### Synthesis of Novel 1, 3 -Oxazepine Compounds from New Azo Schiff bases Containing Thiadiazole Moiety

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#### Abstract

In this work new Azo imines [3-4] and 1,3-oxazepine derivatives [5-10] were prepared starting from azoaldehyde derivative 2-Hydroxy-5-(5-thiole-[1,3,4] -thiadiazol-2-ylazo)-benzaldehyde [2]. Azoaldehyde derivative [2] was prepared via coupling reaction between phenoxide anion of *o*-Hydroxy benzaldehyde [1] and 5-thiole-1,3,4-thiadiazol-2-yl diazonium chloride.The new imines [3] and [4] were prepared by reaction of azoaldehyde derivative [2] with each primary amines 2-Amine-1,3,4-thiadiazole -5- thiol and 5-(Benzylthio)-1,3,4-thiadiazole-2- amine in presence of glacial acetic acid as catalyst in absolute ethanol , respectively. The new 1,3-oxazepine derivatives [5-10] were obtained from treatment of each new Schiff bases derivatives [4] and [5] with each maleic anhydride , phthalic anhydride and 3-Nitrophthalic anhydride ,respectively , in dry benzene.These new prepared compounds might have some biological activity.

All new derivatives were characterized by (C.H.N.) elementary analysis and FT-IR spectra.

الخلاصة: -

تم في هذا البحث تحضير مشتقات ايمينات [3-4] و 3،1- اوكسازيبين [5-10] جديدة من مشتق الازوالديهايد 2-هيدروكسي -5-(5-مركبتو-[4,3,1]- ثايادايازول-2-يل ازو) - بنز الديهايد [2] . حضر مشتق الازوالديهايد [2] من خلال تفاعل الازدواج ما بين ايون الفينوكسايد لمركب *o* - هيدروكسي بنز الديهايد و 5 - مركبتو-4,3,1 - ثايا دايازول - 2- يل دايازونيوم كلورايد . حضرت قواعد شيف الجديدة [3] و[4] عن طريق تفاعل مشتق الازوالديهايد [2] مع كل من 2-امينو -4,3,1 - ثايادايازول - 3 - هيدروكسي بنز الديهايد و 5 - مركبتو-4,3,1 - ثايا دايازول - 2- يل امينو -4,3,1 - ثايادايازول -5- ثايول و5-(بنز ايل ثايو) - 4,3,1 - ثايادايازول - 2 - امين .على التوالي ، بوجود حامض الخليك الثلجي كعامل مساعد في الايثانول المطلق تم الحصول على مشتقات 3،1 – اوكسازيبين الجديدة [5] من معاملة كل من مشتقات قواعد شيف المحضرة [3] و [4] مع كل من انهدريد المالييك و انهدريد الفثاليك و 3-نترو انهدريد الفثاليك ،علي التوالي ،في البنزين الجاف أن هذة المشتقات الجديدة قد تمتلك فعالية حيوية . شخصت جميع المشتقات الجديدة بوساطة التوالي اليوالي الجاف أن هذة المشتقات الجديدة قد تمتلك فعالية حيوية . شخصت جميع المشتقات الجديدة بوساطة التوالي ، التوالي ، التوالي ، علي التوالي .

#### Introduction

For a long time, the synthesis of 1,3- and 1,4-oxazepine rings was based on two limited classical types of reactions, the first reaction is called Valence-bond isomerization which is carried out via irradiation of polyarylpyridine N-oxides. This irradiation results in ring expansion to 1,3-oxazepine in high yield and some deoxygenation to the parent  $\operatorname{amines}^{(1)}$ . The second reaction is called Enamines condensation which is carried out by reaction of Erythro 1,2-diphenyl-2-phenylaminoethanol with dimethylacetylene dicarboxylate in methanol at room temperature to give a mixture of the Michael adduct and tetrahydro-1,4-oxazepin-7-one<sup>(2)</sup>. Recently, cycloaddition reaction, which is a type from a pericyclic reactions is used to synthesis of 1,3-oxazepine ring<sup>(3-6)</sup>. This type of reactions is not limited and gives various 1,3-oxazepine ring derivatives. The type of cycloaddition reaction that used to synthesis of 1,3-oxazepine ring was classified as  $(2+5) \rightarrow 7$  cycloaddition reaction in which two atoms of imine group as two-membered component was added to five-membered component such as maleic or phthalic anhydrides to give a seven-membered heterocycle<sup>(7-9)</sup>.

1,3-Oxazepine is unsaturated seven-membered hetrocycle containing oxygen atom in position (1), nitrogen atom in position (3) in edition of five carbons. Oxazepine derivatives showed various biological activities such as antibacterial<sup>(10)</sup> and inhibitors for some enzymes action<sup>(11)</sup>. Some of oxazepine derivatives are used in another applied fields<sup>(12)</sup>.

Schiff bases or imines are prepared via acid-catalysed condensation reaction of aromatic aldehydes or ketones with primary amines <sup>(13,14)</sup>. Azo Schiff bases are prepared by reaction of azoaldehydes with primary amines <sup>(15)</sup>. Mechanism of Schiff base formation was well known <sup>(13,14)</sup>. Various azo Schiff bases derivatives were prepared and some of them showed biological activity such as anticancer <sup>(16)</sup> antiviral <sup>(17)</sup>, antifungal <sup>(18)</sup>, antibacterial <sup>(19)</sup> and anticonvulsant <sup>(20)</sup>. Thiadiazoles have a variety of potential biological activities <sup>(21,22)</sup>, therefore a large number of thiadiazole derivatives have been prepared <sup>(23,24)</sup>. Many synthesis of 1,3,4-thiadiazoles proceed from thiosemicarbazide or substituted thiosemicarbazide <sup>(25,26)</sup>.

