Accepted Manuscript

Synthesis, Characterization, Crystal Structure of Novel Bis-Thiomethylcyclohexanone Derivatives and Their Inhibitory Properties Against Some Metabolic Enzymes

Abdullah Biçer, Parham Taslimi, Gül Yakalı, İlhami Gülçin, Mehmet Serdar Gültekin, Günseli Turgut Cin

PII:	S0045-2068(18)30994-5
DOI:	https://doi.org/10.1016/j.bioorg.2018.11.001
Reference:	YBIOO 2609
To appear in:	Bioorganic Chemistry
Received Date:	6 September 2018
Revised Date:	3 October 2018
Accepted Date:	1 November 2018



Please cite this article as: A. Biçer, P. Taslimi, G. Yakalı, I. Gülçin, M. Serdar Gültekin, G. Turgut Cin, Synthesis, Characterization, Crystal Structure of Novel Bis-Thiomethylcyclohexanone Derivatives and Their Inhibitory Properties Against Some Metabolic Enzymes, *Bioorganic Chemistry* (2018), doi: https://doi.org/10.1016/j.bioorg. 2018.11.001

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

Synthesis, Characterization, Crystal Structure of Novel Bis-Thiomethylcyclohexanone Derivatives and Their Inhibitory Properties Against Some Metabolic Enzymes

Abdullah Biçer^a, Parham Taslimi^b, Gül Yakalı^{c,d}, Ilhami Gülçin^{b,*},

Mehmet Serdar Gültekin^{b,e} and Günseli Turgut Cin^{a,*}

^aDepartment of Chemistry, Faculty of Science, Akdeniz University, 07058-Antalya, Turkey
^bDepartment of Chemistry, Faculty of Science, Atatürk University, 25240-Erzurum, Turkey
^cSerik Gülsün Süleyman Sural Vocational School of Higher Education, Department of Opticianry Program, Akdeniz University, 07058-Antalya, Turkey
^dCentral Research Laboratory, İzmir Katip Çelebi University, 35620-İzmir, Turkey
^eFaculty of Pharmacy, Ağrı İbrahim Çeçen University, 04100-Agrı, Turkey

*Corresponding authors

E-mail address: gturgut@akdeniz.edu.tr (G. Turgut Cin).

E-mail address: igulcin@atauni.edu.tr, Phone: +904422314375, Fax:+90 4422314109 (Ilhami Gulcin).

ABSTRACT

In this study, a series of novel bis-thiomethylcyclohexanone compounds (3a-3j) were synthesized by the addition of thio-Michael to the bis-chalcones under mild reaction conditions. The bis-thiomethylcyclohexanone derivatives (bis-sulfides) were characterized by ¹H-NMR, ¹³C-NMR, FTIR and elemental analysis techniques. Furthermore, the molecular and crystal structures of **3h**, **3i** and **3j** compounds were determined by single crystal X-ray diffraction studies. In this study, X-ray crystallography provided an alternative and oftencomplementary means for elucidating functional groups at the enzyme inhibitory site. Acetylcholinesterase (AChE) is a member of the hydrolase protein super family and has a significant role in acetylcholine-mediated neurotransmission. Here, we report the synthesis and determining of novel bis-thiomethylcyclohexanone compounds based hybrid scaffold of AChE inhibitors. The newly synthesized bis-thiomethylcyclohexanone compounds showed K_i values of in range of 39.14-183.23 nM against human carbonic anhydrase I isoenzyme (hCA I), 46.03-194.02 nM against human carbonic anhydrase II isoenzyme (hCA II), 4.55-32.64 nM against AChE and 12.77-37.38 nM against butyrylcholinesterase (BChE). As a result, novel bis-thiomethylcyclohexanone compounds can have promising anti Alzheimer drug potential and record novel hCA I, and hCA II enzymes inhibitor.

Keywords: bis-thiomethylcyclohexanone; carbonic anhydrase; acetylcholinesterase; crystal structure

1. INTRODUCTION

Carbon-sulfur bond formation is an important methodology used in the synthesis of sulfurcontaining natural and pharmaceutical products [1-5]. The thio-Michael addition reaction is one of the most important processes for C-S bond formation [6]. Generally, C-S bond formation is performed with the addition of a thiol, 1,4-conjugate on the acceptor α,β unsaturated carbonyl compounds by Lewis acid or deprotonation of thiol [7,8]. It is known that this method plays an important role in the synthesis of bioactive compounds [8-11]. Sulfide and bis-sulfide compounds have been reported to possess important biological activities in organic, pharmaceutical and biological applications (**Scheme 1**) [12-15].

Scheme 1

It is well known that the studies related to enzyme inhibition and activation is one of the most common biological activities. Carbonic anhydrases (CAs) are extensively distributed zinc-comprising metalloenzymes available in all life phyla, which hold pH homeostasis factor in the human body by catalyzing the carbon dioxide (CO_2) hydration action to bicarbonate (HCO3⁻) and proton (H⁺) plus other hydrolytic reactions [16-19]. Also, depending on their localization in diverse organisms, capacity, and catalytic activity to various classes of inhibitors, CAs are divided in seven genetically separate classes, α -, β -, δ -, γ -, ζ -, η - and θ -CAs [20,21]. Indeed, only the α -class is recorded to be present in human bodies, in which sisteen α -CA isoenzymes were explained which differ in their distribution in tissues, subcellular localization, and kinetic and molecular properties [22-24]. The CA isoenzymes are involved in multiple physiological and biochemical processes like calcification, acid-base regulation, ureagenesis, bone resorption, tumorigenicity, and gluconeogenesis, hence representing interesting biochemical aims for the design of CA inhibitors (CAIs) with many biological applications [25,26]. The ubiquitous hCA I isoenzyme is involved in cerebral edema, and retinal, and its inhibition action can be a valuable factor in fighting these situations. The cytosolic hCA II isozyme is involved in epilepsy, edema, glaucoma, and among others [27,28]. The clinical usage of CAIs has been established as anti-glaucoma agents, antiepileptics and diuretics. Also, they frequently used in the treatment of mountain sickness, duodenal and gastric ulcers, neurological disorders, idiopathic intracranial hypertension and osteoporosis [29,30].

On the other hand, Alzheimer's disease (AD) causes an advanced harm on the central neural mechanism, such as the decline in language, behavioral disorders, and memory loss [31,32]. Various anomalies in the brain cells including inflammation, oxidative stress, neuronal cell death and protein aggregates are relevant to the effects of the AD [33]. The

heterogeneous etiology and complexity of AD play a key role in novel drug development to avoid the progress of AD [34-36]. One significant therapeutic palliative-process to minimize the AD effects has been utilized of AChE inhibitors (AChEIs) such as rivastigmine, donepezil, tacrine, and galantamine. Recently, it was reported that some identified AChEIs as promising molecules to treat dementia, withdrawned from the clinical trials due to their toxicity and poor central nervous system stability [37,38]. Therefore, the design, discovery and development of novel AChEIs are still a big challenge for medicinal chemist. Thus, the selectivity of the inhibition has a great importance and presents a significant consideration for developing novel kinds of AChEIs [39-42].

In this study, we synthesized new bis-sulfide derivatives using bisbenzylidenecyclohexanone (bis-chalcone) and evaluated their biological activites. Also, we aimed to investigate the inhibitory effect of these novel bis-thiomethylcyclohexanone derivatives (**3a-j**) using bis-benzylidenecyclohexanone (**2a-j**) derivatives on the hCA I, hCA II, AChE and BChE enzymes for alternative and compared to standard and clinically used inhibitors.

2.RESULTS AND DISCUSSION

2.1. Chemistry

Firstly, the bis-chalcone compounds were synthesized according to previous studies [43,44]. For this purpose, the substituted aromatic aldehydes and cyclohexanones were subjected to the Claisen-Schmidt condensation reaction in ethanol with potassium hydroxide (Scheme 2). Then, suitable reaction conditions for the synthesis of bis-sulfide compounds were determined. For this, the reaction of addition of thiophenol to bis-chalcone using different Lewis acids and bases was studied and the most suitable reaction condition was found to be metallic Na in DCM (Table 1).

Scheme 2 and Table 1

TLC detected the product mixture in experiments with iodine. However, any products were not isolated from this mixture. Also, in the experiments with other catalysts (Table 1, entry 3-7), product formation was not observed. A single product was formed in the experiments with sodium at room temperature in DCM. This compounds was detected by TLC and purified by crystallization (Table 1). The bis-chalcone based sulfides were synthesized by reacting bis-chalcones with thiophenol in the presence of sodium-metal in dichloromethane at room temperature via thio-Michael addition reaction (Scheme 3). Spectrometric data (¹H-NMR,

¹³C-NMR, FTIR and elemental analysis) and physical properties (color and melting point) of these products have been confirmed by comparison with similar studies reported in the literature [45,46].

