

# An Efficient and Solvent Free Synthesis of N-Aryl 2,3-Dihydro-4H naphtho-[2,1-e] 1,3-oxazines

#### Mohammad S Al-Ajely\* and Ahmed M Noori

Department of Chemistry, College of Education for girls, Mosul University, Mosul, Iraq

\*Corresponding author: Mohammad S Al Ajely, Department of Chemistry, College of Education for girls, Mosul University, Mosul, Iraq

# ARTICLE INFO

Received: 🕮 August 03, 2020

Published: 🕮 August 13, 2020

**Citation:** Mohammad S Al-Ajely, Ahmed M Noori. An Efficient and Solvent Free Synthesis of N-Aryl 2,3-Dihydro-4H naphtho-[2,1-e] 1,3-oxazines. Biomed J Sci & Tech Res 29(3)-2020. BJSTR. MS.ID.004815.

# ABSTRACT

Oxazine compounds have proved to have many pharmaceutical applications and most of these compounds now a days are used as drugs. For the importance of this class of heterocyclic compounds we are here investigate the synthesis of new derivatives of 1,3-oxazines using solvent free one pot three component system in a drug discovery program ,so starting from  $\beta$ -Naphthol, formaldehyde and aromatic amines in presence of zarconyl chloride as catalyst. compounds 1-9 were synthesized, Benzo 1,3 diazines (10-14) were also synthesized from their corresponding 1,3 oxazines. These compounds were characterized by IR, some representative by [1] HNMR and were discussed.

Keywords: Aryl; 1,3-Naphthoxazines; Solvent free; synthesis

#### Introduction

Due to their importance in biological applications oxazines have drew attention of many researchers to investigate this type of hetrocyclic compounds. these researchers have succeeded to prepare different types of oxazines using different methodologies [1-10] most of the starting material is anthranilic acid or its derivatives with aldehydes, isocyanate, acetic anhydride or with oxazolones [11]. Some other researchers used phenal, amine and aldehydes to the synthesis of these oxazine compounds [12]. Other researchers used chalcones for the synthesis of this type of compounds [13,14] or from 2-Iodoaryl Azides and Amines [15]. Among the studied reactions of oxazine compounds were opening and reclosing of the oxazine ring which is the other way for the synthesis of new oxazine derivatives [16]. The research work on studying the biological activities of some oxazine compounds revealed that most of these compounds have several pharmacological applications for using these compounds as drugs [17-19]. In addition the latest studies on oxazine compounds showed the ability to stop the cancer cell growth and repairing the DNA strands [20] and according to the above facts we studied the synthesis of some new compounds using green technique for this synthesis .These new compounds will be our goal for studying their biological activities in our ongoing drug discovery program.

#### Experimental

All melting points were uncorrected using electro thermal SMP30 UK melting point apparatus. IR spectra were recorded using Alpha (ATR) instrument. [1] HNMR spectra were recorded using Varian Agilant (USA) 300MHz instrument, DMSO as solvent. All chemical was supplied by sigma –Aldrich, BDH and Fluka companies. Compounds 10 a, b was synthesized following some published procedure [21].

General Procedure for the Synthesis of Naphthaoxazine Copmounds (1-9)

**Table 1:** Physical properties of compounds (1-9).

Comp. No.	Ar	Molecular formula	M.wt m.p.(°C)		Yield (%)	Colour
1	4-chlorophenyl	C <sub>18</sub> H <sub>14</sub> CINO	295.76	98-100	73	Yellow