### **Experimental:**

#### <u>General</u>

- 1) The solvents and liquid reagents were purified when it was necessary; the solid materials were also dried under reduced pressure when it was necessary.
- 2) TLC were performed on pre-coated sheets with 0.25 mm layer of Silica Gel GF254 of the Merck company, the detection was followed by coloring with iodine or  $H_2SO_4$  in ethanol (60%) followed by heating.
- 3) Evaporating of solvents by using Buchi vacuum rotary evaporator type 160.
- 4) Melting points (M.P.) were determined by Stuart melting point apparatus.
- 5) Elemental analysis measured on E.A.300, Euro- Vector, Italy, 2003.
- 6) FT-IR spectra were recorded on FT-IR 8400s, Schimadzu-Spectrophotometer and using KBr discs.

### **Preparation Methods:**

### Synthesis of 2-Hydroxy-5-(5-mercapto-[1,3,4] -thiadiazol-2-ylazo)-benzaldehyde [2] (27)

2-Amino-1,3,4-thiadiazole-5-thiol (2g ,0.015mole) was dissolved in a solution of conc. hydrochloric acid (16mL) and distilled water (10mL). The mixture was cold at (0°C) in an ice bath . a solution of sodium nitrite (1.3g ,0.0188 mole) in (10mL) of distilled water was added dropwise with stirring to the mixture, the temperature of the ice bath was controlled between (0-5°C). A solution of (1.834g ,0.015 mole) of *O*-Hydroxybenzaldehyde [1] in (15mL) of (10%) sodium hydroxide solution was prepared and cold to (5°C) by immersion in an ice bath. The salicyldehyde solution was stirred vigorously ,the cold diazonium salt solution was added very slowly to the salicyldehde solution , a red colour developed and red crystals soon separated .when all the diazonium salt solution was added , the mixture was allowed to stand in an ice bath for 30 min.with occasional stirring. The solution was filtered , washed well with distilled water then with alittle alcohol , recrystallized from ethanol and dried upon filter paper,yield 66%,M.P.178-180°C.

# Synthesis of 4-((Z)-(5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)-2-((Z)-(5-mercapto-1,3,4-thiadiazol-2-ylimino)methyl) phenol [3]

Azobenzaldehyde derivative [2](0.532g, 0.002 mole) was dissolved in absolute ethanol (15 mL) containing a drop of glacial acetic acid ,then 2-Amino-1,3,4-thiadiazole-5-thiol (0.266 g ,0.002 mole) was dissolved in absolute ethanol (15 mL) and added dropwise. The reaction mixture was refluxed with stirring on a water bath at (70 °C) for 2hrs.T.L.C. (ethanol:pet.ether)(1:1) , $R_f = 0.63$ .Then the mixture was allowed to cool down to room temperature , the coloured precipitate was filtered and recrystallized from ethanol ,yield 68% , M.P.220-222°C.

# Synthesis of 2-((Z)-(5-(benzylthio)-1,3,4-thiadiazol-2-ylimino)methyl)-4-((Z)-(5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)phenol [4]

Azobenzaldehyde derivative [2](0.532g, 0.002 mole) was dissolved in absolute ethanol (15 mL) containing a drop of glacial acetic acid ,then 5-(butylthio)-1,3,4-thiadiazol-2-amine (0.378 g ,0.002 mole) was dissolved in absolute ethanol (15 mL) and added dropwise. The reaction mixture was refluxed with stirring on a water bath at ( $70 \ ^{\circ}C$ ) for 2hrs.T.L.C. (ethanol:pet.ether)(1:1) , $R_f = 0.59$ .Then the mixture was allowed to cool down to room temperature , the coloured precipitate was filtered and recrystallized from ethanol ,yield 70% , M.P.160-162°C.

# Synthesis of (Z)-2-(2-hydroxy-5-((Z)-(5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)phenyl)-3-(5-mercapto-1,3,4-thiadiazol-2-yl)-2,3-dihydro-1,3-oxazepine-4,7-dione [5]

A mixture of imine derivative [3]( 0.381 g, 0.001 mole) and maleic anhydride (0.098 g ,0.001 mole) in dry benzene ( 20 mL) was refluxed on a water bath at (75°C) for 4hrs. T.L.C. (benzene:methanol ) (3:1)  $R_f = 0.70$ ., then the mixture was allowed to cool down to room temperature. The resulting solid crystals were filtered and recrystallized from dioxan, yield 65%, M.P.180-182 °C.

# Synthesis of (Z)-3-(2-hydroxy-5-((5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)phenyl)-4-(5-mercapto-1,3,4-thiadiazol-2-yl)-3,4-dihydrobenzo[e][1,3]oxazepine-1,5-dione [6]

A mixture of imine derivative [3]( 0.381g, 0.001 mole) and phthalic anhydride (0.148g ,0.001mole) in dry benzene (20 mL) was refluxed on a water bath at (75°C) for 4hrs. T.L.C. (benzene:methanol) (3:1)  $R_f = 0.74$ ., then the mixture was allowed to cool down to room temperature. The resulting solid crystals were filtered and recrystallized from dioxan, yield 71%, M.P.174-176 °C.