Scheme 3

2.2. Enzyme inhibition studies

Evaluation of the effects of novel bis-sulfide derivatives (3a-j) on both cholinergic enzymes of AChE and BChE, and both hCA isoenzymes was the main objective of this study. The inhibition results are summarised in Table 2 and related Figures 1 and 2. The cytosolic hCA I, and II isoenzymes are present in the human body and are involved in the secretion of electrolytes in a plenty of tissue cells, like the HCO₃⁻ rich aqueous humor in the anterior chamber of the cerebrospinal fluid or the eyes, as well as hold CO₂ homeostasis and pH all the body [47,48]. Indeed, dysregulation of hCA I, and II isoenzymes in tissue cells lead to some pathologic conditions including edema, glaucoma and epilepcy [49,50]. For hCA I isozyme, the Ki values were found in range of 39.14±9.94-183.23±28.18 nM. In this study, the Ki for the positive control CA inhibitor acetazolamide (AZA), a recorded hCA I inhibitor, was 273.61±76.86 nM against hCA I isoenzyme (Table 2). All novel bis-sulfide compounds (3a-j) had effective inhibition effects against hCA I isoenzyme. In addition, between these novel molecules, 2,6-bis((4-chlorophenyl)(phenylthio)methyl)cyclohexanone (3h) and 2,6-bis((3bromophenyl)(phenylthio)methyl)cyclohexanone (3i), which possessed chlorophenyl and bromophenyl groups demonstrated the best hCA I inhibitor (Kis of 39.14±9.94 and 47.92±7.20 nM, respectively) (Figure 1). It is well known that the molecules comprising chlorophenyl and bromophenyl groups are efficient CA inhibitors [51]. As seen in Table 2, IC₅₀ values were found in the range of 38.63-180.34 nM towards hCA I, and 42.84-201.58 nM for hCA II. Novel bis-sulfide derivatives (3a-j) synthesized in this study significantly inhibited the slow cytosolic hCA II isozyme with K_i in the low nanomolar levels. Ki values were obtained between 46.03±10.47 and 194.02±28.55 nM (Table 2). Also, the compounds of 3h and 3i are, in fact, the best inhibitor among these molecules (Ki 46.03±10.47 and 48.18±19.10 nM, respectively) compared to the AZA, which is known as specific inhibitor of CA isoenzymes as a standard (Ki: 229.08±55.14 nM). For hCA I isoform, its excellent inhibitors were (**3h**) and (**3i**), which were the best hCA I inhibitor with IC₅₀: 38.63 nM (r^2 : 0.9812) 44.28 nM (r²: 0.9590), respectively. and IC₅₀: 2,6-Bis((3nitrophenyl)(phenylthio)methyl) cyclohexanone (3c) compound had relativelly weak

inhibition effects when compared to other compounds for this isoenzyme IC₅₀: 180.34 nM (r^2 : 0.9598) for hCA I, IC₅₀: 201.58 nM (r^2 : 0.9902) for hCA II isoenzyme.

Table 2, Figure 1 and 2

In this work, AChE and BChE were also extremely inhibited by novel bis-sulfide compounds (3a-j) at the nanomolar range inhibition. Ki values of this novel compounds were found in the range of 4.55±0.94-32.64±6.86 nM for AChE and 12.77±2.20-37.38±11.53 nM for AChE (Table 2). These inhibition results clearly designated that novel synthesized compounds had efficient both cholinergic enzymes inhibition properties. Indeed, the most potent AChE and BChE inhibition were obtained by novel compound 2,6-bis((3fluorophenyl)(phenylthio)methyl)cyclohexanone (3e) with Ki values of 4.55±0.94 and 12.77±2.20 nM, respectively (Figure 1). Also, tacrine (1,2,3,4-tetrahydroacridin-9-amine), which the first centrally acting cholinesterase inhibitor approved for the treatment of AD, possesed Ki values of 56.37±15.10 and 63.40±13.62 nM against both cholinergic AChE and BChE enzymes, respectively. Eventually, the K_i values of these derivatives for AChE and BChE were calculated from Lineweaver-Burk plots. Additionally, as seen in Table 2, IC₅₀ values are in the range of 7.18-41.77 nM towards AChE and in the range of 17.90-61.38 nM for BChE. 2,6-Bis((4-nitrophenyl)(phenylthio)methyl)cyclohexanone (3d) is weak inhibitor compared to other compounds for these enzymes. IC₅₀ were found as 41.77 nM (r²:0.9732) for AChE and 61.38 nM for BChE ($r^2:0.9682$).

CA inhibitors targeting hCAs are clinically used in recent years for the management of diverse diseases among which obesity, glaucoma, intracranial hypertension epilepsy, and as diuretics [52]. Recently, they started to be utilized for the therapy of hypoxic tumours, were also evaluated as possible drugs for cerebral ischemia, neuropathic pain, and arthritis. Additionally, plenty of work has been done on the synthesis and characterization of CAIs belonging to diverse groups, such as coumarins, carboxylic acids, sulfonamides, phenols, dithiocarbamates, heterocyclic derivatives, etc [53]. Indeed, sulfonamides and their bioisosteres such as the sulfamides and the sulfamates are powerful active site coordinating CAIs, which in deprotonated form, bind to the Zn²⁺ ion present within the active site of enzyme [54]. Many sulfonamide-based drugs, such as methazolamide, acetazolamide, dorzolamide, ethoxzolamide, brinzolamide and celecoxib are recorded. Some of them are in clinical use as antiepileptics (targeting hCAs II, IV and XII), or in clinical trials as antitumor or antimetastatic agents (targeting hCAs IX and XII) [55,56].

Many researches have recorded that by raising ACh level through inhibiting AChE and BChE enzymes in the body offer a potential treatment for therapy AD and hence improve both memory and mental symptoms in that disease [57]. Control AChEIs such as galantamine, donepezil, tacrine, and rivastigmine are involved in improving symptoms for most patients by enhancing cholinergic neurotransmission range in the human body [58]. Thus, these inhibitors are expensive as well as they are long-term utilize give rise to diverse harmful side effects and adverse of symptoms. Recently, naturally occurring BChE and AChE inhibitors have been isolated from the animals and plants and utilized as the natural treatment for the AD [59-61]. In this study, novel bis-sulfide compounds (**3a-j**) have been reported as potent both cholinesterase enzymes inhibitors.

2.3 Crystallographic studies

Thermal ellipsoid views of the **3h**, **3i** and **3j** are shown in Figure 3. Selected bond lengths and angles are illustrated in Table 3, mostly consistent with the similar molecules in the literature [62-65].

Figure 3 and Table 3

As depicted in Figure 3, the asymmetric unit of the molecule **3h** has one-half-molecule and it is completed with a twofold symmetry axis [symmetry code: x, 1–y, z]. Compound **3h** crystallizes in a monoclinic system with space group *lm* (Table 4). The dihedral angle between the phenylthio ring and cyclohexanone ring is $60.0 (3)^\circ$ while the phenylthio rings are inclined to chlorophenyl ring by 34.8 (3)°. The cyclohexanone ring is nearly planar to the molecule plane making dihedral angle is $10.4 (2)^\circ$. In the crystal structure, there is only weak intramolecular interaction (Table 5). The packing diagram of the molecule is formed by stacking interactions along the *a* axis (Figure 4).

Tables 4 and 5

Compound **3i** crystallizes in a triclinic system with space group P-1 (Table 4). The dihedral angle between the phenylthio rings and bromophenyl rings are 25.2(4)° (left groups), 52.8(3)° (right groups). Phenylthio ring (C1/C6) is nearly perpendicular to molecule plane with 87.1(3)°. The dihedral angle between bromophenyl ring (C21/C26) and cyclohexanone ring is 87.3(3)° while the another ring (C8/C13) are inclined to cyclohexanone ring by 57.8 (4)°. In addition, phenylthio rings are nearly perpendicular cyclohexanone ring with 82.8(4)° and 77.2(4)°. In the crystal structure, molecules are linked by strong intermolecular C–H···S hydrogen bond to form an infinite chain along the *b* axis. This hydrogen bond also generates

 $R_2^2(10)$ ring motif with dimeric structure (Figure 4) [65]. Moreover, there is a strong C–H…pi interaction between the C25 atom of the bromophenyl ring and another bromophenyl ring [Cg: C8/C9/C10/C11/C12/C13; C25–Cg 3.374(9) Å, H25…Cg 2.803(9) Å, C25–H25…Cg 120.7(8)°, symmetry code: 1–x, 1–y, 1–z].

Figure 4

Compound **3j** crystallizes in a monoclinic system with Im space group (Table 4). The molecule has crystallographic mirror symmetry with x, 1-y, z symmetry operator. The dihedral angle between the phenylthio ring and cyclohexanone ring is 57.6 (2)° while the phenylthio rings are inclined to bromophenyl ring by 33.3 (2)°. The cyclohexanone ring is nearly planar to the molecule plane making dihedral angle is 9.36 (14)°. The packing structure of the **3j** is stabilised by stacking interactions along the *b* axis (Figure 4).

The expected absolute configuration for molecule 3h and 3j were confirmed by refinement of the Flack parameters (its value 0.01(11) A° for 3h, -0.027(9) A° for 3j). Details of the molecular geometry having two stereogenic centers which is C5 atoms in molecule 3h and 3j, C7 and C20 atoms in molecule 3i reveal that there are two enantiomeric forms (*R* and *S*) named with respect to the majority of chiral centers in the crystal structure of the compounds.

