22 Synthetic route towards mono-spiro-49 and bis-spiro-heterocycles 50.

 $R = Ph, 2-H_3CC_6H_4, 2-H_3COC_6H_4, 2-CIC_6H_4, 2-FC_6H_4, 3-NO_2C_6H_4, 2,4-CI_2C_6H_3, \\ 4-H_3CC_6H_4, 4-CIC_6H_4, 4-FC_6H_4, 1-naphthyl \\$

Scheme 23 Synthetic route towards mono-spiro-51 and bis-spiro-heterocycles 52

revealed noticeable AChE and BChE inhibitory properties and some of them exhibited potency comparable to that of Galantamine.⁵⁹

Similarly, reaction of *N*-acryloyl-3,5-bis(ylidene)-4-piperidones 23 with azomethine ylide generated from phenylglycine and isatin (in 1:1 molar equivalent) in ionic liquid "[bmim]Br" medium afforded the mono-spiro-pyrrolidines 53 due to cycloaddition reaction with the exocyclic ylidene linkage.

However, reaction of the azomethine ylide with 23 in 2:1 molar value equivalent afforded the bis-spiro-pyrrolidines 54. Reaction of 53 with another mol equivalent of azomethine ylide also afforded the bis-spiro-pyrrolidines 54 (Scheme 24). AChE and BChE inhibitory properties were shown by the mono-spiro-53 and bis-spiro-pyrrolidines 54 and some of them exhibited potency comparable to that of Galantamine.⁶⁰

 $R = Ph, 2-H_3CC_6H_4, 2-H_3COC_6H_4, 2-CIC_6H_4, 2-FC_6H_4, 3-NO_2C_6H_4, 2,4-CI_2C_6H_3, \\ 4-H_3CC_6H_4, 4-CIC_6H_4, 4-FC_6H_4, 1-naphthyl \\$

Scheme 24 Synthetic route towards mono-spiro-53 and bis-spiro-heterocycles 54.

$$\begin{array}{c} NH_2 \\ NH_2 \end{array} + \begin{array}{c} O \\ NH_2 \end{array} + \begin{array}{c} O$$

 $R = Ph, \ 4-BrC_6H_4, \ 2-ClC_6H_4, \ 2,4-Cl_2C_6H_3, \ 4-ClC_6H_4, \ 2-H_3CC_6H_4, \ 3-H_3CC_6H_4, \ 4-H_3CC_6H_4, \ 3-NO_2C_6H_4$

Scheme 25 Synthetic route towards dispiropyrrolidines 55.

 $R=H,\ Me,\ Et,\ Pr,\ \emph{i-}Pr,\ cyclo-Pr,\ SO_2-Ph,\ CH_2Ph,\ CO(2,4-Cl_2C_6H_3),\ CO(4-FC_6H_4),\ CO(4-ClC_6H_4),\ CO(4-H_3CC_6H_4);\ R'=4-methylpiperazin-1-yl,\ 4-morpholinyl,\ 4-pyrrolidin-1-yl,\ Et_2N,\ Me_2N$

Scheme 26 Synthetic route towards (allylated-benzylidene)-4-piperidones 57, 59.

R = Ph, 2-naphthyl, 2-thienyl, 3-indolyl, 5-indolyl, 4-[(4-bromobenzyl)oxy]C₆H₄, 4-imidazolyl, 3-methtyl-2-thienyl, 4-methyl-2-thienyl

Fig. 14 3,5-Bis(ylidene)-4-piperidones **60** of anti-inflammatory properties.

Indeno[3,2-b]quinoxalin-11-one were synthesized through condensation of ninhydrin and o-phenylenediamine in refluxing methanol. Ionic liquid mediated "[bmim]Br" multicomponent [3 + 2]-dipolar cycloaddition reaction of azomethine ylide (obtained from the condensation of ι -tryptophan and indeno [3,2-b]quinoxalin-11-one) with 3,5-bis(ylidene)-4-piperidine afforded the dispiropyrrolidines 55 (Scheme 25). AChE and BChE inhibitory properties were shown by the synthesized dispiropyrrolidines 55 relative to Galantamine.