# Synthesis of (Z)-3-(2-hydroxy-5-((5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)phenyl)-4-(5-mercapto-1,3,4-thiadiazol-2-yl)-6-nitro-3,4-dihydrobenzo[e][1,3]oxazepine-1,5-dione [7]

A mixture of imine derivative [3]( 0.381 g, 0.001 mole) and 3-Nitro phthalic anhydride (0.193 g ,0.001mole) in dry benzene ( 20 mL), was refluxed on a water bath at (75°C) for 4hrs. T.L.C. (benzene:methanol ) (3:1)  $R_f = 0.79$ ., then the mixture was allowed to cool down to room temperature. The resulting solid crystals were filtered and recrystallized from dioxan, yield 66%, M.P.200-202 °C.

# Synthesis of (Z)-4-(5-(Benzylthio)-1,3,4-thiadiazol-2-yl)-3-(2-hydroxy-5-((5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)phenyl)-3,4-dihydrobenzo[e][1,3]oxazepine-1,5-dione[8]

A mixture of imine derivative [4]( 0.437 g, 0.001 mole) and maleic anhydride (0.098 g ,0.001mole) in dry benzene ( 20 mL) was refluxed on a water bath at (75°C) for 4hrs. T.L.C. (benzene:methanol) (3:1)  $R_f = 0.63$  then the mixture was allowed to cool down to room temperature . The resulting solid crystals were filtered and recrystallized from dioxan, yield 70%, M.P.280-282 °C.

#### Synthesis of 8-(5-Benzylsulfanyl-[1,3,4]thiadiazol-2-yl)-7-[2-hydroxy-5-(5-mercapto-[1,3,4]thiadiazol-2-ylazo)-phenyl]-7,8-dihydro-6-oxa-8-aza-benzocycloheptene-5,9-dione[9]

A mixture of imine derivative [4]( 0.437 g, 0.001 mole) and phthalic anhydride (0.148 g ,0.001mole) in dry benzene (20 mL) was refluxed on a water bath at (75°C) for 4hrs. T.L.C. (benzene:methanol) (3:1)  $R_f = 0.67$ , then the mixture was allowed to cool down to room temperature. The resulting solid crystals were filtered and recrystallized from dioxan, yield 69%, M.P.190-192 °C.

# Synthesis of (Z)-4-(5-(Benzylthio)-1,3,4-thiadiazol-2-yl)-3-(2-hydroxy-5-((5-mercapto-1,3,4-thiadiazol-2-yl)diazenyl)phenyl)-6-nitro-3,4-dihydrobenzo[e][1,3]oxazepine-1,5-dione [10]

A mixture of imine derivative [4]( 0.437 g, 0.001 mole) and 3-Nitrophthalic anhydride (0.193 g, 0.001 mole) in dry benzene (20 mL) was refluxed on awater bath at (75°C) for 4hrs. T.L.C. (benzene:methanol) (3:1)  $R_f = 0.77$ , then the mixture was allowed to cool down to room

temperature . The resulting solid crystals were filtered and recrystallized from dioxan, yield 70%, M.P.153-155 °C.

Com p. No.	M.P °C	Yield %	M.F	M. Wt g/m ole	C.H.N analysis							
					Calc	ulate	d%	Found%				
					С	Η	Ν	С	Η	Ν		
[2]	178- 180	66	C9H6N4O2 S2	266	-	-	-	-	-	-		
[3]	220- 222	68	C <sub>11</sub> H <sub>7</sub> N <sub>7</sub> O S <sub>4</sub>	381	34.64	1.8 3	25.7 2	35.0 2	2.11	26.0 1		
[4]	160- 162	70	C <sub>18</sub> H <sub>13</sub> N <sub>7</sub> OS <sub>4</sub>	471	45.85	2,7 6	20.8 0	46.1 6	3.17	21,1 1		
[5]	180- 182	65	C <sub>15</sub> H <sub>9</sub> N <sub>7</sub> O 4S <sub>4</sub>	479	37.57	1.8 7	20.4 5	37.1 9	2.13	20.1 6		
[6]	174- 176	71	$\begin{array}{c} C_{19}H_{11}N_{7} \\ O_{4}S_{4} \end{array}$	529	43.10	2.0 8	18.5 2	42.8 1	1.83	18.7 7		
[7]	200- 202	66	$\begin{array}{c} C_{19}H_{10}N_8\\ O_6S_4 \end{array}$	574	39.72	1.7 4	19.5 1	39.4 9	1.95	19.3 4		
[8]	280- 282	70	$\begin{array}{c} C_{22}H_{15}N_{7}\\ O_{4}S_{4} \end{array}$	569	46.39	2.6 3	17.2 2	46.6 6	2.89	16.0 9		
[9]	190- 192	69	$\begin{array}{c} C_{26}H_{17}N_{7} \\ O_{4}S_{4} \end{array}$	619	50.40	2.7 4	15.8 3	50.6 7	2.61	16.0 6		
[10]	153- 155	70	$\begin{array}{c} C_{26}H_{16}N_8\\ O_6S_4 \end{array}$	664	46.98	2.4 0	16.8 6	46.7 1	2.33	17.0 4		

 Table (1): Melting points, percent yields and (C.H.N. )analysis
 of the prepared compounds (2-10)

