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Synthesis of (5,6,7,8)-Membered Rings of (Sulfur ,Nitrogen) via Cyclization of Imine Compounds

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Abstract

In this paper, synthesis of a series of compounds from hetero (Atoms, cycles) like (5,6,7,8-membered) ring via cyclo addition reaction of anil compound to produce compound [1-13], this reactions involved addition of carbonyl compounds like ((succinic acid, malonic acid, any compounds have two terminal from amine and thiol or carboxyl group)) to anil compounds to produce various membered rings. The structure of the newly synthesized compounds [1-13] were confirmed with (C.H.N)- analysis & substantiated with (FT.IR, H.NMR) data & melting points.

Keywords : eight membered , sulphur ,macrocycle , Imine ,heterocycle, five membered.

Introduction :

Hetromacrocycles by far are the largest classical division of organic chemistry .

Hetero cycles bearing nitrogen ,sulphur ,oxygen, constitute the core structure of a number of biologically interesting compounds ,some of them are pyrazoles , imidazoles ,which are structural subunits of several biologically active compounds⁽¹⁻⁴⁾</sup>.

Heterocycles have been used a scaffold to synthesize numerous therapeutic molecules , which are known for their medicinal importance as anticancer antibacterial ,antiseptics, & are known to be involved in a number of biological reactions such as inhibition of DNA ,RNA & protein synthesis⁽⁵⁻⁸⁾.

The utility of anil compounds lay in their usefulness as synthons in the synthesis of bio active molecules, it has ben found that the activity of hetero cycles increases on the incorporation of anil groups ⁽⁹⁻¹³⁾.

All chemicals used were supplied from BDH & Fluka- company, purity

Experimental:

*

99.5 % .

*

All measurements were carried out by :

1 - Melting points : electro thermal 9300 , melting point engineering LTD , U.K

2-FT . IR spectra : fourrier transform infrared shimadzu 8300-(FT . IR), KBr disc was performed by CO.S.Q.C. Iraq

3 - H.NMR-spectra and (C.H.N) – analysis : in center lab – institute of earth and environmental science , al – byat university , Jordan .

Synthesis of compound [1].

Condensation reaction by refluxing ethanolic mixture of equimolar amounts (0.1 mole ,12.0 gm) of p-methyl benzaldehyde & (0.1 mole ,9.7 gm) of 2-amino thiophene were react for (2hrs), the precipitate was filtered & recrystallized from ethanol to produce 83% of anil compounds [1].

Synthesis of compounds [2-5]:

A mixture of compound [1] (0.01 mole, 2.01 gm)was reacted with one of $\{(0.01 \text{ mole}, 1.38 \text{ gm}) \text{ of } 2\text{-mercapto} benzaldehyde}$, (0.01 mole, 1.19 gm of 2-amino benzaldehyde), (0.01 mole, 1.20 gm of salicyldehyde), (0.01 mole, 0.75 gm of alanine)}, respectively, under reflux for (10hrs) in presence of anhydrous 1,5-dioxan (100) ml, the precipitate was filtered, dried, & crystallized from absolute ethanol to produce % (86,84,82,86) respectively from compounds [2,3,4,5].

Synthesis of compounds [6-9]:

A mixture of compound [5] (0.01 mole , 2.58 gm)was reacted with one of $\{(0.01 \text{ mole}, 1.18 \text{ gm}) \text{ of succinic} acid)$, (0.01mole, 1.04 gm of malonic acid), (0.01 mole , 0.78 gm of acetyl chloride) ,(0.01mole , 1.06 gm of benzaldehyde)}, respectively , with reflux for (6hrs) in presence of absolute ethanol (100) ml with drops of sodium ethoxide.

the precipitate was filtered , dried , & crystallized from absolute ethanol to give % (82,85,87,86) respectively, from compounds [6,7,8,9].

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Synthesis of compounds [10,11]:

A mixture of compound [8] (0.01 mole, 3 gm)was reacted with one of $\{(0.01 \text{ mole}, 1.04 \text{ gm}) \text{ of malonic acid }), (0.01 \text{ mole}, 1.18 \text{ gm of succinic acid })\}$ respectively under reflux for (6hrs) in presence of absolute ethanol (100) ml with drops of sodium ethoxide, the precipitate was filtered, dried, & crystallized from absolute ethanol to produce % (87,85) respectively, from compounds [10,11].