2	H <sub>3</sub> C-V 4-methyl-2-pyridinyl	$C_{18}H_{16}N_2O$	276.33	115-118	37	Brown
3	2-benzothiazolyl	$C_{19}H_{14}N_2OS$	318.39	109-114	83	Dark Brown
4	H <sub>3</sub> C 	$C_{19}H_{16}N_2O_3$	320.34	158-164	33	Orange
5	Br	C <sub>18</sub> H <sub>14</sub> BrNO	340.21	113-117	79	Brown
6	Cl NO <sub>2</sub> 2-chloro-4-nitrophenyl	$C_{18}H_{13}ClN_2O_3$	340.76	151-153	59	Yellow
7	4-amino phenyl	$C_{18}H_{16}N_2O$	276.33	218-222	67	Brown
8	4-nitro phenyl	$C_{18}H_{14}N_2O_3$	306.32	160-164	43	Yellow
9	Br 2-bromo phenyl	C <sub>18</sub> H <sub>14</sub> BrNO	340.21	340.21	222	Orange

Formaldehyde (0.2mol.), Zarconyl chloride (ZrOCL2,8H<sub>2</sub>O) (0.2mol.),2-naphthol (0.1mol.) and aromatic amine (0.1mol.). This mixture was grinding by porcelain mortar for 30 min, after that dichloromethane ( $CH_2CL_2$ ) was then added. The organic layer was then separated and washed twice with brine then with water. The organic layer was separated. Evaporation of the solvent by rotary evaporator afforded a crude product which was recrystallized from minimum amount of methanol. Physical properties were listed in the following Table 1.

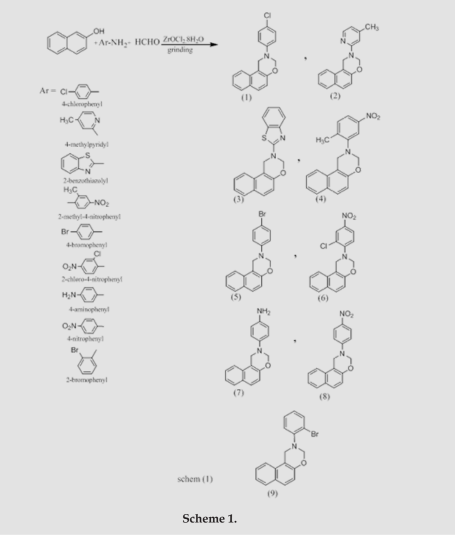
# General Procedure for the Preparation of N-Substituted 2-Methyl, Aryl Benzo-1,3-Diazine-4-one (11-24)

compound 10a or 10b (0.1mol), aromatic or aliphatic amine (0.1mol), This mixture was refluxed for 6h, cooled. The whole mixture was then powered on cold water (50ml) with stirring. The precipitate was filtered off and dried then crystallized from benzene. The physical properties for the synthesized compound were illustrated in Table 2 (Schem 1,2).

Table 2: Physical	properties of compounds	(10-13).
-------------------	-------------------------	----------

Comp. o.	Ar	Molecular formula	M.wt	m.p.(°C)	Yield (%)	Colour
10	2-naphthayl	C <sub>21</sub> H <sub>17</sub> NO	299.37	86-92	83	brown
11	2-benzothiazolyl	$C_{18}H_{14}N_2OS$	306.38	193-196	92	white

12	2-thiazolyl	$C_{14}H_{12}N_2OS$	256.32	171-173	89	Yellowish white
13	4-methyl-2- pyridinyl	$C_{17}H_{16}N_2O$	264.32	154-157	90	white



#### **Results and Discussion**

It is worth to state here that power sonic 405 micro processes -controlled bench-top ultra-sonic cleaner was used for ultrasonic chemical condensation affording nearly the same yield within the same given time which is another green technique for the synthesis of these compounds .As it was mentioned in the experimental part .compounds 10 a,b were prepared and showed the same melting points, for 10a was found; 79-81 published: 81- 82°C, while for 10b it was found 120°C the published one at 121-122°C with identical IR data [21].