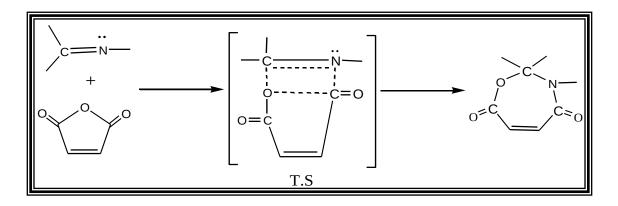
#### **Results and Discussion**

2-Amino-1,3,4-thiadiazole-5-thiol was converted to the corresponding diazonium salt which was directly converted to the coupling product ,azoaldehyde derivative [2], via coupling reaction with 2-Hydroxy benzaldehyde [1]dissolved in (10% v/v) sodium hydroxide solution<sup>(27)</sup>. FT-IR spectrum of compound [2], azoaldehyde derivative, showed appearance of weak absorption band at 1425 cm<sup>-1</sup>atributed to the  $\upsilon$  (N=N). Also ,disappearance of the two absorption bands at 3392cm<sup>-1</sup> and 3279 cm<sup>-1</sup> attributed to the (-NH<sub>2</sub>) stretching frequencies <sup>(14,28)</sup> and appearance of four sharp absorption bands at (3653-3520) cm<sup>-1</sup>atributed to the v(free O-H). The spectrum also showed appearance of four bands at (3443-3220) cm<sup>-1</sup>due to the v (bonding O-H). The weak band at 3160 attributed to the v(N-H). The two absorption band at 3074 and 3000 cm<sup>-1</sup> due to the v(C-H) $cm^{-1}$ aromatic. The bands at 2880cm<sup>-1</sup> and 2760 cm<sup>-1</sup> due to the v(C-H) aliphatic of aldehyde group (-CHO). The weak band at 2630 cm<sup>-1</sup> due to the v (C-S).Furthermore, the absorbtion band at 1674cm<sup>-1</sup> due to the v (C=O) of aldehyde group, we notice that the value was shifted to the lower frequency due to the intramolecular hydrogen bonding with Ortho hydroxy group<sup>(29)</sup>. The strong absorption band at 1545 cm<sup>-1</sup> due to the v (C=N) of the thiadiazole ring. The absorption band at 1480cm<sup>-1</sup> due to the  $v(C^{---}C)$  of aromatic ring. The absorption bands at 968,883,833 and 759 cm<sup>-1</sup> <sup>1</sup> were due to the  $\delta$ (C-H) aromatic out of plane . FT-IR spectrum also showed a strong band at 1240cm<sup>-1</sup> was attributed to the v(C=S) of thion tautomeric form. The spectrum also showed appearance of medium band at 1292 cm<sup>-1</sup> due to the  $\delta$ (O-H) in plane .Schiff bases [3] and [4] have been prepared through condensation reaction between azo aldehyde derivative [2] and each primary amines 2-Amine-1,3,4-thiadiazole -5- thiol and 5-(Benzylthio)-1,3,4-thiadiazole-2- amine respectively, in presence of small amount from glacial acetic acid as catalyst in absolute ethanol.

FT-IR spectrum of imine derivative [3] showed disappearance of the absorption band at 1674cm<sup>-1</sup> attributed to the v(C=O) of aldehyde group, also disappearance of the doublet absorption band at (3392 cm<sup>-1</sup>, 3279 cm<sup>-1</sup>) attributed to the asymmetric and symmetric stretching vibrations of (-NH<sub>2</sub>) group , respectively and appearance of strong absorption band at 1614cm<sup>-1</sup> attributed to the stretching vibration of exocyclic imine group (C=N).

FT-IR spectrum of imine derivative [4] showed disappearance of the absorption band at 1674cm<sup>-1</sup> attributed to the v(C=O) of aldehyde group ,also disappearance of the doublet absorption band at (3392 cm<sup>-1</sup>,3279 cm<sup>-1</sup>) attributed to the asymmetric and symmetric stretching vibrations of (-NH<sub>2</sub>) group ,respectively and appearance of strong absorption band at 1640 cm<sup>-1</sup> attributed to the stretching vibration of exocyclic imine group (C=N).

1,3-Oxazepine derivatives [5-10] have been prepared by using a pericyclic reaction type [2-5] cyclo addition reaction between imine group in compounds [3-4] as two membered component and maleic , phthalic and 3-Nitro phthalic anhydrides as five membered components to give seven - membered 1,3-oxazepine ring. [2-5] cycloaddition reaction is a concerted process proceeds via a single cyclic transition state and thus there is no intermediate in the process .Mechanism of 1,3-oxazepine ring formation has been shown in the following scheme:



#### Scheme(1):Mechanism of a pericyclic reaction type (2+5) cycloaddition reaction between imine group and maleic anhydride

FT-IR spectrum of 1,3-oxazepine derivative [5] showed disappearance of the strong band at 1614cm<sup>-1</sup> attributed to the stretching vibration of exocyclic imine group (C=N) and appearance of two strong absorption bands at1737 cm<sup>-1</sup> and1695cm<sup>-1</sup> attributed to the v(C=O) for lactone and lactam structures inside1,3-oxazepine ring ,respectively.

FT-IR spectrum of 1,3-oxazepine derivative [6] showed disappearance of the strong band at 1614cm<sup>-1</sup> attributed to the stretching vibration of exocyclic imine group (C=N) and appearance of two strong absorption bands at1740 cm<sup>-1</sup> and1685cm<sup>-1</sup> attributed to the v(C=O) for lactone and lactam structures inside1,3-oxazepine ring ,respectively.

FT-IR spectrum of 1,3-oxazepine derivative [7] showed disappearance of the strong band at 1614cm<sup>-1</sup> attributed to the stretching vibration of exocyclic imine group (C=N) and appearance of two strong absorption bands at 1745 cm<sup>-1</sup> and 1689cm<sup>-1</sup> attributed to the v(C=O) for lactone and lactam structures inside1,3-oxazepine ring ,respectively.

