Photo-degradation Study for Some 4-aminoantipyrine derivatives by used (Cs-137) as Gamma-Ray Source در اسة التحطم الضوئي لبعض مشتقات 4-أمينو أنتي بايرين بأستخدام (Cs-137) كمصدر لأشعة كاما

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Abstract

The compounds {4-Benzylidene-amino)-1,5-dimethyl-2-phenyl-1,2-dihydro-pyrazol-3one}, {4-[(4-Hydroxy-benzylidene)-amino]- 1,5-dimethyl-2-phenyl-1,2-dihydro-pyrazol-3-one} and {4-[(2-Hydroxy-benzylidene)-amino]- 1,5-dimethyl-2-phenyl-1,2-dihydro-pyrazol-3one}were prepared as [4-aminoantipyrine] derivatives by condensation reaction . These compounds were characterized by FTIR spectra , UV.-Vis. Spectra, and melting point measurement .The bond constant (K) for active group that characterized by IR spectrum was calculated .Also ,solvent effect on UV.-Vis. Spectrum was studied which appeared the prepared solution compounds were influenced by solvent polarity .

Photo study for compounds was carry out by used Cs-137 as Gamma Ray source for (24 hour) . IR spectra and UV.-Vis. Spectra after irradiation didn't appear clear bands for compounds spectra that refer all compounds were gotten a photo-degradation because of they absorbed Gamma Ray. The mean external exposure [X_(R)], the dose rate in air [D_{aire} (rad)],and the dose rate in compound [D_{compound} (rad)] were calculated .

الخلاصة

أجريت الدراسة الضوئية للمركبات المحضرة وذلك بتعريضها لأشعة كاما لمدة (24 ساعة) بأستخدام السيزيوم-137 كمصدر مشع ، بعدها تمت مقارنة أطياف الأشعة تحت الحمراء وأطياف الأشعة الفوق البنفسجية –المرئية للمركبات بعد التشعيع بالأطياف قبل التشعيع حيث لوحظ ان الأطياف بعد التشعيع لم تظهر حزم واضحة لأي من المركبات المدروسة وذلك يدل على أن جميع المركبات حصل لها تحلل ضوئي ناتج عن أمتصاصها لأشعة كاما. كذلك حسبت قيم الجرع الممتصةمن قبل المركبات(D_{compound}) والجرع الممتصة في الهواء (D_{air}) ومعدل التعرض الخارجي (X_(R)) لأشعة كاما.

Introduction:

Schiff base of 4-aminoantipyrine and its complexes have a variety of applications in biological, clinical, analytical and pharmacological areas. Studies of a new kind of chemotherapeutic Schiff bases are now attracting the attention of biochemists. Earlier work reported that some drugs showed increased activity, when administered as metal complexes rather than as organic compounds^[1,2,3,4,5].

Gamma - Ray irradiation doses do not require catalysts, so there are no catalyst residues in the final product to interfere with physical properties. There is no heat treatment to degrade thermally sensitive components, and the dosage of irradiation can be controlled easily.

The main chains may be degraded or cross linked by the gamma -ray irradiation, with both processes usually taking place concurrently^[6].

Two radioactive isotopes of cesium ¹³⁷Cs and ¹³⁴Cs came 1986 from Chernobyl in Ukraine and deposited in many countries. These nuclides have the half - lives 30.0 a and 2.06 a, respectively. These isotopes have been used in adsorption, suspension and sedimentation research. Because ¹³⁴Cs was not with in depositions before 1986, it has met the particles, with which it has been detected, after the end of April in the year 1986 ^[7]. The measured ratio ¹³⁴Cs/¹³⁷Cs (into the 1986 level corrected) was $57(\pm 2)$ % (Aaltonen, et al., 1990) or 56 % (De Cort, et al., 1998) ^[8,9]. If the ratio is smaller, then a part of ¹³⁷Cs isotopes has deposited before 1986. By this way Froelich and Walling

(1994) found the age information for sediment and soil layers and Rafferty et al. (2000) for soil layers^[10]. The activity of ¹³⁴Cs may be measured directly today^[11,12,13,14,15]. However, even relatively small cesium -134 amounts are spectrometrically detected by using a coincidence method (Lee and Chung, 1991, Chung et al., 1992)^[16].