5.3. Anti-inflammatory active agents

Acute lung injury is life threatening and usually associated with acute inflammatory factors IL-6, IL-1β (interleukin) and TNF-(tumor necrosis factor). The 3,5-bis(3-allyl-4hydroxybenzylidene)-4-piperidones 57 were synthesized through acid-catalyzed condensation [HCl(gas)/AcOH] of 4piperidones with 3-allyl-4-hydroxybenzaldehyde (56). Mean-3-aryliden-4-piperidone 59 obtained via while, was

condensation of the 3-(ylidene)-1-(cyclopropyl)-4-piperidone **58** (O-THP protected benzaldehyde, obtained from reaction of **56** with 3,4-dihydro-2*H*-pyran) with various aldehydes (basecatalyzed reaction, NaOH/EtOH) (Scheme 26). Piperidones **57** and **59** revealed inhibitory properties against IL-6 and TNF- α supporting the possibility as a potential treatment to prevent inflammation associated with lung injury.⁶²

Anti-inflammatory properties of 3,5-bis(ylidene)-4-piperidones **60** (Fig. 14) were supported by the *in vivo* carrageenin-induced paw oedema of rats (i.p., 0.01 mmol per kg body weight). Some of the tested agents revealed higher efficacy than that of indomethacin (non-steroidal anti-inflammatory drug) after 3.5 h of administration.⁶³

2-{3,5-[Bis(ylidene)-4-oxopiperidin-1-yl]}-2-oxoethylmorpholine-4-carbodithioates **61** where synthesized through reaction of sodium morpholine-4-carbodithioate with the appropriate 1-chloroacetyl-3,5-bis(ylidene)-4-piperidones in methanol-water (1:1) at 60 °C (Scheme 27). Some of the synthesized **61** revealed promising down-regulation properties of TNF-α-induced NF-κB activation in KBM5 cell. NF-κB possesses a dual role due to its capability to regulate many growth factors and cytokines responsible for anti-apoptosis, angiogenesis and metastasis. Suppression of NF-κB is an important therapeutical pathway against cancer and inflammation of cells. ⁶⁴

Dispiro-heterocyles 62 were synthesized by the azomethine ylide [3 + 2]-cycloaddition reaction with 3,5-bis(ylidene)-4-piperidones in refluxing EtOH (Scheme 28). Single crystal X-ray studies supported the structure. Anti-inflammatory

$$\begin{split} \text{R} &= \text{Ph, } 3,4\text{-}(\text{MeO})_2\text{C}_6\text{H}_3, \, 2\text{-}\text{CIC}_6\text{H}_4, \, 3\text{-}\text{CIC}_6\text{H}_4, \, 4\text{-}\text{BrC}_6\text{H}_4, \, 4\text{-}\text{MeC}_6\text{H}_4, \, 4\text{-}\text{C}_2\text{H}_5\text{C}_6\text{H}_4, \, 2,4\text{-}\text{CI}_2\text{C}_6\text{H}_3, \\ 3\text{-}\text{FC}_6\text{H}_4, \, 4\text{-}\text{FC}_6\text{H}_4, \, 3,5\text{-}\text{F}_2\text{C}_6\text{H}_3, \, 3,4,5\text{-}(\text{MeO})_3\text{C}_6\text{H}_2, \, 2\text{-}\text{F}_3\text{CC}_6\text{H}_4, \, 4\text{-}\text{MeOC}_6\text{H}_4, \, 2,6\text{-}\text{F}_2\text{C}_6\text{H}_3, \, 2,5\text{-}\text{F}_2\text{C}_6\text{H}_3, \\ 2,5\text{-}(\text{MeO})_2\text{C}_6\text{H}_3, \, 3,5\text{-}\text{Me}_2\text{C}_6\text{H}_3, \, 2,5\text{-}\text{Me}_2\text{C}_6\text{H}_3, \, 4\text{-}^\text{t}\text{BuC}_6\text{H}_4, \, 4\text{-}\text{F}_3\text{CC}_6\text{H}_4 \end{split}$$

Scheme 27 Synthetic route towards 3,5-[bis(ylidene)-4-piperidone-1-yl]-2-oxoethylmortholine-4-carbodithioates 61.