FT-IR spectrum of compound [7] also showed appearance of two strong absorption bands at 1560cm<sup>-1</sup> and 1345cm<sup>-1</sup> attributed to the asymmetric and symmetric stretching vibrations of (-NO<sub>2</sub>)

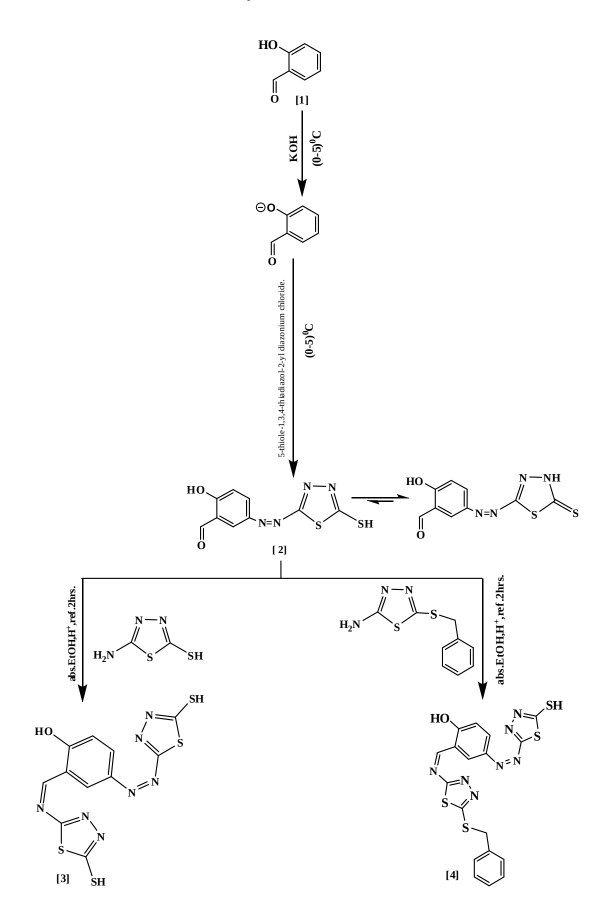
group<sup>(30)</sup>. Also appearance of strong absorption band at 877cm<sup>-1</sup> due to the stretching vibration of (C-NO<sub>2</sub>) bond is good evidence for the structure given to the product<sup>(31)</sup>.

FT-IR spectrum of 1,3-oxazepine derivative [8] showed disappearance of the strong band at  $1640 \text{cm}^{-1}$  attributed to the stretching vibration of exocyclic imine group (C=N) and appearance of two strong absorption bands at 1730 cm<sup>-1</sup> and 1708cm<sup>-1</sup> attributed to the v(C=O) for lactone and lactam structures inside1,3-oxazepine ring ,respectively.

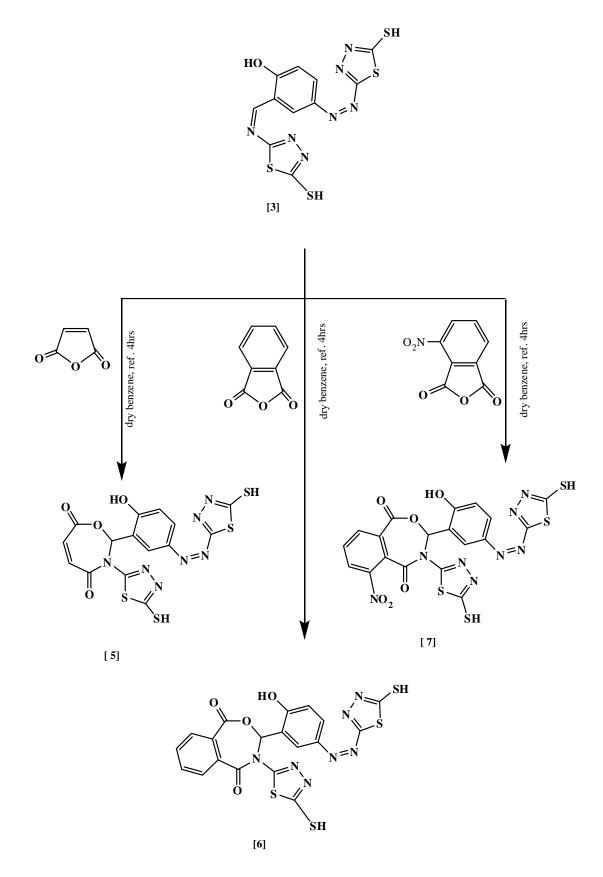
FT-IR spectrum of 1,3-oxazepine derivative [9] showed disappearance of the strong band at 1640 cm<sup>-1</sup> attributed to the stretching vibration of exocyclic imine group (C=N) and appearance of two strong absorption bands at 1720 cm<sup>-1</sup> and 1670 cm<sup>-1</sup> attributed to the v(C=O) for lactone and lactam structures inside1,3-oxazepine ring, respectively.

FT-IR spectrum of 1,3-oxazepine derivative [10] showed disappearance of the strong band at  $1640 \text{cm}^{-1}$  attributed to the stretching vibration of exocyclic imine group (C=N) and appearance of two strong absorption bands at 1780 cm<sup>-1</sup> and 1701cm<sup>-1</sup> attributed to the v(C=O) for lactone and lactam structures inside1,3-oxazepine ring ,respectively.