Today, Ge –detectors with high purity Ge –crystals of 2 kg are readily available and detectors with crystals up to 5 kg can be produced. Use of large area crystals results in high detection efficiencies, which can serve to reduce detection limits or to make faster measurements of low activity samples possible. The second major development that has contributed to lowering detection limits has been research on identifying and reducing the background components in gamma-ray spectrometry systems^[17].

In this research, photo stability for 4-aminoantipyrine derivatives were studied, because these compounds are important in medicine manufacture and the field of dyes. These compounds irradiated by gamma ray as type of cosmic ray. For benefit from the results to limitation the keep condition of these materials ,also we can to increase the photo stability by product metal complexes from these compounds.

Experimental:

All of the materials and solvents in table 1 were used as received .

Chemical material	Purity %	Provided Company
4-aminoantipyrine	95	Fluka A.G.
Benzylaldehyde	98	B.D.H.
Salicylaldehyde	98	B.D.H.
P-Hydroxy benzylaldehyde	98	Fluka A.G.
Ethanol	99	B.D.H.
Methanol	99	B.D.H.
Benzene	97	B.D.H.
Cyclohexane	99	B.D.H.
CCl ₄	95	B.D.H.

Table(1) : Chemical Materials were used

IR spectra were recorded as KBr discs using Testscan Shimadzu model 8400 FT-IR spectrophotometer in range (4000-400) cm⁻¹. Electronic spectra of the prepared compounds were measured in the region (200-800)nm for 10^{-3} M solution in ethanol using Shimadzu 160 spectrophotometer, with 1.000 ± 0.001 cm matched quartz cell. Gallen kamp capillary melting point apparatus was used to measure the melting points of the compounds.

Preparation of (A1):

A solution of 4-aminoantipyrine (0.89 g , 4.4 mmole) in ethanol (20ml) was added to benzylaldehyde (0.466 g , 4.4 mmole) dissolved in ethanol (15ml) adding anhydrous potassium carbonate . The mixture was allowed to reflux with stirred for three hours. The potassium carbonate was filtered off from the reaction mixture and the solvent was evaporated. The yellow solid separated was filtered and recrystallized from ethanol/H₂O.(see Fig. 1.)

Preparation of (A2&A3) :

The preparation of these compounds a similar way which that showed at the (A1) compound preparation ,but the Salicylaldehyde and P-Hydroxy benzylaldehyde used respectively , the physical properties summarized in table 2.(see Fig. 2&3).

Irradiation for Compounds:

Irradiation for compounds was carry out in nuclear physics lab at physics department /college of science/Kerbala university, by used Cs-137 as irradiation source .

The sample was put in photocell after that the photocell was put about 10cm far from the Cs-137 source and the irradiation reaction was left for 24h.

After that the sample was characterized by used FTIR and UV./Vis. Spectra.

Irradiation reaction for (A1) compound was carry out at 4/5/2009, (A2) compound at 5/5/2009, and (A3) compound at 6/5/2009.

Results and Discussion:

All compounds were characterized (Before & After) irradiation by used FT-IR $\,$, UV-Vis spectroscopy , melting point and solubility.

IR Spectra:

The IR spectrum for (A1) before irradiation, Fig.4 appears the bands at (1122,3043,2928 cm-1) ascribed to v(C-N) ,aromatic v (C-H) and aliphatic v (C-H) stretching respectively. The intense strong band at (1638cm-1) assigned to v(C=N) stretching. The other bands were listed in table 3.

The IR spectrum for (A2) before irradiation , Fig. 5 appears the band at (3454 cm⁻¹) can be attributed to v(O-H) stretching, the v(C=N) stretching vibration have been showed at (1610 cm⁻¹), the weak band at (3068 cm⁻¹) assigned to v(C-H) of aromatic frequencies and appears the band at (1445cm-1) ascribed to δ (C-H) aliphatic frequency .The other bands were listed in table 3.

While the IR spectrum for (A3) before irradiation, Fig. 6 appears the band at (3464 cm-1) assignable to intramolecular hydrogen bonded –OH group . The spectrum of the compound shows v(C=N) band in the region (1625cm-1). The other bands were listed in table 3.

After irradiation, the IR spectra of all compounds haven't been showed any clear band, figures (7,8&9). That may be return the bonds destroyed during irradiation of each compound.