# N-Aryl 2,4H(1,2E) (1,3) Napha Oxazine Compounds (1-9)

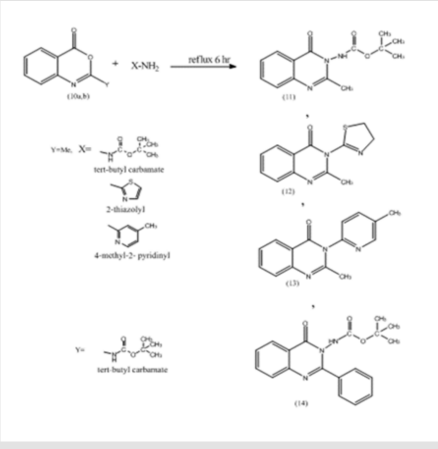
These compounds were synthesized using similar procedure

[22] and were characterized by IR and showed the following main absorption bands ( $v_{max}$  cm<sup>-1</sup>) :3020-3092 for CH, sharp bands at (1654- 1624) for C=N, 1589 -1454 for C=C, C=c Aromatic . While C=N appeared at 1390-1328, C-O-C at 1250-1022 other band were illustrated in Table 3. [1] HNMR for individual compounds were as follow:

#### Compound (1)

In which the aryl group is p-Chloro phenyl. The Aromatic protons were 6 types as follows :7.99, 7.83, 7.7.81ppm. for carbon 8 of naphthyl ring, 7.71, 7.7ppm. for proton 5. this ring 7.54, 7.40ppm for protons 6,7 while protons 4,3 appeared at 7.12, 7.11 and 7.054, 7.02ppm. Phenyl protons appeared as AB (q) at, 6.79, 6.78 ppm.  $CH_2$  protons of the oxazine ring between oxygen and nitrogen appeared at 5.54, 5.5ppm. and the other  $CH_2$  protons appeared at 4.96,

4.77ppm. protons of naphthyl ring protons appeared as two triplet signals which can be easily differentiated from other protons.



Scheme 2.

Table 3: Physical properties of compounds (10-13).

Comp. No.	Ar		IR υ cm <sup>-1</sup> (neet)								
comp. No.		Ar C-H	C=N	C==C ArC=C,	C-N	С-О-С	Others				
1	4-chlorophenyl	3059		1588,1482	1366	1220,1094	C-Cl 736				
2	H <sub>3</sub> C- 4-methyl-2-pyridinyl	3048	1628	1586 1505,1454	1328	1204,1157	C-H alph 2910				
3	2-benzothiazolyl	3020	1654	1572,1462	1381	1256,1066	C-S 749				
4	$\underbrace{H_3C}_{2\text{-methyl-4-nitroaphenyl}}NO_2$	3051		1588 1508,1464	1390	1215,1046	N-0 sym1259 asym1508				
5	Br	3061		1586 1551,1478	1368	1219,1064	C-Br 486				

8	4-nitro phenyl	3063	 1589 1502,1481	1323	1235,1060	NO- sym1268 asym1589
7	4-amino phenyl	3054	 1510,1468	1375	1222,1059	3278 3364
	2-chloro-4-nitrophenyl		1549			N-H <sub>2</sub>
0		0072	1503,1461	1027	1200,1022	asym1503
6		3092	 1587	1329	1225,1022	N-0 sym1273

# Compound (2), Ar is (4-methyl-2-pyridyl)

This compound also showed 6 signals for the aromatic protons resonated at 8.1, 8.0, 7.5, 7.4, 6.85ppm, for 8,5,6, 7,3,4 protons while the phenyl protons appeared at 7.9, 7.89 ppm. for AB like system, 7.53, 7.52 ppm. for the rest protons nearly equivalents oxazine protons, CH2 resonated at 5.7, 5.68 ppm. for protons 4,3 respectively.

#### Compound (6) Ar is (2-Bromo phenyl)

Protons NMR chart for this compound also showed the following signals: 8.0, 8.02, 7.5, 7.45, 7.65, 6 .91 ppm. for naphthyl ring protons of 8,5 protons, 6,7 and 3,4 protons respectively for oxazine ring protons 6.1, 5.05ppm while the phenyl ring protons 7.61 ortho to carbon bearing Br substituent, 6.67, 7.21, 6.65ppm. for the next protons the value of 6.65ppm. is for proton of the ring adjacent to the nitrogen attachment.