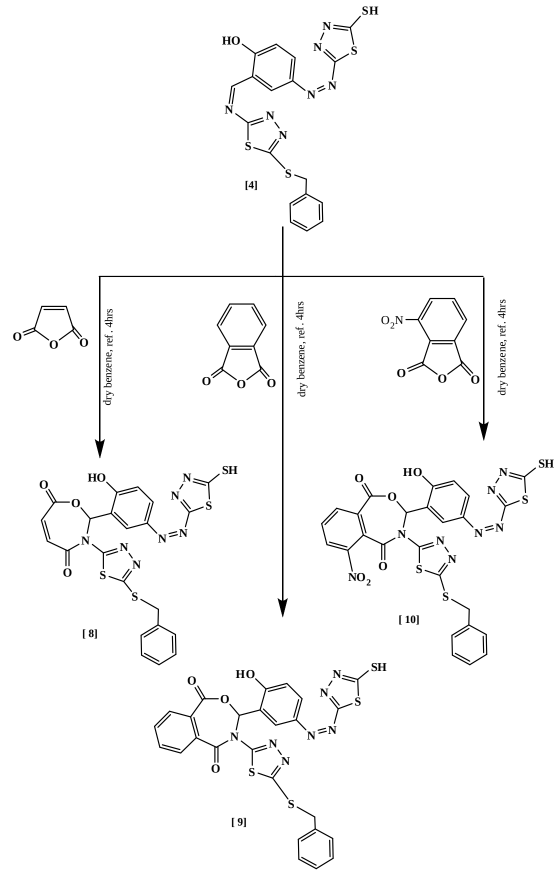
FT-IR spectrum of compound [10] also showed appearance of two strong absorption bands at 1560cm<sup>-1</sup> and 1335cm<sup>-1</sup> attributed to the asymmetric and symmetric stretching vibrations of (-NO<sub>2</sub>) group<sup>(29)</sup>. Also appearance of strong absorption band at 866cm<sup>-1</sup> due to the stretching vibration of (C-NO<sub>2</sub>) bond is good evidence for the structure given to the product.



Scheme (2): Reactions proceeding



Scheme (3): Reactions proceeding

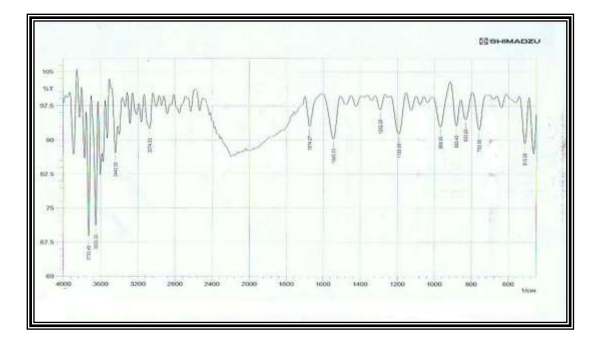


Scheme (4): Reactions proceeding

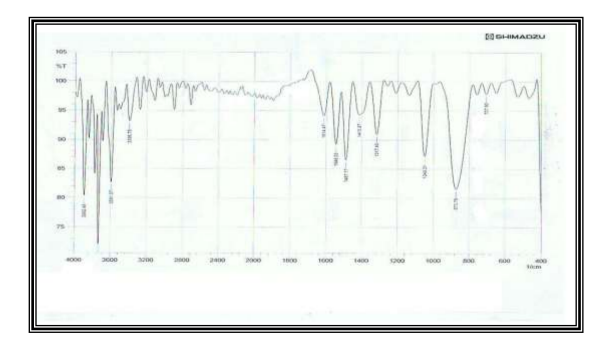
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Comp. no.	υ(-OH)	υ(-NH₂)	υ(N-H)	υ(C-H) arom.	U (C-H) aliph.	υ(S-H)	υ(C=O)	υ(C=N)	∪ (C <sup></sup> C) arom.	δ (O-H) in plane	υ(N=N)	υ(C-O)	υ(C= S)	υ(C-H) arom. ο.ο.p.	υ(C- NO₂)
[2]	3520-3653(sp) free(OH) 3220-3443(sp)bonding (OH)	-	3160(w)	3000(w) 3074(m)	2760- 2940 (w)	2550(w)	1674(s)	1545(s) endo	1480(m)	1292(m)	1425(w)	1070(m) 1035(w) phenol	1193 (s)	968(s) 883(s)	-
[3]	3591(sp) free(OH) 3260-3560(sp)bonding (OH)	-	3200(w)	3000(w) 3100(w) 3140(m)	2680- 2980 (w)	2500(w)	-	1614(s) exo 1545(s) endo	1487(s)	1317(s)	1413(w)	1250(w) 1130(w)	1205 (m)	833(s) 759(s)	-
[4]	3520-3600(sp) free(OH) 3260-3400(sp)bonding (OH)	-	3200(w)	3072(w) 3132(w)	2700- 2980 (w)	2530(w)	-	1640(s) exo 1543(s) endo	1600(w) 1498(s)	1321(s)	1423(w)	1240(w) 1136(m)	1209 (s)	889(s) 819(s) 758(s)	-
[5]	3498-3600(sp) free(OH) 3240-3420(sp)bonding (OH)	-	3195(w)	2993(w) 3060 (w) 3120(w)	2720- 2940 (w)	2560(w)	1737(s) lactone 1995(s) lactam	- 1546(s) endo	1485(s)	1315(s)	1430(w)	1220 1070 Phenol 1060 lactone	1160 (m)	981(m) 945(w) 877(s)	-
[6]	3529-3640(sp) free(OH) 3260-3500(sp)bonding (OH)	-	3240(w)	3000(w) 3150(w)	2720- 2940 (w)	2560(w)	1740(s) lactone 1685(s) lactam	- 1540(s) endo	1485(s)	1315(s)	1423(w)	1139(br) Phenol+lac tone	1200 (w)	870(s) 829(m) 775(w)	
[7]	3600(sp) free(OH) 3200-3520(sp)bonding (OH)	-	3160(w)	3000(w) 3040(w) 3100(w)	2665- 2960 (w)	2561(w)	1745(s) lactone 1689(s) lactam	- 1535(s) endo	1491(s)	1315(s)	1421(w)	1140(br) Phenol+lac tone	1220 (w)	830(m) 790(w)	877(s)
[8]	3533-3650(sp) free(OH) 3200-3443(sp)bonding (OH)	-	3140(w)	3000(w) 3070(w) 3100(w)	2725- 2930 (w)	2560(w)	1730(s) lacton 1708(s) lactam	- 1535(s) endo	1489(s)	1311(s)	1430(w)	1210(m) 1100(m) Phenol 1070(m) lactone	1175 (m)	856 (s) 810(s) 760(m)	
[9]	3525-3610(sp) free(OH) 3200-3427(sp)bonding (OH)	-	3160(w)	3000(w) 3040(w) 3080(w) 3120(w)	2677- 2960 (w)	2677(w)	1720(s) lacton 1670(s) lactam	- 1533(m) endo	1490(s)	1309(s)	1420(w)	1226(m) 1141(m) Phenol 1090(m) lactone	1180 (s)	966(s) 915(s) 761(m)	
[10]	3500-3610(sp) free(OH) 3200-3477(sp)bonding (OH)	-	3190(w)	3000(w) 3050(w) 3100(w)	2840- 2900 (w)	2545(w)	1780(s) lactone 1701(s) lactam	- 1545(s) endo	1490(s)	1313(s)	1425(w)	1240(m) 1163(m) Phenol 1080 lactone	1210 (w)	840(w) 810(w) 785(w) 760(w)	866(s)