 $R = Ph, 4-BrC_6H_4, 4-ClC_6H_4, 4-FC_6H_4, 4-H_3CC_6H_4, 4-H_3COC_6H_4, 2-thienyl; R' = H, Me$

Scheme 28 Synthetic route towards dispiro-heterocycles 62 of anti-inflammatory properties.

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 $R = Ph, 4-CIC_6H_4, 4-H_3CC_6H_4, 4-NO_2C_6H_4; R' = H, COPh, CO-4-C_6H_4-CH_3, CO-4-C_6H_4O(CH_2)_2NMe_2, CO-4-C_6H_4O(CH_2)_2NEt_2, CO-4-C_6H_4O(CH_2)_2(1-piperidinyl), CO-4-C_6H_4O(CH_2)_2(4-morpholinyl), CO-4-C_6H_4O(CH_2)_2N^{\dagger}Me_3. \\ C_6H_4O(CH_2)_2N^{\dagger}Me_3. \\ I^{-}$

Fig. 15 3,5-Bis(ylidene)-4-piperidones 63 with noticeable antimycobacterial properties.

properties of the synthesized agents were established by the carrageenin-induced paw edema method of rats (i.p., 50 mg per kg body weight). Some of the synthesized agents revealed higher anti-inflammatory potency than that of indomethacin after 4 h of administration.⁶⁵

5.4. Antimycobacterial active agents

Infectious diseases are still one of the main human global health problems. Tuberculosis (TB) is one of the top ten human life threatening globally. It is the second cause of mortality after

HIV/AIDS due to single infectious pathogen. Various pathogenic agents (*Mycobacterium* sp.) have been identified causing TB. Many drugs have been discovered for treating patients with TB but due to the side effects and drug resistant TB strains, novel therapeutical agents are still needed.⁶⁶

A series of 3,5-bis(ylidene)-4-piperidones 63 were identified with noticeable antimycobacterial properties (M. tuberculosis $H_{37}Rv$). Some of them revealed properties in the rat liver mitochondria respiration with swelling in mitochondria⁶⁷ (Fig. 15).

 $R = 4 - H_3CC_6H_4$, $4 - H_3COC_6H_4$, $2 - CIC_6H_4$, $3 - FC_6H_4$, $2, 4 - CI_2C_6H_3$

Scheme 29 Spiro-heterocycles 65-67 of antimycobacterial properties

R = Ph, 2-BrC₆H₄, 4-BrC₆H₄, 2-ClC₆H₄, 2,4-Cl₂C₆H₃, 4-ClC₆H₄, 2-H₃CC₆H₄, 3-H₃CC₆H₄, 4-H₃CC₆H₄, 2-H₃COC₆H₄, 3-H₃COC₆H₄, 4-H₃COC₆H₄, 3-NO₂C₆H₄; R' = H, OCF₃

Scheme 30 Dispiropyrrolidines 68 of antifungal properties.

R = Ph, 2-ClC₆H₄, 3-ClC₆H₄, 4-ClC₆H₄, 3-BrC₆H₄, 4-BrC₆H₄, 4-MeOC₆H₄, 4-MeO₆H₄, 4-MeO₆ 3-F₃CC₆H₄, 3-O₂NC₆H₄, 4-O₂NC₆H₄,4-MeSC₆H₄, 2-Cl-5-F₃CC₆H₃, 2,4-(MeO)₂C₆H₃, 3,4-(MeO)₂C₆H₃, 3,5-Cl₂C₆H₃, 4-OH-3,5-(MeO)₂C₆H₂,

Fig. 16 3,5-Bis(ylidene)(-N-methyl-4-piperidones) 69 of anti-plasmodial properties.