#### Compound (8) where Ar is (p-Amino phenyl)

This compound showed the following resonating signals :8.1, 8.02, 7.54, 7.4, 7.64, 6.9ppm. for 8,5,6,7 and 4,3 protons respectively. Oxazine protons showed two signals (with and opposite side of ring plane) at 6.1, 6.05 ppm. for CH<sub>2</sub> between N, O atoms and near

Table 4: IR data for com	npounds (11 <b>-</b> 14).

N atom respectively. The phenyl protons appeared as two types of protons; near nitrogen oxazine attachment at 6.1, 6.05ppm, near NH<sup>2</sup> on both sides 6.45, 6.46ppm. while the phenyl NH<sub>2</sub> protons resonated at 5.5.

## Compound (9) where Ar is (o-methyl, P-Nitro phenyl)

This compound was also giving 3 types of protons of the NMR spectra of naphthyl ring protons resonating at 8.1, 8.02, 7.54, 7.41, 7.65, 6.91ppm. for 8,7,6,5 and 3,4 protons. The oxazine ring protons were resonating at 5.0, 6ppm. for CH<sub>2</sub> near N atom and between N, O atoms respectively. Phenyl ring protons of two types resonating at 7.89, 7.87ppm. near NO<sub>2</sub> group, 6.0ppm. near nitrogen attachment of the oxazine ring, CH<sub>2</sub> of oxazine near nitrogen 5.0 ppm, 6.1ppm. for the CH<sub>2</sub> protons of the oxazine between N, O atoms while the protons of the phenyl ring appeared at 2.1 ppm.

# N-substituted, Methyl or Phenyl Benzoxazine -4- one Compounds (11-14)

The detail of IR data for these compounds were shown in Table 4. The NMR spectra of some selected samples were characterized by the following resonating signals:

Comp. No.	v		IR v cm <sup>-1</sup> (neet)						
Y	Х	N-H	Ar C-H	alph C-H	C=0	c <sub></sub> cec=N	C <sub></sub> C <sub>Ar C=C</sub> ,	C-N others	
11	O CH <sub>3</sub>						1569	1266	
	N C O C CH <sub>3</sub>	3233	3011	2977	1740	1668		C-0	
CH <sub>3</sub>	<i>tert</i> -butyl carbamate						1506,1435	1151	
12	S S	3279	3130	2924	1741	1689	1637	1257	
CH <sub>3</sub>	2-thiazolyl	5275	5150	2724	1741	1009	1512,1445	1237	
13	CH3	3271	3031	2810	1671	1585	1585,1500,1428	1289	
CH <sub>3</sub>	4-methyl-2- pyridinyl								

14	CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub>	3326	3067	2977	1742	1639	1590 1507,1436	1233 C-0
-Ph	<i>tert</i> -butyl carbamate							1153

# Compound (11) 2-mrthyl –N-t-butyl -4-oxo benzo -1,3 – Diazino Carbamate

## Compound (12) N-2thiozolyl-4 benzo-1,3-Diazine-4-one

The benzene ring protons were found at 8.12, ppm. for carbon 5 proton, 7.65, for carbon 8 proton. The other protons of this ring were resonated at 7.86, 7.65, 7.61ppm for protons at carbon, 7,6 respectively, protons of the t-Butyl group appeared at 1.48ppm, 3.16 ppm for  $CH_3NH$  at 10ppm.

Benzene ring protons appeared at 8.4, 8.0, 7.62ppm assigned to protons 2,5,8 and at 7.7, 7.6ppm. for protons 7,6 respectively. The thiozolyl protons appeared at 7.2 (near nitrogen proton), the other at 6.7 ppm, 3.18ppm. for  $CH_3$  of Diazine ring protons and 2.35ppm. for pyridyl protons (Figures 1,2).