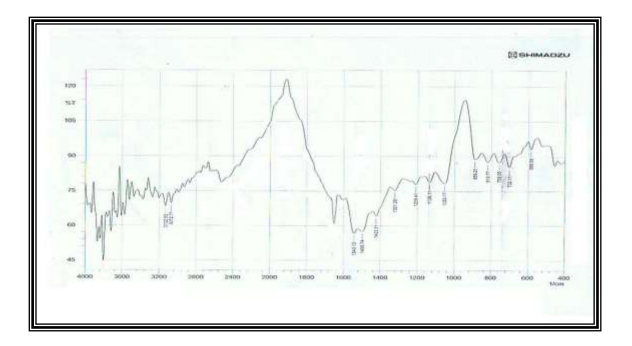
s=strong, w=weak, m=medium, sp=sharp, br=broad, o.o.p.= out of plane



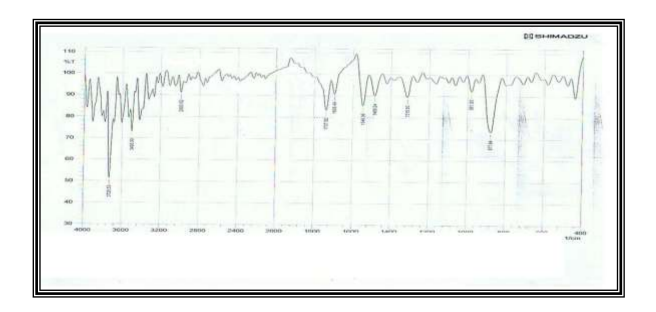
FT-IR spectrum of compound [2]



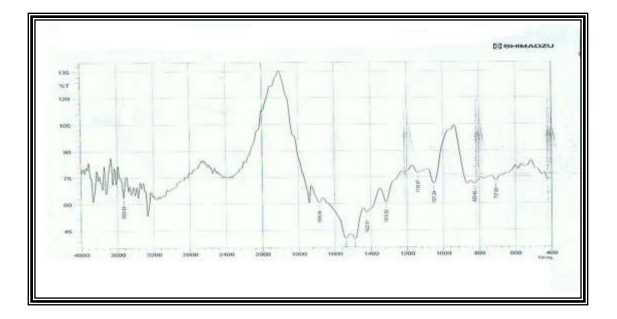
FT-IR spectrum of compound [3]



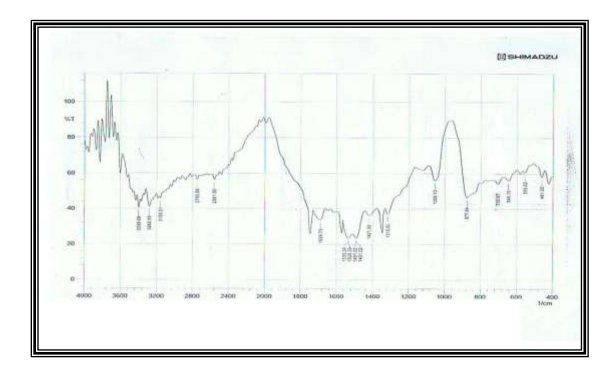
FT-IR spectrum of compound [4]



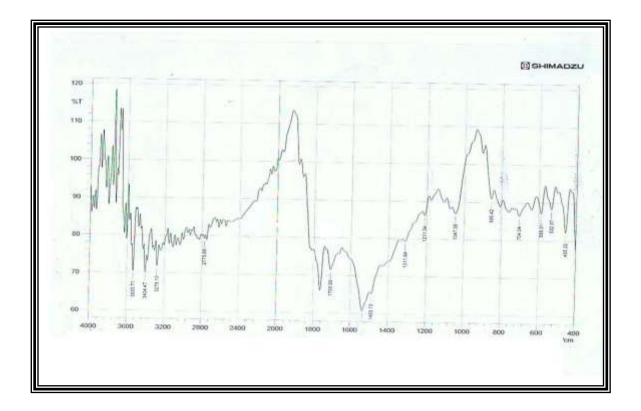
FT-IR spectrum of compound [5]