Calculation of Bond Constant [K]:

Bond constant of each active group appears in IR spectrum was calculated by using equation $1^{(18)}$:

 $v = (1/2\pi) (k/\mu)^{0.5}$ (1)

v : vibration frequency

- k : bond constant
- μ : reduced mass

The reults of (k) data were listed in table 4.

UV-Vis spectra:

The electronic spectrum of (A1) before irradiation, Fig. 10 exhibits the peaks at (306 nm) (32679 cm⁻¹) (ϵ_{max} =4000 molar⁻¹.cm⁻¹) and (346nm) (2890 cm⁻¹) (ϵ_{max} =4000 molar⁻¹.cm⁻¹) assigned to ($\pi \rightarrow \pi^*$) and ($n \rightarrow \pi^*$) transitions respectively ⁽¹⁹⁾. Table 5.

The electronic spectrum of (A2) before irradiation, Fig. 11 displays tow peaks (307nm) (32573 cm⁻¹) (ϵ_{max} =4000 molar⁻¹. cm⁻¹) and (336nm) (29761 cm⁻¹) (ϵ_{max} =3783 molar⁻¹. cm⁻¹) attributed to ($\pi \rightarrow \pi^*$) and ($n \rightarrow \pi^*$) transitions respectively ⁽¹⁹⁾. Table 5.

Figure 12, showed the electronic spectrum of (A3) before irradiation. The two peaks at (251nm) (39840 cm⁻¹) ($\varepsilon_{max} = 4000 \text{ molar}^{-1}$.cm⁻¹) and (377nm) (26525 cm⁻¹) ($\varepsilon_{max} = 4000 \text{ molar}^{-1}$.cm⁻¹) .these peaks were assigned to ($\pi \rightarrow \pi^*$) and ($n \rightarrow \pi^*$) transitions respectively ⁽¹⁹⁾. Table 5.

After irradiation, the electronic spectra of all compounds were disappeared because these compounds destroyed .Fig.(13,14&15).

Solvent effect on UV-Vis. Spectra:

The solvent effect on UV-Vis. Spectra of all compounds were studied by using different solvents polarity. This study appeared the following results:

Compound (A1):

- 1- Red shift of $(n \rightarrow \pi^*)$ transition at (332,346,353,364 nm) by using CCl4,ethanol,methanol and benzene respectively.
- **2-** Blue shift of $(n \rightarrow \pi^*)$ transition at (353,346,332,321 nm) by using methanol, ethanol, CCl4 and cyclohexane respectively . Fig.(10,16,17,18&19).

The electronic spectra of (A2 & A3) also effected with polarity as same as compound (A1). See **Fig.(11,20,21,22,23,12,24,25,26&27)**.

Effect of Irradiation on Compounds

The study of irradiation for these compounds appeared that all compounds were degradation by Gamma Ray.

The mean external exposure [$X_{(R)}$], the dose rate in air [D_{aire} (rad)],and the dose rate in compound [$D_{compound}$ (rad)] were calculated by following equations ⁽²⁰⁾:

$$D_{air} (rad) = 0.87 * X_{(R)} \dots (2)$$

$$D_{\text{compound}}$$
 (rad) = 0.87 * [$\mu_{\text{mm}}/\mu_{\text{ma}}$] * $X_{(R)}$ (3)

 μ_{ma} : the mass absorption coefficient for Cs-137 in air.(1.79x10⁻² m²/Kg) μ_{mm} : the mass absorption coefficient for compound .

 $\mu_{\rm mm}({\rm m}^2/{\rm Kg}) = \Sigma \ {\rm w}_i \ \mu_i \ \ldots \ (4)$

 w_i : weight fraction of element in the compound.

 μ_i : mass absorption coefficient of element.

$$X_{(R)} = \Gamma * [A/d^2]....(5)$$

 Γ : exposure constant.(0.333 R.m²/h.ci).

A : radio active source at experimented time.

 A_0 : radio active source at industrial time. (100 µci).

 λ : decay constant = [ln2/T_{1/2}]. T_{1/2}:half time.(30 year).

 t_1 : experimental time .(4,5,and 6/5/2009 respectively).

 t_0 :industrial time. (1/12/2006).

d : the distance between the sample and Cs-137 source.(10cm).

The results of The mean external exposure [$X_{(R)}$], the dose rate in air [D_{aire} (rad)], and the dose rate in compound [$D_{compound}$ (rad)] data were listed in table 6.