To examine the chemical details of enzymes function, the first step is to determine what functional groups are required for enzyme activity and to ascertain these functional groups [66]. In this study, X-ray crystallography provides an alternative and oftencomplementary means for elucidating functional groups at the enzyme active site. According to enzyme inhibition studies, comprising chlorophenyl and bromophenyl groups in 3h and 3i respectively are efficient both CA isoenzymes inhibitors. The X-ray crystallography reveals that the electron density of these groups in compounds less than similar groups in molecule 3j. Therefore, the enzymes inhibition effects can be directly related to electron density of binding site of molecules [67].

3. CONCLUSION

The novel bis-thiomethylcyclohexanone compounds were investigated for AChE, BChE, hCA I, and hCA II enzymes inhibition effects. As we explained above, novel bis-sulfide compounds (**3a-j**) can be good candidate drugs, the same as AChE, BChE, and carbonic anhydrase inhibitor compounds, for therapy of some diseases like AD, epilepsy, glaucoma, gastric and duodenal ulcers, mountain sickness, osteoporosis or neurological disorders.

4. EXPERIMENTALS

4.1. General

All chemicals were obtained from Sigma-Aldrich, Fluka, Merck or Alfa Aesar and were used without further purification. Melting points were determined on a Gallenkamp melting point apparatus. FTIR spectra were recorded on Bruker Tensor27 FTIR spectrometer using the KBr disc in the range of 4000-400 cm⁻¹. ¹H-NMR and ¹³C-NMR spectra were recorded on a Varian 200 MHz or Bruker Avance DPX 400 MHz spectrometer using CDCl₃ as a solvent. Splitting patterns are designated as follows; s, singlet; d, doublet; m, multiplet. Chemical shift (δ) values are given in ppm. Elemental analysis was done on a LECO CHNS-932 instrument.

4.2. Chemistry

Bis-chalcones (**2a-j**) were synthesized following literature procedures by the condensation of cyclohexanone with aromatic aldehydes (**1a-j**) [43,44].

4.2.1. General procedure for synthesis of bis-thiomethylcyclohexanone derivatives (3a-j)

Bis-chalcone (1 eq.) was dissolved in 20 mL of DCM and Na (catalitic amount) added. After five minutes shaking in DCM, to reaction mixture was slowly added thiophenol (2.4 eq.) and the reaction was stirred at room temperature overnight. After completion of reaction, the mixture evaporated in vacuo. Pure bis-sulfide derivatives were obtained by recrystallization from methanol.

2,6-Bis(phenyl(phenylthio)methyl)cyclohexanone (**3a**). Yield 62%; White solid; mp: 168-169°C; FT-IR (KBr, cm⁻¹) v: 3040-3025 (arom. C-H stretch.), 2958-2861 (alif. C-H stretch.), 1724 (C=O stretch.), 1622 (C=C stretch.), 780-700 (C-S stretch.); ¹H-NMR(400 MHz, CDCl₃, δ, ppm): 7.19-7.06 (m, 20H), 4.67 (d, *J*=7.6 Hz, 2H, CH-S), 2.89 (m, 2H, CH-C=O), 2.65 (m, 2H), 2.10 (m, 2H), 1.70-1.68 (m, 2H); ¹³C-NMR(100 MHz, CDCl₃, δ, ppm): 208.6, 142.1, 134.9, 132.6, 128.8, 128.2, 128.1, 127.2, 126.8, 57.5, 52.9, 33.1, 25.6; Elemental Analysis Calculated: C, 77.73; H, 6.07; S, 12.96. Found: C, 77.64; H, 6.12; S, 12.83.

2,6-Bis(naphthalen-2-yl(phenylthio)methyl)cyclohexanone (3b). Yield 43%; White solid; mp 166-167°C; FT-IR (KBr, cm⁻¹) v: 3051(arom. C-H stretch.), 2923-2854 (alif. C-H stretch.), 1707 (C=O stretch.), 1630 (C=C stretch.), 742-692 (C-S stretch.); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.63-7.00 (m, 24H), 4.79 (d, *J*= 7.6 Hz, 2H, CH-S), 3.00-2.97 (m, 2H, CH-C=O), 2.73-2.60 (m, 2H), 2.07-2.05 (m, 2H), 1.76-1.72 (m, 2H); ¹³C-NMR (100 MHz,

CDCl₃, δ, ppm): 208.3, 139.5, 134.9, 133.1, 132.5, 128.8, 128.0, 127.9, 127.6, 127.2, 126.8, 126.4, 126.0, 125.6, 57.6, 53.0, 33.2, 25.6; Elemental Analysis Calculated: C, 80.80; H, 5.72; S, 10.77. Found: C, 80.83; H, 5.80; S, 10.68.

2,6-Bis((3-nitrophenyl)(phenylthio)methyl)cyclohexanone (3c). Yield 36%; white solid; mp 138-139°C; FT-IR (KBr, cm⁻¹) v: 3081 (arom. C-H), 2943-2895 (alif. C-H), 1710 (C=O), 1534 (C=C), 1450 (NO₂ asym.), 1350 (NO₂ sym.), 732-687 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.78 (d, *J*=8.0 Hz, 2H), 7.69 (s, 2H), 7.23 (d, *J*=8.0 Hz, 2H), 7.20-7.00 (m, 12H,), 4.44 (d, *J*= 9.6 Hz, 2H, CH-S), 3.00-2.97 (m, 2H), 2.90-2.87 (m, 2H), 2.15-2.00 (m, 2H), 1.91-1.88 (m, 1H), 1.70-1.60 (m, 1H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 209.3, 147.8, 144.6, 134.3, 133.8, 132.6, 129.2, 128.7, 128.4, 122.5, 121.7, 57.3, 52.7, 35.1, 25.5; Elemental Analysis Calculated: C, 65.75; H, 4.79; S, 10.96. Found: C, 65.82; H, 4.90; S, 10.95.

2,6-Bis((4-nitrophenyl)(phenylthio)methyl)cyclohexanone (3d). Yield 44%; White solid; mp 162-164°C; FT-IR (KBr, cm⁻¹) v: 3083-3075 (arom. C-H), 2924-2857- (alif. C-H), 1711 (C=O), 1599 (C=C), 1519 (NO₂ asym.), 1347 (NO₂ sym.), 749-694 (C-S); ¹H-NMR(400 MHz, CDCl₃, δ, ppm): 7.86 (d, *J*=8.4 Hz, 4H), 7.31-7.05 (m, 14H), 4.47 (d, *J*=8.4 Hz, 2H, CH-S), 2.98-2.92 (m, 2H, CH-C=O), 2.81-2.79 (m, 2H), 2.15-2.12 (m, 2H), 1.86-1.71 (m, 2H); ¹³C-NMR(100 MHz, CDCl₃, δ, ppm): 209.2, 149.8, 146.0, 133.7, 132.6, 129.2, 128.8, 128.4, 123.3, 57.0, 53.0, 34.1, 25.5; Elemental Analysis Calculated: C, 65.75; H, 4.79; S, 10.96. Found: C, 65.82; H, 4.92; S, 11.00.

2,6-Bis((3-fluorophenyl)(phenylthio)methyl)cyclohexanone (3e). Yield 40%; White solid; mp 168°C; FT-IR (KBr, cm⁻¹) v: 3056-3035 (arom. C-H), 2924-2855 (alif. C-H), 1697 (C=O), 1590 (C=C), 1146 (Ar-F), 763-690 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.16 (m, 10H), 6.99 (m, 2H), 6.77 (m, 2H), 6.75 (m, 2H), 6.73 (m, 2H), 4.50 (d, *J*= 8.0 Hz, 2H, CH-S), 2.93-2.86 (m, 2H, CH-C=O), 2.73-2.70 (m, 2H), 2.08-2.04 (m, 1H), 1.75-1.63 (m, 3H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 208.6, 163.9, 145.0, 144.7, 133.1, 129.4, 128.9, 127.7, 123.8, 114.9, 113.7, 57.4, 52.7, 33.9, 25.5; Elemental Analysis Calculated: C, 72,45; H, 5.28; S, 12.08. Found: C, 72.60; H, 5.43; S, 12.20.

2,6-Bis((4-fluorophenyl)(phenylthio)methyl)cyclohexanone (3f). Yield 38%; White solid; mp 144-145°C; FT-IR (KBr, cm⁻¹) v: 3068-3054 (arom. C-H), 2956-2857 (alif. C-H), 1706

(C=O), 1603 (C=C), 1226 (Ar-F), 750-690 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.19-7.09 (m, 10H), 6.97 (m, 4H), 6.72 (t, *J*= 8.4 Hz, 4H), 4.46 (d, *J*= 8.8 Hz, 2H, CH-S), 2.90-2.83 (m, 2H, CH-C=O), 2.73-2.70 (m, 2H), 2.07-2.03 (m, 1H), 1.75-1.60 (m, 3H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 208.4, 160.4, 137.9, 133.1, 129.6, 128.9, 127.6, 115.0, 114.8, 57.8, 52.3, 34.2, 25.6; Elemental Analysis Calculated: C, 72,45; H, 5.28; S, 12.08. Found: C, 72.53; H, 5.42; S, 11.95.