Synthesis of compounds [12,13]:

A mixture of p-methyl benzal dehyde (0.1mole ,1.2 gm)with P-chloro acetanilide(0.1 mole , 1.69gm) in ethanol (100) ml& 2ml of (3% sodium hydroxide solution)with stirring for (5hrs) at room temperature ,then refluxed for (8hrs), , the precipitate was filtered , dried ,& crystallized from ethanol to produce 88 % of compounds [12]. To prepare compound [13], mixture of compound [12] (0.01 mole , 2.71 gm) & hydrazine(0.01 mole , 0.50 gm) under reflux for (7hrs) in presence of absolute ethanol (100) ml, the precipitate was filtered , dried ,& crystallized from ethanol to produce % 86 of compound [13].

Scheme Reaction :





Results & Discussion :

In this study, we wish to report on anew approach for preparation of hetero atoms cycles (S,N,O) & hetero cycles (5,6,7,8-membered) ring from compounds [1-13].

Their FT.IR-Spectrum showed an absorption band at (1620) cm⁻¹ in compound [1] due to the (CH=N) anil group ,which disappear & other bands are appear at {(1685-1698) cm⁻¹ for amide⁽¹⁵⁾ group Q

(-C-N),(1530-1545) cm⁻¹ for (C-N) endocycle & bands due to(C-S, C-NH, C-O, CH-NH)} in formed compounds [2-13] also new bands appeared such as (C=CH) due to alkene in compounds [9,12], bands at (1710-1725)cm⁻¹ due to carbonyl of ketone in forned cycles in compounds [6-11], & other bands are summarized in table (1) & figure (1-4).

Their H.NMR-Spectra showed signal at 8.89 \int for proton of azomethine group (CH=N) in compound [1] which disappear & new signals appear at (5.96 \int for CH-S)⁽¹⁶⁾ in compound [2], (3.9 \int for CH-O) in

0

compound[4], $(3.09 \int - 3.19 \int$ for CH-NH in cyclic compounds[3,5-11,13], $(9.72 \int$ for proton of amide HN-C-) in compound [12] as result of formed cycles, & other data of functional groups show in the following, Table (2) & figure (5-8).

Their (C.H.N)- analysis & melting points , it was found from compared the calculated data with experimentally data of these compounds ,the results were compactable , the data of analysis , M.F & melting points are listed in table (3) .

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ComP.	I.R _(KBR) (Important Groups)					
no.						
[1]	(CH=N) azomethine group : 1620					
[2]	(O=C-N) amide of endocyclic :1698,(C-N) endocyclic :1537,(C-S) endocyclic :675,1404, (C=C) aromatic:1581.					
[3]	(O=C-N) amide of endocyclic :1690,(C-N) endocyclic :1540 ,(NH): 3320.					
[4]	(O=C-N) amide:1698,(C-N) endocyclic:1540,(C-O-C):1050.					
[5]	(O=C-N) amide:1685,(C-N) endocyclic:1535,(NH): 3330, (CH) aliphatic:2930.					
[6]	Â					
	(O=C-N) amide:1690,(C-N) endocyclic :1530 ,(
[7]	Å					
	(O=C-N) amide:1680,(C-N) endocyclic :1498 ,() ketone: 1717, (CH) aliphatic :2925 .					
[8]	(O=C-N) amide:1690,(C-N) endocyclic :1544 ,(CH) aliphatic :2930.					
[9]	(O=C-N) amide:1695,(C-N) endocyclic :1545 ,(NH):3320,(=CH) alkene:3080 .					
[10]	Â					
	(O=C-N) amide:1686,(C-N) endocyclic :1537 ,() ketone: 1720, (CH) aliphatic :2920 .					
[11]	ĝ					
	(O=C-N) amide:1690,(C-N) endocyclic :1540 ,(
[12]	(O=C-N) amide:1695,(=CH) alkene: 3050 .					
[13]	(C=N) azomethine:1620,(N-N) endocyclic :1400 ,(NH) : 3330, (CH) aliphatic :2940 .					

Table (1): (FT.IR)-data (cm⁻¹) of compounds [1-13].

Table (2): H.NMR-data(β_{ppm}) of compounds [1-13].