R = 1-piperidinyl, 4-morpholinyl, NEt₂, NMe₂; R' = Ph, 4-ClC₆H₄, 4-NO₂C₆H₄, 4-H₃CC₆H₄

Scheme 31 N-acyl-3,5-bis(ylidene)-4-piperidones 70 of antimalarial properties.

Solvent-free microwave irradiation of 1-methyl-4-piperidone with aromatic aldehyde in presence of pyrrolidine afforded the corresponding 3-arylidene-4-piperidones 64. Azomethine ylide (derived from condensation of isatin with sarcosine, proline or benzylamine) cycloaddition to 64 in refluxing methanol yielded the corresponding spiro-heterocycles 65–67. Single crystal X-ray studies supported the synthesized structures. Antimycobacterial properties were observed by the synthesized spiro-heterocycles 65-67 against M. tuberculosis, multi-drug resistant M. tuberculosis and M. smegmatis⁶⁸ (Scheme 29).

5.5. Antifungal active agents

Dispiropyrrolidines 68 were synthesized through multicomponent domino azomethine ylide (formed from condensation of 2-amino-3-phenylpropanoic acid and isatin) dipolar cycloaddition reaction with the appropriate 3,5bis(arylidene)-4-piperidones in ionic liquid (Scheme 30). Antifungal properties of 68 were revealed against Candida albicans ATCC 10231. Some of the synthesized agents showed potent inhibitory properties relative to fluconazole (standard reference).69

Antimalarial active agents

Malaria is one of the most widely distributed infectious diseases in tropical and subtropical regions. *Plasmodium* sp. which is a protozoan organism transmitted to humans due to mosquito bites. Although many drugs are known to combat malaria

 $X = CI, Br, I; R = 2-F, 2,4-CI_2, 4-NO_2$

Scheme 32 Synthetic route towards 1-ethoxycarbonyl-3,5-bis[(3'-indolyl)methylene]-4-piperidines 71 lipase inhibitory properties.

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emerging drug resistance means that new effective agents are still needed.70

Anti-plasmodial properties were exhibited by 3,5-bis(ylidene)-N-methyl-4-piperidones 69 against chloroquine-sensitive Pf3D7, chloroquine-resistant PfINDO, and artemisininresistant PfMRA-1240 strains⁷¹ (Fig. 16).

A series of N-acyl-3,5-bis(ylidene)-4-piperidones 70 were synthesized via dehydrohalogenation of the unsubstituted piperidone with the corresponding acid chloride (Scheme 31). Some of the synthesized agents revealed potent inhibitory properties against Plasmodium falciparum D6 and C235 (drug resistant) strains which also subjected to Plasmodium berghei revealing higher efficacy than chloroquine and mefloquine (standard reference drugs).72

5.7. Antiobesity active agents

1-Ethoxycarbonyl-3,5-bis[(3'-indolyl)methylene]-4-piperidones 71 were synthesized through base catalyzed condensation (piperidine in refluxing toluene) of 1-ethoxycarbonyl-4piperidine with 3-indolecarboxaldehyde followed by alkylation with various alkyl halides in refluxing DMF/K₂CO₃ (Scheme 32). Lipase inhibitory properties were revealed by the synthesized agents relative to orlistat (standard reference).73

Conclusion

Curcumin is an important natural compound with broad spectrum biological properties. Limitation of clinical application of curcumin is mainly due to its poor bio-availability in vivo. 3,5-Bis(ylidene)-4-piperidone scaffolds are considered a curcumin mimic with diverse promising bio-properties. The distinguished biological observations of curcumin mimics can be considered for optimizing high potent hits/leads accessible in drug discovery program. Many articles dealing with the biological properties of this scaffold have appeared. It has been also intensively utilized for construction of diverse potentially bioactive heterocycles.

Conflicts of interest

There is no conflict to declare.

Acknowledgements

This work was supported financially by National Research Centre, Egypt, project ID: 12060101.

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