2,6-Bis((3-chlorophenyl)(phenylthio)methyl)cyclohexanone (3g). Yield 35%; White solid; mp 115-116°C; FT-IR (KBr, cm⁻¹) v: 3064-3058 (arom. C-H), 2948-2857 (alif. C-H), 1709 (C=O), 1570 (C=C), 1072 (Ar-Cl), 745-688 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.20-7.10 (m, 10H), 7.06 (m, 2H), 7.02 (m, 2H), 6.98 (m, 2H), 6.88 (m, 2H), 4.48 (d, *J*= 8.0 Hz, 2H, CH-S), 2.91-2.85 (m, 2H, CH-C=O), 2.71-2.68 (m, 2H), 2.08-2.04 (m, 1H), 1.77-1.62 (m, 3H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 208.5, 144.2, 134.0, 133.1, 129.3, 129.0, 128.1, 127.7, 127.1, 126.3, 57.3, 52.7, 33.7, 25.5; Elemental Analysis Calculated: C, 68.21; H, 4.97; S, 11.37. Found: C, 68.25; H, 5.12; S, 11.29.

2,6-*Bis((4-chlorophenyl)(phenylthio)methyl)cyclohexanone (3h).* Yield 52%; White solid; decomposition after 240°C; FT-IR (KBr, cm⁻¹) v: 3068-3054 (arom. C-H), 2926-2857 (alif. C-H), 1704 (C=O), 1600 (C=C), 1091 (Ar-Cl), 746-688 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.15 (m, 10H), 7.02 (d, *J*=8.8 Hz, 4H), 6.93 (d, *J*=8.4 Hz, 4H), 4.46 (d, *J*= 8.4 Hz, 2H, CH-S), 2.90-2.83 (m, 2H, CH-C=O), 2.74-2.70 (m, 2H), 2.08-2.03 (m, 1H), 1.74-1.61 (m, 3H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 209.1, 140.7, 134.0, 133.1, 132.5, 129.3, 129.0, 128.2, 127.7, 57.7, 52.4, 34.2, 25.6; Elemental Analysis Calculated: C, 68.21; H, 4.97; S, 11.37. Found: C, 68.12; H, 5.13; S, 11.22.

2,6-*Bis((3-bromophenyl)(phenylthio)methyl)cyclohexanone (3i)*. Yield 36%; White solid; mp 120-121°C; FT-IR (KBr, cm⁻¹) v: 3062-3058 (arom. C-H), 2917-2856 (alif. C-H), 1709 (C=O), 1566 (C=C), 1069 (Ar-Br), 747-688 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.26-6.91 (m, 18H), 4.49 (d, *J*= 8.4 Hz, 2H, CH-S), 2.91-2.85 (m, 2H, CH-C=O), 2.71-2.68 (m, 2H), 2.07-2.00 (m, 1H), 1.74-1.62 (m, 3H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 208.5, 144.5, 134.0, 133.1, 131.0, 130.0, 129.7, 129.0, 127.8, 126.8, 122.3, 57.3, 52.7, 33.7, 25.5; Elemental Analysis Calculated: C, 58.90; H, 4.29; S, 9.82. Found: C, 58.84; H, 4.33; S, 9.96.

2,6-Bis((4-bromophenyl)(phenylthio)methyl)cyclohexanone (3j). Yield 48%; White solid; mp 166-167°C; FT-IR (KBr, cm⁻¹) v: 3066-3052 (arom. C-H), 2937-2853 (alif. C-H), 1706 (C=O), 1577 (C=C), 1068 (Ar-Br), 751-688 (C-S); ¹H-NMR (400 MHz, CDCl₃, δ, ppm): 7.18-6.89 (m, 14H), 6.87 (d, *J*= 8.4 Hz, 4H), 4.44 (d, *J*= 8.8 Hz, 2H, CH-S), 2.89-2.82 (m, 2H, CH-C=O), 2.74-2.70 (m, 2H), 2.07-2.00 (m, 1H), 1.77-1.60 (m, 3H); ¹³C-NMR (100 MHz, CDCl₃, δ, ppm): 209.1, 141.2, 133.9, 133.1, 131.2, 129.7, 129.0, 127.7, 120.6, 57.6, 52.4, 34.2, 25.6; Elemental Analysis Calculated: C, 58.90; H, 4.29 S, 9.82. Found: C, 58.86; H, 4.42; S, 9.76.

4.3. X-ray Crystallography

Single crystal X-ray diffraction data of **3h**, **3i** and **3j** were collected at room temperature on an Rigaku-Oxford Xcalibur diffractometer with an Eos-CCD detector, operated at 50 kV and 40 mA using graphite monochromated Mo-K α ($\lambda = 0.71073$ Å) radiation. Utilising CrysAlis^{*Pro*} software package, data collections and reductions along with absorption corrections were performed [68]. Structure solutions were performed using SHELXT [69] embedded in the Olex2 [70]. Refinement of coordinates and anisotropic thermal parameters of non-hydrogen atoms were carried out by the full-matrix leastsquares method in SHELXL [71]. For the compounds, all hydrogen atoms were placed in geometrically idealized positions (C-H=0.97 A° for methylene groups and C-H=0.93 A° for aromatic groups). Detailed crystallographic data and refinement results are summarized in Table 4.

4.4. Biochemical Studies

Both hCA isoenzymes inhibition effects of novel bis-sulfide compounds (**3a-j**) was measured according to Verpoorte et al. [72] conforming to previous studies [73-77] and recorded at 348 nm spectrophotometrically using p-nitrophenylacetate substrate (PNA) [78]. On the other hand, AChE and BChE inhibitory effects of novel bis-sulfide compounds (**3a-j**) were determined according to procedure of Ellman et al. [79] conforming to previous studies [80-85] and recorded at 412 nm spectrophotometrically using acetylthiocholine iodide and butyrylthiocholine iodide as substrates for the enzymatic reaction. 5,5'-Dithio-bis(2-nitrobenzoic) acid compound was used for the measurement of the AChE and BChE activities, respectivelly [86].

Supplementary material

Crystallographic data as .cif files for the structures reported in this paper have been deposited at the Cambridge Crystallographic Data Center with CCDC 1865288, 1865295 and 1865296 for the molecule **3h**, **3i** and **3j** respectively. Copies of the data can be obtained free of charge at

www.ccdc.cam.ac.uk/conts/retrieving.html (or from the Cambridge Crystallographic Data Center, 12, Union Road, Cambridge CB2 1EZ, UK). Email: deposit@ccdc.cam.ac.uk.

Acknowledgement: The authors acknowledge Scientific Research Projects Coordination Unit of Akdeniz University (Project Number FDK-2016-1541) for financial support and Faculty of Science, Ataturk University.

REFERENCES

- Z. Qiao, X. Jiang, Recent developments in sulfur-carbon bond formation reaction involving thiosulfates, Org. Biomol. Chem. 15 (2017) 1942-1946.
- [2] K. L. Dunbar, D. H. Scharf, A. Litomska, C. Hertweck, Enzymatic carbon-sulfur bond formation in natural product biosynthesis, Chem. Rev. 117 (2017) 5521-5577.
- [3] E. H. Krenske, R. C. Petter, K. N. Houk, Kinetics and thermodynamics of reversible thiol additions to mono- and diactivated michael acceptors: Implications for the design of drugs that bind covalently to cysteines, J. Org. Chem. 81 (2016) 11726-11733.
- [4] M. Feng, B. Tang, S.H. Liang, X. Jiang, Sulfur containing scaffolds in drugs: Synthesis and application in medicinal chemistry, Curr. Top. Med. Chem. 16 (2016) 1200-1216.
- [5] D. Enders, K. Luettgen, A.A. Narine, Asymmetric sulfa-Michael additions, Synthesis 7 (2007) 959-980.
- [6] P. Espeel, F.E. Du Prez, One-pot multi-step reactions based on thiolactone chemistry: A powerful synthetic tool in polymer science, Eur. Polym. J. 62 (2015) 247-272.
- [7] J.P. Phelan, J.A. Ellman, Conjugate addition-enantioselective protonation reactions, Beilstein J. Org. Chem. 12 (2016) 1203-1228.
- [8] R. Dalpozzo, G. Bartoli, G. Bencivenni, Asymmetric organocatalytic reactions of α,β-unsaturated cyclic ketones, Symmetry 3 (2011) 84-125.
- [9] N. Azizi, A. Khajeh-Amiri, H. Ghafuri, M. Bolourtchian, A highly efficient, operationally simple and selective thia-Michael addition under solvent-free condition, Green Chem. Lett. Rev. 2 (2009) 43-46.
- [10] G. Yerli, H. Gezegen, M. Ceylan, Iodine-catalyzed addition of 2-mercaptoethanol to chalcone derivatives: Synthesis of the novel β-mercapto carbonyl compounds, Org. Commun. 5 (2012) 70-76.
- [11] S.S. Pandit, S.S. Pardhe, Y.B. Pandit, Guanidine hydrochloride: An efficient catalyst for the synthesisof 2-hydrazolyl-4-thiazolidinone derivatives under solvent free conditions, Iran. Chem. Commun. 5 (2017) 227-236.
- [12] B.L. Zhao, D.M. Du, Enantioselective squaramide-catalyzed trifluoromethylthiolation-sulfur-Michael/Aldol cascade reaction: One-pot synthesis of CF3S containing spiro cyclopentanonethiochromane, Org. Lett. 19 (2017) 1036-1039.