Table (2): Physical properties of compound prepared

Comp.	Formula	Color	m.p .	Yield		Solubility				
				%	H ₂ O	Et- OH	Meth- OH	Benzene	Cyclo hexane	CCl ₄
A1	C ₁₈ H ₁₇ N ₃ O	Yellow- Crystal	216	75	X	+	+	±	±	±
A2	C ₁₈ H ₁₇ N ₃ O ₂	Brown- Powder	248	89	X	+	+	±	±	±
A3	C ₁₈ H ₁₇ N ₃ O ₂	Yellow- Needles	262	65	X	+	+	±	±	±

Soluble = +, Insoluble = x, Slightly soluble = \pm

Table (3): IR spectral data(wave number)cm-1 of compounds [A1],[A2] and[A3]

Comp	v(C-H) aromatic	(C-H) aliphatic	v(О-H)	v(C=C) aromatic	v(C=N)	Additional bands
A1	3043	v (2928)		1564	1638	1446w δ(C-N) 1720 ν(C=O) 1564s,1047m,838s(Ring vibration)
A2	3068	δ (1445)	3454	1600	1610	1440m δ(C-N) 1718 v(C=O) 1508s,1030m,831s(Ring vibration) 1147s δ(C-O) 1369m (O-H)in plane 761s (O-H)out plane
A3	3047	δ (1490)	3464	1590	1625	1485w δ(C-N) 1748 v(C=O) 1578s,1040m,831s(Ring vibration) 1130s δ(C-O) 1400m (O-H)in plane 758s (O-H)out plane

Table (4): Bond Co	onstant data(dyne.cm ⁻¹)	of active groups

Comp.	K(C-H)	K(C-H)	K(C=C)	K(C=N)	K(C-O)	K(C-N)	K(O-H)
	aromatic	aliphatic					
A1	5.04x10 ⁵	4.67×10^5	8.66x10 ⁵	10.22×10^5		7.97x10 ⁵	
A2	5.12×10^5	1.13×10^{5}	9.06x10 ⁵	9.88x10 ⁵	5.32x10 ⁵	7.90x10 ⁵	6.62x10 ⁵
A3	5.05x10 ⁵	1.21×10^{5}	8.95x10 ⁵	10.06x10 ⁵	5.16x10 ⁵	8.40x10 ⁵	6.66x10 ⁵

Table (5). CV-Visible data of Compounds								
Comp.	λ (nm)	Wave number (cm ⁻¹)	ϵ_{max} . (molar ⁻¹ . cm ⁻¹)	Assignments				
		· · · ·	· · · · · ·					
A1	306	32679	4000	$\pi \rightarrow \pi^*$				
	346	2890	4000	$\pi \rightarrow \pi^*$ n $\rightarrow \pi^*$				
A2	307	32573	4000	$\pi \rightarrow \pi^*$				
	336	29761	3783	$\pi \rightarrow \pi^*$ n $\rightarrow \pi^*$				
A3	251	39840	4000	$\pi \rightarrow \pi^*$ n $\rightarrow \pi^*$				
	377	26525	4000	n→π*				

Table (5): UV-Visible data of Compounds

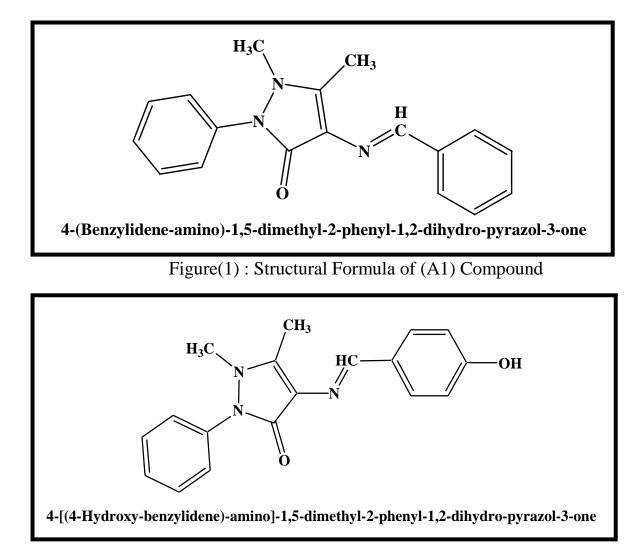
	Table (0). Data of absorption doses for Gamma Ray							
Comp.	Α	X _(R)	D _{air}	D _{compound}				
	(µci)	$(R.h^{-1})$	(rad)	(rad)				
A1	94.552	31.486x10 ⁻⁴	27.392x10 ⁻⁴	25.189x10 ⁻³				
A2	94.546	31.483x10 ⁻⁴	27.390x10 ⁻⁴	25.630x10 ⁻³				
A3	94.540	31.481x10 ⁻⁴	27.388x10 ⁻⁴	25.628x10 ⁻³				