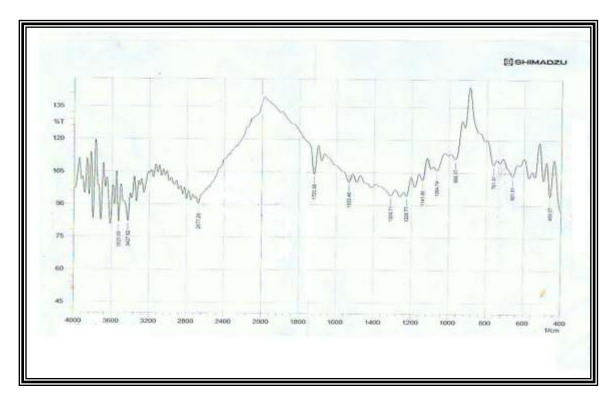
FT-IR spectrum of compound [6]



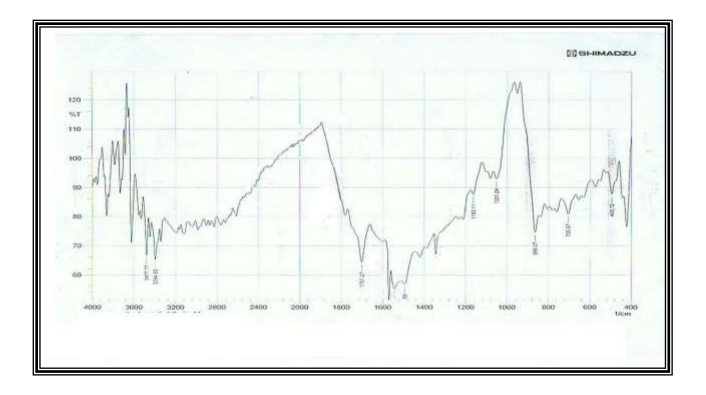
FT-IR spectrum of compound [7]



FT-IR spectrum of compound [8]



FT-IR spectrum of compound [9]



FT-IR spectrum of compound [10]

### References

- 1. 1. O. Buchardt, C.L. Pedersen and N. Harrit, J.Org. Chem., 37(23) 3592 (1972)
- 2. O. Tsuge, K. Oe and T. Ohnishi, Heterocycles, 19 (9) 1609 (1982).
- 3. M. A. Al-Hahithi, University of Sharjah, Journal of pure and applied science, 3(3) 25 (2006).
- 4. N. M. Al-Jamali, *Ph. D. Thesis*, University of Baghdad, 2008.
- 5. R. T. Haiwal, J. Kerbala University, 6(4) 216 (2008).
- 6. Z. H. Abood, J. Kerbala University, 7(1) 297 (2009).
- 7. O. H. Abid, National Journal of Chemistry, 3, 480 (2001).
- 8. I. S. Al-Shaibany, A. A. Mukhluss and Z. H. Abood, *J. Kerbala University*, proceeding of the 5<sup>th</sup> Sci. of K. U., 2009.
- 9. Z. H. Abood, J. Kerbala University, 8(1) 354 (2010).

10. M. H. Serrano-Wu, D. R. St. Laurent and Y. Chen., *Bioorganic and Medicinal Chemistry Letters*, 12 (19) 2757(2002).

11. L. Smith, W. C. Wong and A. S. Kiselyov, *Accepted for Publication 11 July*, 2006, *Bioorganic and Medicinal Chemistry Letters*, 2006, Available online, 2006.

- 12. P. G. Blain, *Toxicological Reviews*, 22(2) 103 (2003).
- 13. A. Al-Juboori, *M.Sc.Thesis*, Al-Mustansiriya University ,2005.
- 14. N. A. Salih, Ph.D. Thesis, Al-Nahrain University (2005).
- 15. A.A.Jarrahpour, M.Motamedifar, K.Pakshir, N.Hadi and M.Zarei, *Molecules*, 9,815 (2004).
- 16. S.B.Desai, B.Desai and K.R.Desai, *Heterocycl. Commun.*, 7,83 (2001).
- 17.P.H.Wang ,J. G. Keck ,E.J.Lien and M.M.C .Lai ,J .Med.Chem., 33,608(1990)..

18. W.M.Singh and B.C.Dash, Pesticides , 22,33 (1988).

19. P.G.More, R.B.Bhalvankar and S.C.Pattar, J.Indian Chem. Soc., 78,474(2001).

20.M.Verma, S.N.Pandeya, K.N.Singh and J.P.Stables, Acta Pharm., 54, 49(2004).

21. R.Noto, P.Meo, M.Geto and G.Weber, J.Heterocyclic Chem., 933,863 (1996).

22. A. S. Al-Ani, M.Sc. Thesis, Al-Mustansiriya University 2004.

23. Z. H. Abood, R. T. Haiwal and I. L. Kadum, J. Kerbala University, 6(4) 140 (2008).

24. F.Al-Omran, R.M.Mohareb and A.Abou El-Khair, J. Heterocyclic Chem., 39, 877 (2002).

25. C.Ainsworth , J.Am. Chem. Soc., 78, 1937 (1956).

26. J.A.Mohan, G.S.Anjaneyulu and R.Kiran, *Indian J.Chem.*, 27B, 128 (1988). 27.A.I.Vogel, "A text book of Practical organic chemistry ", 3<sup>nd</sup> eddition Longman group limited, London, 1973, p.622.

28. A.K.Gupta and H.K.Mishra, J. Indian Chem. Soc., 8, 508 (1981).

Chemistry'',2<sup>nd</sup> 29.D.H.Williams, I.Fleming, "Spectroscopic Organic method in edditionMcGraw-Hill Book Co.Limited,London(1973).