- [13] X. Gu, Y.Z. Zhu, Therapeutic applications of organosulfur compounds as novel hydrogen sulfide donors and/or mediators, Expert Rev. Clin. Pharmacol. 4 (2011) 123-133.
- [14] M. Feng, B. Tang, S.H. Liang, X. Jiang, Sulfur containing scaffolds in drugs: synthesis and application in medicinal chemistry, Curr. Top. Med. Chem. 16 (2016) 1200-1216.
- [15] S. Parcell, Sulfur in human nutrition and applications in medicine, Altern. Med. Rev. 7 (2002) 22-44.
- [16] A. Akıncıoğlu, Y. Akbaba, H. Göçer, S. Göksu, İ. Gülçin, C.T. Supuran, Novel sulfamides as potential carbonic anhydrase isoenzymes inhibitors, Bioorg. Med. Chem. 21 (2013) 1379-1385.
- [17] K. Aksu, M. Nar, M. Tanç, D. Vullo, İ. Gülçin, S. Göksu, F. Tümer, C.T. Supuran, Synthesis and carbonic anhydrase inhibitory properties of sulfamides structurally related to dopamine, Bioorg. Med. Chem. 21 (2013) 2925-2931.
- [18] M. Güney, A. Coşkun, F. Topal, A. Daştan, İ. Gülçin, C.T. Supuran, Oxidation of cyanobenzocycloheptatrienes: Synthesis, photooxygenation reaction and carbonic anhydrase isoenzymes inhibition properties of some new benzotropone derivatives, Bioorg. Med. Chem. 22 (2014) 3537-3543.
- [19] S. Göksu, A. Naderi, Y. Akbaba, P. Kalın, A. Akıncıoğlu, İ. Gulcin, S. Durdaği, R.E. Salmas, Carbonic anhydrase inhibitory properties of novel benzylsulfamides using molecular modeling and experimental studies, Bioorg. Chem. 56 (2014) 75-82.
- [20] M. Boztaş, Y. Çetinkaya, M. Topal, İ. Gülçin, A. Menzek, E. Şahin, M. Tanc, C.T. Supuran, Synthesis and carbonic anhydrase isoenzymes I, II, IX, and XII inhibitory effects of dimethoxybromophenol derivatives incorporating cyclopropane moieties, J. Med. Chem. 58 (2015) 640-650.
- [21] A. Yıldırım, U. Atmaca, A. Keskin, M. Topal, M. Çelik, İ. Gülçin, C.T. Supuran, N-Acylsulfonamides strongly inhibit human carbonic anhydrase isoenzymes I and II, Bioorg. Med. Chem. 23 (2015) 2598-2605.
- [22] A. Akıncıoğlu, H. Akıncıoğlu, I. Gülçin, S. Durdağı, C.T. Supuran, S. Göksu, Discovery of potent carbonic anhydrase and acetylcholine esterase inhibitors: Novel sulfamoylcarbamates and sulfamides derived from acetophenones, Bioorg. Med. Chem. 23 (2015) 3592-3602.
- [23] P. Taslimi, İ. Gulcin, B. Ozgeris, S. Goksu, F. Tumer, S.H. Alwasel, C.T. Supuran, The human carbonic anhydrase isoenzymes I and II (hCA I and II) inhibition effects of trimethoxyindane derivatives, J. Enzyme Inhib. Med. Chem. 31 (2016) 152-157.
- [24] M. Küçük, İ. Gulcin, Purification and characterization of carbonic anhydrase enzyme from black sea trout (*Salmo trutta* Labrax Coruhensis) kidney and inhibition effects of some metal ions on the enzyme activity, Environ. Toxicol. Pharmacol. 44 (2016) 134-139.
- [25] P. Taslimi, İ. Gülçin, N. Öztaşkın, Y. Çetinkaya, S. Göksu, S.H. Alwasel, C.T. Supuran, The effects of some bromophenol derivatives on human carbonic anhydrase isoenzymes, J. Enzyme Inhib. Med. Chem. 31 (2016) 603-607.

- [26] D. Ozmen Ozgun, C. Yamali, H.İ. Gül, P. Taslimi, İ. Gülçin, T. Yanik, C.T. Supuran, Inhibitory effects of isatin mannich bases on carbonic anhydrases, acetylcholinesterase and butyrylcholinesterase, J. Enzyme Inhib. Med. Chem. 31 (2016) 1498-1501.
- [27] A. Sujayev, E. Garibov, P. Taslimi, İ. Gülçin, S. Gojayeva, V. Farzaliyev, S.H. Alwasel, C.T. Supuran, Synthesis of some tetrahydropyrimidine-5-carboxylates, determination of their metal chelating effects and inhibition profiles against acetylcholinesterase, butyrylcholinesterase and carbonic anhydrase, J. Enzyme Inhib. Med. Chem. 31 (2016) 1531-1539.
- [28] K. Aksu, B. Özgeriş, P. Taslimi, A. Naderi, İ. Gülçin, S. Göksu, Antioxidant activity, acetylcholinesterase and carbonic anhydrase inhibitory properties of novel ureas derived from phenethylamines, Arch. Pharm. 349 (2016) 944-954.
- [29] M. Huseynova, P. Taslimi, A. Medjidov, V. Farzaliyev, M. Aliyeva, G. Gondolova, O. Şahin, B. Yalçın, A. Sujayev, E.B. Orman, A.R. Özkaya, İ. Gülçin, Synthesis, characterization, crystal structure, electrochemical studies and biological evaluation of metal complexes with thiosemicarbazone of glyoxylic acid, Polyhedron 155 (2018) 25-33.
- [30] M. Tuğrak, H.İ. Gül, H. Sakagami, İ. Gülçin, C.T. Supuran, New azafluorenones with cytotoxic and carbonic anhydrase inhibitory properties: 2-Aryl-4-(4-hydroxyphenyl)-5H-indeno[1,2b]pyridin-5-ones, Bioorg. Chem. 81 (2018) 433-439.
- [31] F. Topal, İ. Gulcin, A. Dastan, M. Guney, Novel eugenol derivatives: potent acetylcholinesterase and carbonic anhydrase inhibitors, Int. J. Biol. Macromol. 94 (2017) 845-851.
- [32] M. Işık, S. Beydemir, A. Yılmaz, M.E. Naldan, H.E. Aslan, İ. Gülçin, Oxidative stress and mRNA expression of acetylcholinesterase in the leukocytes of ischemic patients, Biomed. Pharmacother. 87 (2017) 561-567.
- [33] U.M. Kocyigit, Y. Budak, M.B. Gürdere, Ş. Tekin, T. Kul Köprülü, F. Ertürk, K. Özcan, İ. Gülçin, M. Ceylan, Synthesis, characterization, anticancer, antimicrobial and carbonic anhydrase inhibition profiles of novel (3aR,4S,7R,7aS)-2-(4-((E)-3-(3-aryl)acryloyl) phenyl)-3a,4,7,7a-tetrahydro-1H-4,7-methanoisoindole-1,3(2H)-dione derivatives, Bioorg. Chem. 70 (2017) 118-125.
- [34] H.I. Gül, A. Demirtas, G. Ucar, P. Taslimi, İ. Gülçin, Synthesis of Mannich bases by two different methods and evaluation of their acetylcholine esterase and carbonic anhydrase inhibitory activities, Lett. Drug Des. Discov. 14 (2017) 573-580.
- [35] Ç. Bayrak, P. Taslimi, İ. Gülçin, A. Menzek, The first synthesis of 4-phenylbutenone derivative bromophenols including natural products and their inhibition profiles for carbonic anhydrase, acetylcholinesterase and butyrylcholinesterase enzymes, Bioorg. Chem. 72 (2017) 359-366.
- [36] A. Aktaş, P. Taslimi, I. Gülçin, Y. Gök, Novel NHC Precursors: Synthesis, characterization and carbonic anhydrase and acetylcholinesterase inhibitory properties, Arch. Pharm. 350 (2017) e1700045.