Table (6): Data of absorption doses for Gamma Ray

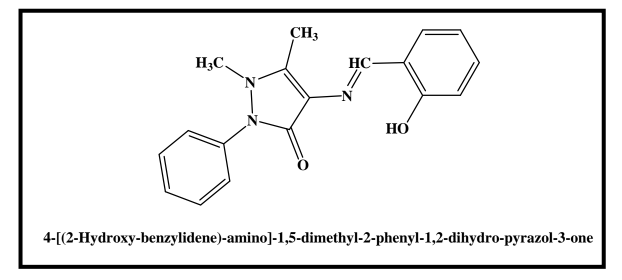
References:

- I- Chandra, S.; Jain, D.; Sharma, A. K. and Sharma, P. . "Coordination Modes of a Schiff Base Pentadentate Derivative of 4-Aminoantipyrine with Cobalt(II), Nickel(II) and Copper(II) Metal Ions: Synthesis, Spectroscopic and Antimicrobial Studies ". J. Molecules 14, 174-190 (2009).
- 2- Raman, N.; Thalamuthu, S.; Dhaveethuraja, J.; Neelakandan , M.A. and Sharmila Banerjee ." DNA Cleavage and Antimicrobial Activity Studies on Transition Metal (II) Complexes of 4-Amino -antipyrine Derivative ". J. Chil. Chem. Soc, 53 (1) (2008).
- 3- Raman, N. ; Dhaveethuraja, J. and Sakthivel, A. . " Synthesis, spectral characterization of Schiff base transition metal complexes: DNA cleavage and antimicrobial activity studies " . J. Chem. Sci., 119(4): 303–310 (2007).
- 4- Sadasivan, V. and Alaudeen, M. . " Synthesis and Crystal Structure of the Zinc(II) complex of 5-(2,3-dimethyl-1-phenyl -3-pyrazolin-5-one-4-ylhydrazono) hexahydropyrimidine -2-thioxo- 4,5,6-trione ". *Indian J.Chem.*,46A:1959-1962 (2007).
- 5- Vaghasiya, Y. K. ; Nair, R. ; Soni, M. ; Baluja, S. and Chanada, S. . " Synthesis, structural determination and antibacterial activity of
- compounds derived from vanillin and 4-aminoantipyrine " . J. Serb. Chem. Soc. 69 (12) :991-998 (2004).
- 6- Lee, I. J.; Choi, H. W.; Nho, Y. C. and Suh, D. H. . "Gamma-Ray Irradiation Effect of Polyethylene on Dimaleimides as a Class of New Multifunctional Monomers". J.Appl. Polymer Sci., Vol. 88: 2339–2345 (2003).
- 7- Saxén, R. and Ilus E. . "Discharge of ¹³⁷Cs and ⁹⁰Sr by Finnish rivers to the Baltic Sea in 1986-1996". J. Environ. Radio. 54: 275-291(2003).
- 8- Aaltonen, H.; Saxén, R. and Ikäheimonen, T. K. . "Airborne and deposited radioactivity. in Finland in 1987". *Report STUK A75* (1990).
- 9- De Cort, M.; Dubois, G.; Fridman, Sh. D.; Germenchuk, M.G.; Izrael, Yu. A.; Janssens, A.; Jones, A. R.; Kelly, G. N.; Kvasnikova, E. V.; Matveenko, I. I.; Nazarov, I. M.; Pokumeiko, Yu. M.; Sitak, V. A.; Stukin, E. D.; Tabachny, L. Y.; Tsaturov, Y. S. and Avdyushin, S.I. ." Atlas of Caesium Deposition on Europe after the Chernobyl Accident ". *EUR report nr.* 16733, EC,Of&Mac222;ce for Of & Mac222; cial Publications of the EuropeanCommunities, Luxembourg (1998).