Comps	H.NMR _(DMF) (Important peaks)
[1]	8.89 {1H,(CH=N)} proton of azomethine group.
[2]	6.34-7.8 (Ar-H) , 5.96 (CH-S).
[3]	6.6-7.8 (Ar-H) ,3.11 (CH-NH) .
[4]	6.36-7.3 (Ar-H) , 3.9 (CH-O) .
[5]	3.09 (CH-NH), 9.96 (CH ₂ —C—N).
[6]	3.1 (1H ,CH-N), 12.2 (O=C-CH ₂ -) ,10.2 (CH ₂ -C-N).
[7]	3.19 (1H,CH-N), 12.29 (2H, O=C-CH ₂).
[8]	3.1 (1H, CH-N), 10.1 (CH2-C-N), 10.5 (CH3-C-N).
[9]	2.3 (1H ,CH=C), 3.4 (CH-NH), 6.4-7.2 (Ar-H).
[10]	3.12 (1H,CH-N), 12.3 (2H, O=C-CH ₂ -C=O).
[11]	3.3(1H,CH-N),12.59 (CH ₂ — C), 12.72(0=C-CH ₂ -C=O).
[12]	O II 9.72 (
[13]	1.2 (2H,CH ₂ -C), 3.2 (CH-NH), 6.4-7.2 (Ar-H), 1.2 (CH ₃).

Comps	M.F	M.P Nan	Name of compounds	Calculation/Found		
		(C°)		С%	Н%	N%
[1]	$C_{12}H_{11}N_1S_1$	161	2-(4 ⁻ Toluine)- thiophenidine .	71.641	5.472	6.965
				71.342	5.211	6.654
[2]	$C_{19}H_{15}NOS_2$	242	2-(4 ⁻ Toluine)- 3-thiophenidine-	67.655	4.451	4.154
			5,6- benzo-1,3-Thiazane-4-one.	67.462	4.318	4.310
[3]	$C_{19}H_{16}N_2OS$	218	2-(4 ^T oluine)- 3-thiophenidine-	71.25	5.00	8.750
			5,6- benzo-pipyrimidine-4-one.	71.012	5.021	8.592
[4]	$C_{19}H_{15}NO_2S$	235	2-(4 ⁻ Toluine)- 3-thiophene-1-	71.028	4.672	4.361
			oxo-5,6- benzo-pipyrimidine-4-	71.320	4.711	4.451
			one.			
[5]	$C_{14}H_{14}N_2OS$	195	2-(4 ⁻ - Toluine)- 3-thiophene	65.116	5.426	10.852
			Imidazoline-4-one.	65.014	5.201	10.312
[6]	$C_{18}H_{16}N_2O_3S$	238	3-(2 ⁻ Thiophene) -2-(4 ⁻ Toluine)-	63.529	4.705	8.235
			1,5-(2 ⁻ ,5 ⁻ - dione–azane)–	63.342	4.611	8.301
			imadazol-4-one.			
[7]	$C_{17}H_{14}N_2O_3S$	222	3-(2 ⁻ Thiophene)-2-(4 ⁻ -Toluine)-	62.576	4.294	8.588
			1,5-(2 ⁻ ,4 ⁻ -di one –azolidine)–	62.328	4.271	8.401
			imadazol-4-one.			
[8]	$C_{16}H_{16}N_2O_2S$	200	2-(4 ⁻ - Toluine)-3-thiophene-1-	64.00	5.333	9.333
			aceto- Imidazoline-4-one.	64.018	5.350	9.114
[9]	$C_{21}H_{18}N_2OS$	210	3-(2 ⁻ Thiophene) -2-(4 ⁻ Toluine)-	72.832	5.202	8.092
			1,5-(2,4,6-Tri one -azecane)-	72.672	5.151	8.001
			imadazol-4-one.			
[10]	$C_{19}H_{16}N_2O_4S$	240	3-(2 ⁻ Thiophene)-2-(4 ⁻ -Toluine)-	61.956	4.347	7.608
			1,5-(2 ⁻ ,4 ⁻ ,6 ⁻ -Tri one –azepane)–	61.813	4.238	7.516
			imadazol-4-one.			
[11]	$C_{20}H_{18}N_2O_4S$	229	2-(4 ⁻ Toluine)- 3-thiophene-5-	62.827	4.712	7.329
			styrene- Imidazoline-4-one.	62.719	4.623	7.113
[12]	$C_{16}H_{14}N_1O_1Cl$	165	N-(4-Chloro phenyl)-3-Toluine	70.718	5.156	5.156
			acrylamide.	70.651	5.08	5.201
[13]	C ₁₆ H ₁₆ N ₃ Cl	176	4-[(5 ⁻ -Toluine-4 ⁻ ,5 ⁻ -dihydro	67.250	5.604	14.711
			pyrazol-3 ⁻ -yl)amino] chloro	67.161	5.587	14.511
			benzene.			

Table (2): physical properties & (C.H.N)- analysis of compounds [1-13].

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