- [37] M. Zengin, H. Genç, P. Taslimi, A. Kestane, E. Güçlü, A. Ögütlü, O. Karabay, İ. Gülçin, Novel thymol bearing oxypropanolamine derivatives as potent some metabolic enzyme inhibitors-their antidiabetic, anticholinergic and antibacterial potentials. Bioorg. Chem. 81(2018) 119-126.
- [38] K. Aksu, H. Akıncıoğlu, A. Akıncıoğlu, S. Göksu, F. Tümer, İ. Gulçin, Synthesis of novel sulfamides incorporating phenethylamines and determination of their inhibition profiles against some metabolic enzymes, Arch. Pharm. 351(9) (2018) e1800150.
- [39] P. Taslimi, A. Sujayev, E. Garibov, N. Nazarov, Z. Huyut, S.H. Alwasel, İ. Gülçin, The synthesis of new cyclic thioureas and evaluation of their metal-chelating activity, acetylcholinesterase, butyrylcholinesterase and carbonic anhydrase inhibition profiles, J. Biochem. Mol. Toxicol. 31 (2017) e21897.
- [40] N. Öztaşkın, P. Taslimi, A. Maraş, S. Göksu, İ. Gülçin, Novel antioxidant bromophenols with acetylcholinesterase, butyrylcholinesterase and carbonic anhydrase inhibitory actions, Bioorg. Chem. 74 (2017) 104-114.
- [41] A. Akıncıoğlu, E. Kocaman, H. Akıncıoğlu, R.E. Salmas, S. Durdağı, İ. Gülçin, C.T. Supuran, S. Göksu, The synthesis of novel sulfamides derived from β-benzylphenethylamines as acetylcholinesterase, butyrylcholinesterase and carbonic anhydrase enzymes inhibitors, Bioorg. Chem. 74 (2017) 238-250.
- [42] P. Taslimi, S. Osmanova, İ. Gulçin, S. Sardarova, V. Farzaliyev, A. Sujayev, R. Kaya, F. Koc, S. Beydemir, S.H. Alwasel, O.I. Kufrevioglu, Discovery of potent carbonic anhydrase, acetylcholinesterase and butyrylcholinesterase enzymes inhibitors: the new amides and thiazolidineones synthesized on an acetophenone base, J. Biochem. Mol. Toxicol. 31 (2017) e21931.
- [43] N. Nohut Maslakci, A. Biçer, G. Turgut Cin, A. Uygun Oksüz, Electrochromic properties of some bis-chalcone derivatives-based nanofibers, J. Appl. Polym. Sci. 135 (2018) 46010.
- [44] G. Yakalı, A. Biçer, C. Eke, G. Turgut Cin, Solid state structural investigations of the bis(chalcone) compound with single crystal X-ray crystallography, DFT, gamma-ray spectroscopy and chemical spectroscopy methods, Radiat. Phys. Chem. 145 (2018) 89-96.
- [45] N.K. Konduru, S. Dey, M. Sajid, M. Owais, N. Ahmed, Synthesis and antibacterial and antifungal evaluation of some chalcone based sulfones and bisulfones, Eur. J. Med. Chem. 59 (2013) 23-30.
- [46] C. Guha, N. Sepay, T. Halder, A.K. Mallik, Remarkable Diastereoselectivity of the Thia-Michael Reaction on α,α'-di[(E)-benzylidene]alkanones: Exclusive formation of a *meso* product, Synlett 29 (2018) 1161-1166.
- [47] S.B. Öztürk Sarıkaya, F. Topal, M. Şentürk, İ. Gülçin, C.T. Supuran, In vitro inhibition of αcarbonic anhydrase isozymes by some phenolic compounds, Bioorg. Med. Chem. Lett. 21 (2011) 4259-4262.

- [48] M. Şentürk, İ. Gülçin, Ş. Beydemir, Ö.İ. Küfrevioğlu, C.T. Supuran, In vitro inhibition of human carbonic anhydrase I and II isozymes with natural phenolic compounds, Chem. Biol. Drug Des. 77 (2011) 494-499.
- [49] A. Innocenti, İ. Gülçin, A. Scozzafava, C.T. Supuran, Carbonic anhydrase inhibitors. Antioxidant polyphenol natural products effectively inhibit mammalian isoforms I-XV, Bioorg. Med. Chem. Lett. 20 (2010) 5050-5053.
- [50] A. Innocenti, S.B. Öztürk Sarıkaya, İ. Gülçin, C.T. Supuran, Carbonic anhydrase inhibitors. Inhibition of mammalian isoforms I-XIV with a series of natural product polyphenols and phenolic acids, Bioorg. Med. Chem. 18 (2010) 2159-2164.
- [51] P. Taslimi, H.E. Aslan, Y. Demir, N. Öztaşkın, A. Maraş, İ. Gulçin, Ş. Beydemir, S. Göksu, Diarilmethanon, bromophenols and diarilmetan compounds: discovery of potent aldose reductase, α-amylase and α-glycosidase inhibitors as new therapeutic approach in diabetes and functional hyperglycemia, Int. J. Biol. Macromol. 119 (2018) 857-863.
- [52] T.A. Coban, S. Beydemir, İ. Gülcin, D. Ekinci, A. Innocenti, D. Vullo, C.T. Supuran, Sildenafil is a strong activator of mammalian carbonic anhydrase isoforms I-XIV, Bioorg. Med. Chem. 17 (2009) 5791-5795.
- [53] M. Şentürk, İ. Gülçin, A. Daştan, Ö.İ. Küfrevioğlu, C.T. Supuran, Carbonic anhydrase inhibitors. Inhibition of human erythrocyte isozymes I and II with a series of antioxidant phenols, Bioorg. Med. Chem. 17 (2009) 3207-3211.
- [54] A. Akıncıoğlu, M. Topal, İ. Gülçin, S. Göksu, Novel sulfamides and sulfonamides incorporating tetralin scaffold as carbonic anhydrase and acetylcholine esterase inhibitors, Arch. Pharm. 347 (2014) 68-76.
- [55] H.İ. Gül, K. Kucukoglu, C. Yamali, S. Bilginer, H. Yuca, İ. Ozturk, P. Taslimi, İ. Gülçin, C.T. Supuran, Synthesis of 4-(2-substitutedhydrazinyl)benzenesulfonamides and their carbonic anhydrase inhibitory effects, J. Enzyme Inhib. Med. Chem. 31 (2016) 568-573.
- [56] H.İ. Gül, M. Tuğrak, H. Sakagami, P. Taslimi, İ. Gülçin, C.T. Supuran, Synthesis and bioactivity studies on new 4-(3-(4-substitutedphenyl)-3a,4-dihydro-3h-indeno[1,2-c]pyrazol-2-yl) benzenesulfonamides, J. Enzyme Inhib. Med. Chem. 31 (2016) 1619-1624.
- [57] F. Özbey, P. Taslimi, İ. Gulcin, A. Maraş, S. Goksu, C.T. Supuran, Synthesis, acetylcholinesterase, butyrilcholinesterase, carbonic anhydrase inhibitory and metal chelating properties of some novel diaryl ether, J. Enzyme Inhib. Med. Chem. 31 (2016) 79-85.
- [58] N. Öztaşkın, Y. Çetinkaya, P. Taslimi, S. Göksu, İ. Gülçin, Antioxidant and acetylcholinesterase inhibition properties of novel bromophenol derivatives, Bioorg. Chem. 60 (2015) 49-57.
- [59] İ. Gulçin, M. Abbasova, P. Taslimi, Z. Huyut, L. Safarova, A. Sujayev, V. Farzaliyev, S. Beydemir, S.H. Alwasel, C.T. Supuran, Synthesis and biological evaluation of aminomethyl and alkoxymethyl derivatives as carbonic anhydrase, acetylcholinesterase and butyrylcholinesterase inhibitors, J. Enzyme Inhib. Med. Chem. 32 (2017) 1174-1182.