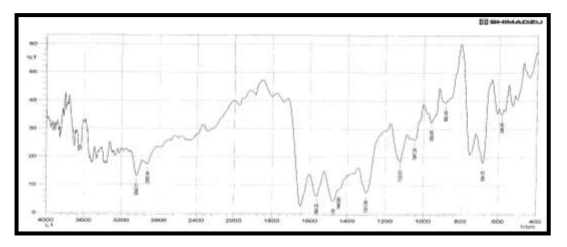
- 10- Froelich, W. and Walling, D.E. . " Use of Chernobyl derived radiocaesium to investigate contemporary overbank sedimentation on the flood plains of Carpathian rivers . Variability in Stream Erosion and Sediment Transport (Proceedings of the Canberra Symposium, Dac.1994) ". *IAHS Publ.* (224): 161-169 (1994).
- 11- Kasi, S. S. H. . " Mathematics for ground magazines in catchment modeling . XXI Nordic Hydrological Conference, Uppsala 26 -30.6.2000, ed. T. Nilsson ". *NHP Report Vol. 2(46): 355-358* (2000).
- 12- Kasi, S. S. H. . "Soil water content more accurately by nuclear methods. XXI Nordic Hydrological Conference, Uppsala 26-30.6.2000, ed. T. Nilsson ". NHP Report Vol. 2(46): 549-554 (2000).
- 13- Kasi, S. S. H. . " Cesium deposition in soil and its effects . Proceedings of the 8th International Symposium on Radiation Physics. Radiat " . *Phys. Chem. 61: 673-675* (2001).
- 14- Rafferty, B.; Brennan, M.; Dawson, D. and Dowding, D. . "Mechanisms of ¹³⁷Cs migration in coniferous forest soils". *J. Environ. Radio.48: 131-143* (2000).
- 15- Kasi, S. S. H. . " Cesium -137 deposition in Europe and the Large River Basins of Europe ". http://www.helsinki.fi/people/servo.kasi/CsEurLRiverB.zip, a *.zip packed *.cdr Corel-7 map (2002).
- 16-Lee, C. J. and Chung, C. . " Determination of ¹³⁴Cs in environmental samples using a coincidence gamma ray spectrometer". *Appl. Radiat. Isot.* 42, 783-788 (1991).
- 17- Hult, M. . " Low level gamma ray spectrometry using Ge-detectors ". *Metrlogia J.* 44 (4) (2007).
- Douglas, A. Skoog and Donald, M. West . "Fundamental of Analytical Chemistry 7th Edition ", U.S.A. (1997).
- 19- Lever, A.B.P. . "Inorganic Electronic Spectroscopy 2nd Edition", New York (1984).
- 20- Tsoulfanidis . "Measurement and Detection of Radiation 2nd Edition ",McGraw-Hill series in nuclear engineering, New York (1983).



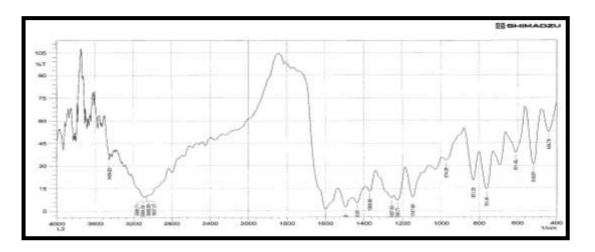
Figure(2) : Structural Formula of (A2) Compound



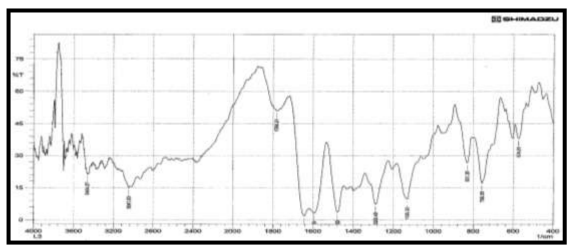
Figure(3) : Structural Formula of (A3) Compound



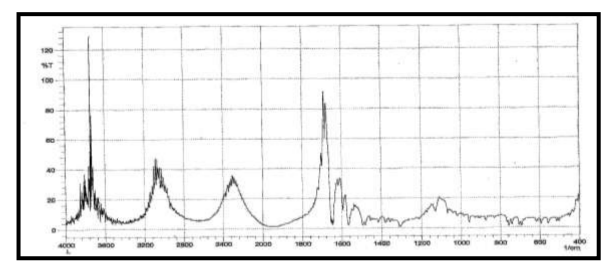
Figure(4):FT-IR Spectrum of (A1)Compound before irradiation