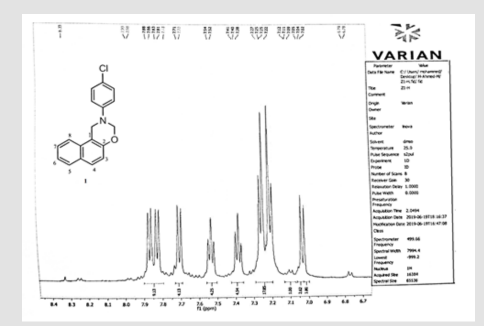
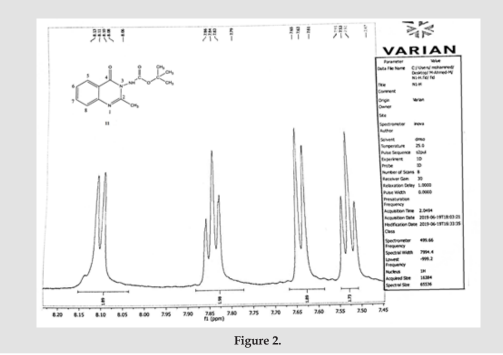


Figure 1.



#### Conclusions

We conclude from the above study that all the synthesized compounds were a really oxazine compounds that is comes from their structure's elucidation using the spectral data for both green techniques used on their synthesis. The screening effects will be studded and their results will be our next paper.

# Acknowledgment

We greatly acknowledge the Iraqi Ministry of higher Education for providing Ahmed M. Noori a scholarship to perform this work which is part of his MSc Thesis.

#### References

- 1. EV Gromachevskaya, TP Kosulina, VG Kulnevich, VP Smolyakov (1997) Chemistry and Technology of Furan Compounds, Krasnodar 21.
- EV Gromachevskaya, TP Kosulina, GD Krapivin, VG Kulnevich (1993) Reaction of o-aminophenyldiphenylcarbinol with unsaturated aldehydes, Chemistry of Heterocyclic Compounds 29: 1109-1110.
- EV Gromachevskaya, VG Kulnevich, TP Kosulina, VS Pustovarov (1988), 4H-3,1-benzoxazines. 2.Synthesis of 2,4-substituted 1,2-dihydro-4H-3,1-benzoxazines, Chemistry of Heterocyclic Compounds 24: 692-697.
- 4. TP Kosulina, EV Gromachevskaya, VG Kulnevich (1993) Reaction of o-aminophenyldiphenylmethanol with  $\alpha$ -acetylenic ketone, Chemistry of Heterocyclic Compounds 29: 1232-1233.
- F Kienzle (1983) A facile synthesis of 1,4-dihydro-1-alkyl-2H-3,1-benzoxazines and related compounds, Tetrahedron Letters 24(21): 2213-2216.
- W Nijhuis, W Verboom, S Harkema, DN Reinhoudt (1989) Synthesis of 3,1-benzozazines and 3,1-benzothiazines, Recueil des Travaux Chimiques des Pays-Bas 108: 147.
- 7. RS Tiwari A Uperti, RK Satsangi (1981) Synthesis of 2-Hippuryl-3-Aryl-Quinazolinones, Journal of the Chemical Society of Pakistan 3: 215.
- 8. EP Papadopoulos, CD Torres (1982) Convenient preparation of N-substituted 2-amino 4H-3, 1-benzoxazin-4-one and 3-substituted 2, 4(1H, 3H) quinazoinediones, Heterocyclic chemistry 19(2): 269-272.
- 9. U Rose (1991) 2-Aryl-substituted 4H-3,1-benzoxazin-4-ones as novel active substances for the cardiovascular system, Journal of Heterocyclic Chemistry 28(8): 2005-2012.