- [60] C. Yamali, H.İ. Gül, A. Ece, P. Taslimi, İ. Gulçin, Synthesis, molecular modeling and biological evaluation of 4-[5-aryl-3-(thiophen-2-yl)-4,5-dihydro-1H-pyrazol-1-yl]benzenesulfonamides towards acetylcholinesterase, carbonic anhydrase I and II enzymes, Chem. Biol. Drug Des. 91 (2018) 854-866.
- [61] M. Rezai, Ç. Bayrak, P. Taslimi, İ. Gulçin, A. Menzek, The first synthesis, antioxidant and anticholinergic activities of 1-(4,5-dihydroxybenzyl)pyrrolidin-2-one derivative bromophenols including natural products, Turk. J. Chem. 42 (2018) 808-825.
- [62] S. Naveen, B. Dinesh, K. Abiraj, D. Channe Gowda, M. A. Sridhar, J. Shashidhara Prasad, Synthesis and crystal structure of tert-butyl 2-((phenylthio)-carbonyl) pyrrolidine-1-carboxylate, J. Chem Crystallogr. 37 (2007) 721-725.
- [63] A.N. Chekhlov, Crystal structure of 1-(2-pyridiniomethyl)-2,4-bis(phenylsulfonyl)benzene chloride ethanol solvate, Russ. J. Gen. Chem. 74 (2004) 944-948.
- [64] R.N. Osorio-Yanez, C. Crisostomo-Lucas, E. Santacruz-Juarez, R. Reyes-Martineza, D. Morales-Morales, Acta Cryst. Sect. E 70 (2014) 0529.
- [65] J. Bernstein, R.E Davis, L. Shimoni, N.L. Chang, Hydrogen-bond Pattern Functionality and Graph Sets, Angew. Chem. Int. Ed. Engl. 34 (1995) 1555-1573.
- [66] J.F. Darby, M. Atobe, J.D. Firth, P. Bond, G.J. Davies, P. O'Brien, R.E. Hubbard, Increase of enzyme activity through specific covalent modification with fragments, Chem. Sci. 8 (2017) 7772-7779.
- [67] X.Y. Meng, H. X. Zhang, M. Mezei, M. Cui, Molecular docking: a powerful approach for structure-based drug discovery, Curr. Comput. Aided Drug Des. 7 (2011) 146-157.
- [68] Agilent Technologies, CrysAlis PRO and CrysAlis RED, Yarnton, Oxfordshire England, 2002.
- [69] G.M. Sheldrick, SHELXT-Integrated space-group and crystal-structure determination, Acta Crystallogr. Sect. A 71 (2015) 3-8.
- [70] O.V. Dolomanov, L.J. Bourhis, R.J. Gildea, J.A.K. Howard and H. Puschmann, OLEX2: A complete structure solution, refinement and analysis program, J. Appl. Cryst. 42 (2009) 339-341.
- [71] G.M. Sheldrick, Crystal structure refinement with SHELXL, Acta Crystallogr. Sect. C 71 (2015)3-8.
- [72] J.A. Verpoorte, S. Mehta, J.T. Edsall, Esterase activities of human carbonic anhydrases B and C, J. Biol. Chem. 242 (1967) 4221-4229.
- [73] P. Taslimi, İ. Gulçin, Antioxidant and anticholinergic properties of olivetol, J. Food Biochem. 42 (2018) e12516.
- [74] F. Türker, D. Barut Celepci, A. Aktaş, P. Taslimi, Y. Gök, M. Aygün, İ. Gulçin, Metacyanobenzyl substituted benzimidazole: Synthesis, characterization, crystal structure and carbonic anhydrase, α-glycosidase, butyrylcholinesterase, acetylcholinesterase inhibitory properties, Arch. Pharm. 351 (2018) e201800029.

- [75] İ. Gulçin, P. Taslimi, Sulfonamide inhibitors: A patent review 2013-present, Exp. Opin. Ther. Pat. 28 (2018) 541-549.
- [76] B. Yiğit, M. Yiğit, D. Barut Celepci, Y. Gök, A. Aktaş, M. Aygün, P. Taslimi, İ. Gulçin, Novel benzylic substituted imidazolinium, tetrahydropyrimidinium and tetrahydrodiazepinium saltspotent carbonic anhydrase and acetylcholinesterase inhibitors, ChemistrySelect 3 (2018) 7976-7982.
- [77] A. Behcet, T. Çağlılar, D. Barut Celepci, A. Aktaş, P. Taslimi, Y. Gök, M. Aygün, R. Kaya, İ. Gulçin, Synthesis, characterization and crystal structure of 2-(4-hydroxyphenyl)ethyl and 2-(4-nitrophenyl)ethyl substituted benzimidazolium bromide salts: their inhibitory properties against carbonic anhydrase and acetylcholinesterase, J. Mol. Struct. 1170 (2018) 160-169.
- [78] F. Turkan, Z. Huyut, P. Taslimi, İ. Gulçin, The in vivo effects of cefazolin, cefuroxime, and cefoperazon on the carbonic anhydrase in different rat tissues, J. Biochem. Mol. Toxicol. 32 (2018) e22041.
- [79] G.L. Ellman, K.D. Courtney, V. Andres, R.M. Featherston, A new and rapid colorimetric determination of acetylcholinesterase activity, Biochem. Pharmacol, 7 (1961) 88-95.
- [80] P. Taslimi, C. Çağlayan, F. Farzaliyev, O. Nabiyev, A. Sujayev, F. Türkan, R. Kaya, İ. Gulçin, Synthesis and discovery of potent carbonic anhydrase, acetylcholinesterase, butyrylcholinesterase and α-glycosidase enzymes inhibitors: the novel N,N'-bis-cyanomethylamine and alkoxymethylamine derivatives, J. Biochem. Mol. Toxicol. 32 (2018) e22042.
- [81] M.U. Koçyigit, P. Taslimi, F. Gurses, S. Soylu, S. Durna Dastan, İ. Gulçin, The effects of wireless electromagnetic fields on the activities of carbonic anhydrase and acetylcholinesterase enzymes in various tissues of rats, J. Biochem. Mol. Toxicol. 32 (2018) e22031.
- [82] P. Taslimi, E. Sujayev, F. Turkan, E. Garibov, Z. Huyut, F. Farzaliyev, S. Mamedova, İ. Gulçin, Synthesis and investigation of the conversion reactions of pyrimidine-thiones with nucleophilic reagent and evaluation of their acetylcholinesterase, carbonic anhydrase inhibition and antioxidant activities, J. Biochem. Mol. Toxicol. 32 (2018) e22019.
- [83] S. Burmaoglu, A.O. Yilmaz, P. Taslimi, O. Algul, D. Kılıç, İ., Gulçin, Synthesis and biological evaluation of phloroglucinol derivatives possessing α -glycosidase, acetylcholinesterase, butyrylcholinesterase, carbonic anhydrase inhibitory activity, Arch. Pharm. 351 (2018) e1700314.
- [84] Y. Sarı, A. Aktaş, P. Taslimi, Y. Gök, C. Caglayan, İ. Gulçin, Novel N-propylphthalimide and 4vinylbenzyl substituted benzimidazole salts: synthesis, characterization and determination of their metal chelating effects and inhibition profiles against acetylcholinesterase, and carbonic anhydrase enzymes, J. Biochem. Mol. Toxicol. 32 (2018) e22009.
- [85] F. Erdemir, D. Barut Celepci, A. Aktaş, P. Taslimi, Y. Gök, H. Karabıyık, İ. Gulçin, 2-Hydroxyethyl substituted NHC precursors: Synthesis, characterization, crystal structure and carbonic anhydrase, α-glycosidase, butyrylcholinesterase, and acetylcholinesterase inhibitory properties, J. Mol. Struct. 1155 (2018) 797-806.

[86] P. Taslimi, H. Akıncıoğlu, İ. Gulçin, Synephrine and phenylephrine act as α-amylase, αglycosidase, acetylcholinesterase, butyrylcholinesterase and carbonic anhydrase enzymes inhibitors, J. Biochem. Mol. Toxicol. 3 (2017) e21973. Acctiontic

	Entry	Catalyzer	Temperature	Solvent	Time	Yield
	1	I ₂ (%10 mol)	Room temperature	DCM	1 day	Isomer mixture
	2	$I_2(\%10 \text{ mol})$	Reflux	DCM	1 day	Isomer mixture
	3	FeCl ₃ (%2 mol)	Room temperature	DCM	1 day	No reaction
	4	AlCl ₃ (%2 mol)	Room temperature	DCM	1 day	No reaction
	5	$Al_2O_3(5 mol)$	Reflux	EtOH	1 day	No reaction
L	6	t-BuOK (2 mol)	Room temperature	BuOH	1 day	No reaction
	7	NEt ₃ (5 mol)	Reflux	DCM	1 day	No reaction
	8	Na (1 mol)	Room temperature	DCM	1 day	Product observed

Table 1. Optimization of reaction conditions for addition of thiophenol to bis-chalcones

Table 2. Inhibition patrameters including half maximal inhibitory concentration (IC_{50}) and inhibition constants (Ki) of a dousen of novel bissulfide compounds (**3a-j**) against human carbonic anhydrase I, and II isoenzymes (hCA I, and II), acetylcholinesterase (AChE), and butyrylcholinesterase (BChE) enzymes

Compounds				IC ₅₀	(nM)						K _i (nM)	, 	
Compounds	hCA I	r ²	hCA II	r ²	AChE	r ²	BChE	r ²	hCA I	hCA II	AChE	BChE	AChE/BChE
3 a	109.43	0.9736	138.55	0.9308	12.73	0.9563	28.93	0.9892	87.53±14.07	127.44±31.84	8.63±1.83	19.73±3.64	0.43
3b	147.05	0.9927	170.82	0.9673	10.03	0.9604	31.53	0.9722	183.23±28.18	157.99±50.72	7.14±1.05	22.84±4.93	0.31
3c	180.34	0.9598	201.58	0.9902	19.39	0.9912	42.04	0.9831	167.34±22.72	194.02±28.55	12.74±2.95	20.51±3.02	0.62
3d	117.03	0.9819	143.84	0.9819	41.77	0.9732	61.38	0.9682	137.03±43.94	154.93±36.04	32.64±6.86	37.38±11.53	0.87
3 e	71.66	0.9816	80.03	0.9772	7.18	0.9208	17.90	0.9801	80.48±17.11	88.32±18.13	4.55±0.94	12.77±2.20	0.35
3f	55.08	0.9743	74.22	0.9609	15.71	0.9714	24.83	0.9492	50.66±9.17	80.12±21.98	11.42±1.98	18.53±5.94	0.61
3g	61.38	0.9891	68.52	0.9923	15.02	0.9843	20.28	0.9618	64.83±19.33	61.34±13.81	13.56±2.64	14.82±4.08	0.91
3h	38.63	0.9812	42.84	0.9904	23.83	0.9672	28.11	0.9910	47.92±7.20	48.18±19.10	17.81±4.66	19.73±3.98	0.90
3i	44.28	0.9590	64.03	0.9891	9.83	0.9840	29.43	0.9733	39.14±9.94	46.03±10.47	6.06±2.01	17.43±2.25	0.34
3ј	69.43	0.9711	72.71	0.9690	17.47	0.9531	31.47	0.9562	54.37±11.10	60.52±18.52	14.37±3.33	20.28±4.90	0.70
AZA*	265.41	0.9954	248.54	0.9714	-	-	-	-	273.61±76.86	229.08±55.14	-	-	-
\mathbf{TAC}^{Ψ}	-	-	-	-	92.20	0.9796	105.16	0.9711	-	-	56.37±15.10	63.40±13.62	0.88