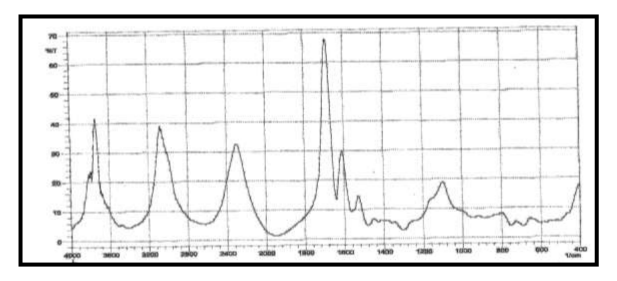
Figure(5):FT-IR Spectrum of (A2) Compound before irradiation



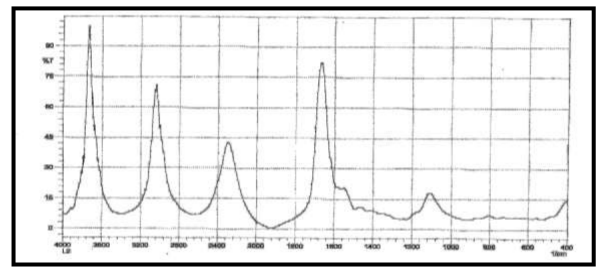
Figure(6):FT-IR Spectrum of (A3) Compound before irradiation



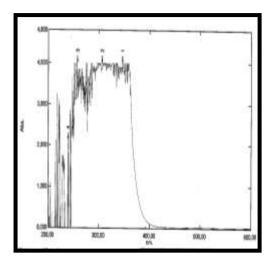
Figure(7):FT-IR Spectrum of(A1)Compound after irradiation



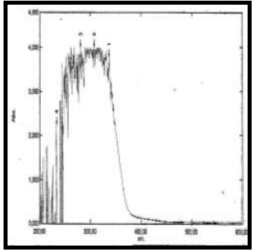
Figure(8):FT-IR Spectrum of (A2)Compound after irradiation



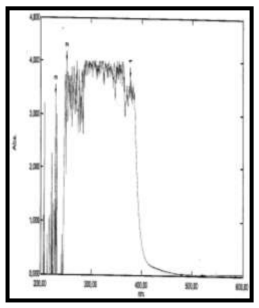
Figure(9):FT-IR Spectrum of(A3) Compound after irradiation



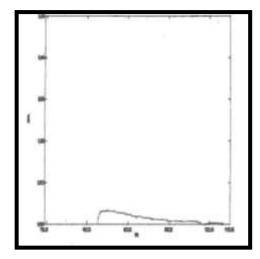
Figure(10):UV-Vis.Spectrum of(A1) Before irradiation



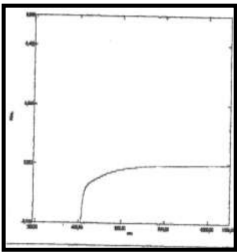
Figure(11):UV-Vis.Spectrum of(A2) Before irradiation



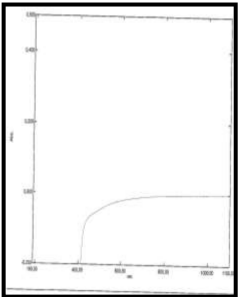
Figure(12):UV-Vis.Spectrum of(A3) Before irradiation



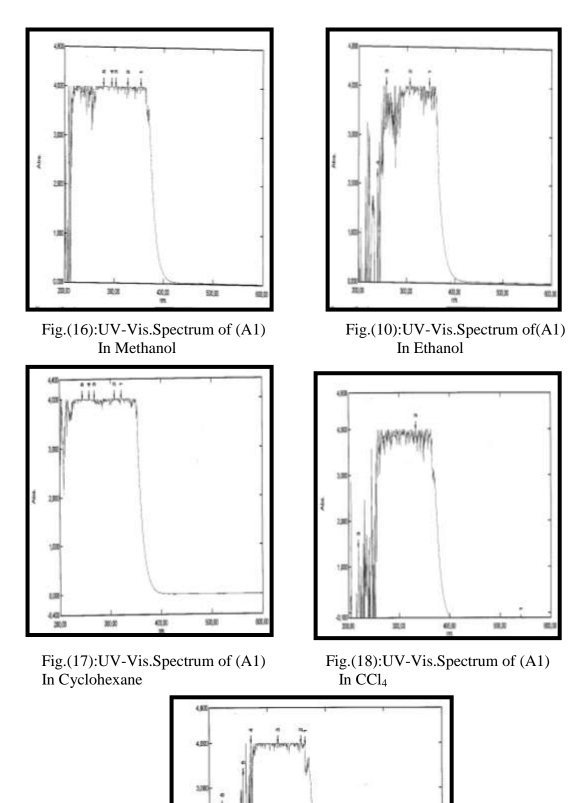
Figure(13):UV-Vis.Spectrum of(A1) After irradiation