#### ISSN: 2574-1241

#### DOI: 10.26717/BJSTR.2020.29.004815

Mohammad S Al-Ajely. Biomed J Sci & Tech Res



This work is licensed under Creative Commons Attribution 4.0 License

Submission Link: https://biomedres.us/submit-manuscript.php

- 10. H Wamhoff, S Herrmann, S Stoelben, M Nieger (1993) A new, general approach for the synthesis of heteroannulated 3,1-oxazin-4-ones, Tetra-hedron 49(3): 581-594.
- 11. M O Osman Habib, M Hussein Hassan, B Evelin B Moawad, E-LMekabaty Ahmedl (2012) Synthetic Approaches and Biological Evaluation of Some New Sulfonate Ester-Containing Imidazolone Derivatives American Journal of Organic Chemistry 2(4): 79-86.
- 12. Cheng-Xi Zhang, Yu-Yuan Deng, Yi-Yang Zhang, Po Yang, Yi Gu (2015) Study on products and reaction paths for synthesis of 3,4-dihydro-2H-3-phenyl-1,3-benzoxazine from phenol, aniline and formaldehyde Chinese. Chemical Letters 26(3): 348-352.
- 13. ZA Sallal (2014) Synthesis and characterization of new Oxazine , Thiazine and Pyrazol derived from chalcones, Baghdad Science Journal 11(2): 477-485.
- 14. G Chaitra, RM Rohini (2018) Synthesis of 1,3-Oxazine derivative from Chalcone and Screening for their Anti-Oxidant and Anti-Inflammatory activity. International Research Journal of Pharmaceutical and Biosciences 4(6): 20.
- 15. Wu Youcan Zhang, Zhiping Yin, Hai Wang, Xiao-Feng (2019) Pd/C-Catalyzed Carbonylative Synthesis of 2. Aminobenzoxazinones from 2-Iodoaryl Azides and Amines, Organic Letters pp. A-E.
- AM Noori (2019) Synthesis of some New Oxazin Compounds and the study of their reactions, MSc Thesis, Mosul university, Mosul-Iraq.
- Nadeem Siddiquia, Ruhi Alia, M Shamsher Alama, Waquar Ahsana (2010) Pharmacological Profile of Benzoxazines: A Short Review J. Chem. Pharm. Res 2(4): 309-316.
- Zilong Tang, Zhonghua Zhu, Zanwen Xia, Hanwen Liu, Jinwen Chen, et al. (2012) Synthesis and Fungicidal Activity of Novel 2,3-Disubstituted-1,3-benzoxazines, Molecules 17(7): 8174-8185.
- D Patronea, F Nicholas, P Batesa, J Patronea, D Pelza (2016) Identification and Optimization of Anthranilic Acid-Based Inhibitors of Replication Protein A, Chem Med Chem 11(8): 893-899.
- 20. JC de Wouter Bruijna, A Jos Hagemanb, Carla Araya-Cloutiera, Harry Gruppena, Jean-Paul Vinckena (2018) QSAR of 1,4-benzoxazin-3-one antimicrobials and their drug design perspectives, Bioorganic & Medicinal Chemistry 26(23-24): 6105-6114.
- MA Abd-Elhakeem, AM Elsayed (2013) Synthesis and antimicrobial activity of some new 2,3-disubstituted quinazoline-4(3H)-ones derivatives, Journal of Chemical and Pharmaceutical Research 5(5): 275-279.
- 22. Amol H Kategaonkar, Swapnil S. Sonar, Rajkumar U (2010) An Efficient Synthesis of 3,4-Dihydro-3-substituted-2H-naphtho[2,1-e] [1,3] oxazineDerivatives Catalyzed by Zirconyl (IV) Chloride and Evaluationof its Biological Activities, Bull. Korean Chem. Soc 31(6): 1657-1660.



ISSN: 2574

#### Assets of Publishing with us

- Global archiving of articles
- Immediate, unrestricted online access
- Rigorous Peer Review Process
- Authors Retain Copyrights
- Unique DOI for all articles

https://biomedres.us/