*Acetazolamide (AZA) was used as a standard inhibitor for both carbonic anhydrases I, and II (hCA I and II) isoenzymes

⁴Tacrine (TAC) was used as a standard inhibitor for both acetylcholinesterase (AChE), and butyrylcholinesterase (BChE) enzymes

		Bond Len	igths (A)		
3h		3i	8	3j	
O1-C4	1.214(7)	O1-C19	1.206(8)	Br1-C9	1.899(5)
Cl1-C9	1.735(5)	S2-C20	1.861(8)	01-C1	1.221(8)
S1-C5	1.852(4)	S2-C27	1.782(9)	S1-C5	1.857(5)
S1-C12	1.778(5)	Br2-C23	1.864(8)	S1-C12	1.771(5)
		Br1-C12	1.912(9)		
		S1-C1	1.770(9)		
		S1-C7	1.834(8)		
		Bond An	igles (°)		
<u>3h</u>		<u> </u>		3j	
C5-S1-C12	101.4(2)	C1-S1-C7	101.1(4)	C12-S1-C5	101.6(2)
01-C4-C3	123.6(2)	C20-S2-C27	102.3(4)	S1-C5-C2	106.1(3)
CI1-C9-C8	120.0(5)	01-C19-C18	123.2(8)	01-C1-C2	122.7(3)
		Br1-C12-C13	117.3(7)	Br1-C9-C8	120.4(4)
		DIZ-023-024	119.8(0)		

Table 3. Selected experimental parameters of the novel bis-sulfide compounds (3h, 3i and 3j)

Crystal data	3h	3i	3j
Empirical formula	C ₃₂ H ₂₈ Cl ₂ OS ₂	$C_{32}H_{28}Br_2OS_2$	$C_{32}H_{28}Br_2OS_2$
Formula weight	563.56	652.48	652.48
Temperature (K)	293(2)	292(2)	294(2)
Crystal system	monoclinic	triclinic	monoclinic
Space group	lm	P-1	lm
Unit cell dimensions			
<i>a</i> (Å)			
<i>b</i> (Å)	5.1312(5)	10.399(2)	5.0984(4)
<i>c</i> (Å)	25.692(2)	10.431(3)	25.8212(19)
a (9	10.9554(11)	14.373(4)	11.2384(6)
	90	108.01(2)	90
β(9	97.941(9)	101.662(19)	98.313(6)
y (9	90	97.05(2)	90
Volume (Å ³)	1430.4(2)	1423.2(6)	1463.94(18)
Z	2	2	2
D_{calc} (g/cm ⁻³)	1.308	1.523	1.480
Absorption coefficient (mm ⁻¹)	0.397	3.019	2.935
F (000)	588	660.0	660.0
Crystal size (mm)	$0.659 \times 0.395 \times 0.053$	0.335 × 0.242 × 0.166	0.241 × 0.211 × 0.092
······································	$-6 \le h \le 3$,	$-12 \le h \le 12$,	$-6 \le h \le 6$,
Index ranges	$-31 \le k \le 27,$	$-12 \le k \le 12,$	$-31 \le k \le 31$,
	$-13 \le l \le 13$	$-17 \leq l \leq 17$	$-8 \le l \le 13$
Reflections collected/unique	2733/1791	7467/7467	4554/2187
Data / restrains / parameters	1791/2/172	7467/12/335	2187/2/172
Goodness of fit on F ²	1.029	0.848	1.023
Einal D indians $[I > 2-(I)]$	$R_1 = 0.0465,$	$R_1 = 0.0543,$	$R_1 = 0.0340,$
Final K indices $[I > 26(I)]$	$wR_2 = 0.1151$	$wR_2 = 0.1462$	$wR_2 = 0.0603$
<i>P</i> indices (all data)	$R_1 = 0.0523$	$R_1 = 0.1367$	$R_1 = 0.0616$
A mulces (an uata)	$wR_2 = 0.1226$	$wR_2 = 0.1704$	$wR_2 = 0.0685$
Largest difference peak and hole (e Å ⁻³)	0.23/-0.19	0.39/-0.39	0.24/-0.28
	24		

Table 4. Crystal data and structure refinement parameters for the novel bis-sulfide compounds (3h, 3i and 3j)

Compounds	D-H···A	D–H	Н…А	D···A	D-H···A
3h	C2-H2BS1	0.97	2.74	3.1821(5)	108
	C15-H15BS1	0.970(11)	2.719(8)	3.178(8)	109.5(7)
2;	C17-H17AO1i	0.969(12)	2.592(10)	3.544(10)	167.4(8)
51	C17-H17BS2	0.971(12)	2.683(8)	3.103(8)	106.6(7)
	C25-H25Cg2	0.932(10)	2.803(9)	3.374(9)	120.7(8)
3ј	C3-H3BS1	0.970(6)	2.757(5)	3.192(4)	108.0(4)

Table 5. Interactions geometry (Å,°) for the novel bis-sulfide compounds (3h, 3i and 3j)

FIGURE LEGENDS

- Figure 1. Determination of inhibition constant values (Ki) of novel bis-sulfide compounds (3a-j) obtained from Lineweaver-Burk graphs against human carbonic anhydrases isoenzyme I and II (hCA I and II) (3i), and acetylcholinesterase (AChE) and butrylcholinesterase (BChE) enzymes (3e)
- Figure 2. Inhibition constant values (Ki) of novel bis-sulfide compounds (3a-j) for human carbonic anhydrases isoenzyme I, and II (hCA I, and II) (A) and acetylcholinesterase (AChE) and butrylcholinesterase (BChE) enzymes (B)
- Figure 3. Thermal ellipsoid view of the molecules with the atom numbering scheme and 75% probability
- Figure 4. Packing structure of the novel bis-sulfide compound of 3h via stacking interactions along the *a* axis, (3i) via strong C-H...O interctions along the *b* axis, (3j) via stacking interactions along the b axis



















Scheme 1. Biological properties of sulfide and bis-sulfide compounds

MAN



a Ar =C₆H₅-, **b** Ar=2-Naphthyl-, **c** Ar=3-NO₂C₆H₄-, **d** Ar= 4-NO₂C₆H₄-, **e** Ar=3-FC₆H₄-, **f** Ar=4-FC₆H₄-, **g** Ar=3-ClC₆H₄-, **h** Ar=4-ClC₆H₄-, **i** Ar= 3-BrC₆H₄-, **j** Ar=4-BrC₆H₄-

Scheme 2. Synthesis of bis-chalcone derivatives (2a-j)



a Ar =C₆H₅-, **b** Ar=2-Naphthyl-, **c** Ar=3-NO₂C₆H₄-, **d** Ar=4-NO₂C₆H₄-, **e** Ar=3-FC₆H₄**f** Ar=4-FC₆H₄-, **g** Ar=3-ClC₆H₄-, **h** Ar=4-ClC₆H₄-, **i** Ar=3-BrC₆H₄-, **j** Ar=4-BrC₆H₄-

Scheme 3. Synthesis of the novel bis-sulfide derivatives (3a-j)

Highlight

- A novel serie of bis-thiomethylcyclohexanone derivatives was synthesised.
- These compounds have been characterized by ¹H-NMR, ¹³C-NMR, FTIR and elemental analysis.
- The molecular structure and crystal packing of the compounds **3h**, **3i** and **3j** were obtained.
- Their inhibition effects on hCA I, hCA II, AChE and BChE enzymes were determined.

Synthesis, Characterization, Crystal Structure of Novel Bis-Thiomethylcyclohexanone Derivatives and Their Inhibitory Properties Against Some Metabolic Enzymes



In this study, synthesis, characterization, crystal structure and biological activivtes of a dousen of novel bis-thiomethylcyclohexanone derivatives were achieved. Also, for evaluation of novel copounds inhibition effects were inwestigated against the carbonic anhydrase I, and II isoenzymes, acetylcholinesterase and butyrylcholinesterase enzymes.