Figure(14):UV-Vis.Spectrum of(A2) After irradiation



Figure(15):UV-Vis.Spectrum of(A3) After irradiation



152

400,00

Fig.(19):UV-Vis.Spectrum of(A1) In Benzene

300,08

900.00

531.0

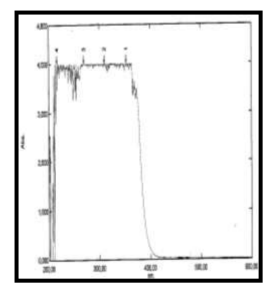


Fig.(20):UV-Vis.Spectrum of (A2) In Methanol

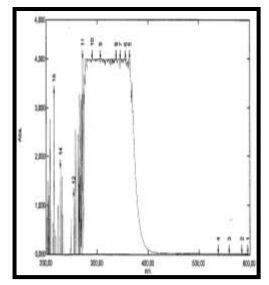


Fig.(22):UV-Vis.Spectrum of(A2) In Cyclohexane

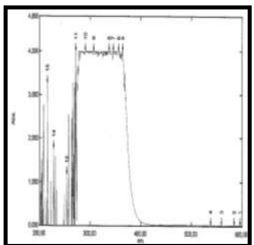


Fig.(23):UV-Vis.Spectrum of(A2)In Benzene

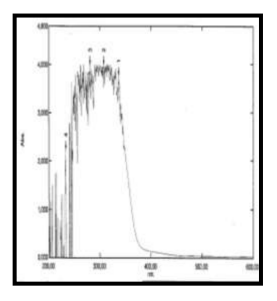


Fig.(11):UV-Vis.Spectrum of(A2) In Ethanol

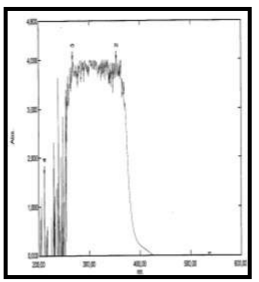


Fig.(21):UV-Vis.Spectrum of (A2) In CCl₄

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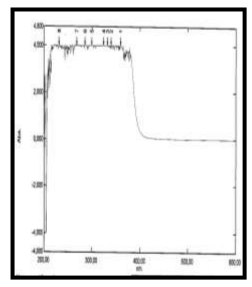


Fig.(24):UV-Vis.Spectrum of (A3) In Methanol

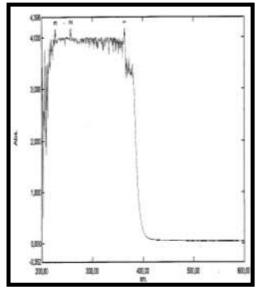


Fig.(25):UV-Vis.Spectrum of (A3) In Cyclohexane

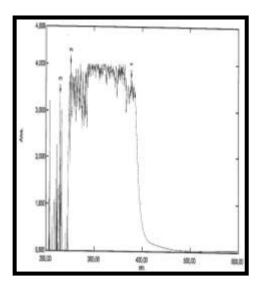


Fig.(12):UV-Vis.Spectrum of(A3) In Ethanol

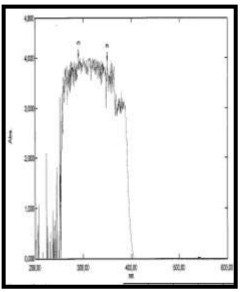


Fig.(26):UV-Vis.Spectrum of(A3) In CCl₄

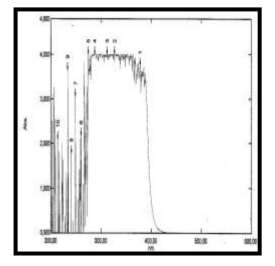


Fig.(27):UV-Vis.Spectrum of(A3) In Benzene