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Synthesis, Characterization, Thermal Studies, and Antioxidant Activities of Azo Dye[2-[(3-Hydroxyphenyl)diazinyl]-1,2-Benzothiazol-3(2H)-one-1,1- Dioxide]and Metal Ion Complexes

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Abstract

 The new azo dye was synthesized *via* the reaction of the diazonium salt form of 3 aminophenol with 2-hydroxyquinoline. This dye was then used to access a series of complexes with the chlorides of manganese, iron, zinc, cadmium, and vanadium sulfate. The prepared ligand and its complexes were characterized by FT-IR spectroscopy, UV-visible spectroscopy, mass spectrometry, thermogravimetric analysis, differential scanning calorimeter, and microelemental analysis. Conductivity, magnetic susceptibility, metal content, and chlorine content of the complexes were also measured. The ligand and cadmium complex were identified using¹H NMR and ¹³C NMR spectroscopy. The results showed that the shape of the ligand is a trigonal planner, and the complex shapes are tetrahedral, except for vanadium, which isa square-based pyramid. Additionally, the findings demonstrated that the complexes include water inside the coordination field and that each and every one of them is a non-electrolyte. The dye used the complexes prepared from it to determine their ability to inhibit free radicals by measuring their ability as antioxidants using DPPH as a free radical.D-ascorbic acid was employed as a standard substance in determining the value of IC_{50} , as it was found that the ligand had a high ability to inhibit free radicals. The ability to inhibit the complexes varied according to the value of IC₅₀, and the results are as follows: $H_2L > D$ -ascorbic acid $\geq Zn$ -complex \geq Fe $complex > V-complex > Cd-complex > Mn-complex.$

Keywords:3-Aminophenol, Antioxidant,Azodye complexes, 2-Hydroxyquinoline ,Thermodynamic parameter.

تحضير،تشخيص،د ارسة التحلل الح ارري وانشطة مضادات االكسدة لصبغة ازو)-2)-3 هايدروكسي فنيل(دايازنيل(-1،2-بنزوثايزول-3-اون -1،1ثنائي اوكسايد(مع معقدات ايونات الفلزات

فاطمة الزهراء شاكر حاتم و عباس علي صالح الحمداني

قسم الكيمياء, كلية العلوم للبنات, جامعة بغداد, بغداد, العراق

الخالصة

 حضرت صبغة االزو الجديدة من تفاعل ملح الديازونيوم من -3امينو فينول مع -2هايدروكسي كوينولين استخدمة الصبغة لتحضير سلسلة من المعقدات مع كلوريدات كل من المنغنيز والحديد والخارصين والكادميوم وكبريتات الفناديوم، شخص الليكاند والمعقدات المحضرة بطيف االشعة تحت الحم ارء وطيف االشعة فوق البنفسجة-المرئية وطيف الكتلة والتحاليل الحرارية الوزنية والمسعر التفاضلي والتحليل الدقيق للعناصر.وقياس التوصيلية والحساسية المغناطيسية ومحتوى الفلز ومحتوى الكلور للمعقدات. تم التحقق من الليكاند ومعقد الكادميوم بطيف الرنين النووي المغناطيسي للبروتون والكاربون. أظهرت النتائج ان الليكاند ثالثي السن والمعقدات رباعية السطوح عدا الفناديوم كان هرم مربع القاعدة. فضالً عن احتواء المعقدات على الماء داخل مجال التناسق وان جميعها غير الكتروليتية . استخدم الصبغة المعقدات المحضرة منها لتحدد قابليتها على كبح الجذور الحرة من خالل قياس قابليتها كمضادات اكسدة باستخدام مادة DPPH كجذر حر وحامض االسكوربك كمادة قياسية وتحديد قيمة 50IC حيث وجد ان الليكاند يمتلك قابلية عالية على كبح الجذور الحرة والمعقدات تفاوتت قابليتها على الكبح حسب قيمة 50IC وكانت النتائج كما يلي:

 $(H₂L > Ascorbic acid > Zn-complex > Fe-complex > V-complex > Cd-complex > Mn$ complex)

الكلمات المفتاحية: مضادات االكسدة, -3امينو فينول , معقدات اصباغ االزو, الثوابت الثرموديناميكية و-2 هيدروكسيكوينولين.

1.Introduction

 Azo dyes are among the most widely used, useful, and important types of chemical compounds, with diverse applications in science and technology[1-2]. The presence of an azo moiety (-N=N-) coupled with two monocyclic or polycyclic moieties identifies azo compounds[3]. Aromatic or heterogeneous systems are either unique or homologous[4]. Because of their distinctive physical and chemical characteristics and biological activities, they have a broad variety of uses in the pharmaceutical, cosmetic, food, dyeing, and textile industries [5]. However, their coloring job remains their most popular and favorite tool. The medicinal significance of azo compounds is widely recognized for their antibacterial, antifungal, and anti-HIV properties [6]. The majority of azo dyes are used to color textiles, and it is estimated that 10% of the dyes used in these dyeing procedures are not. They stick to the fibers and end up in wastewater treatment[7]. Some azo dyes are carcinogenic,andthis is due to degradation products such as benzidine,which activate various tumors in humans and animals. Aromatic amines found in many azo dyes affect human health, cause allergic reactions, and cause other diseases.Different metal quinoline complexes have been shown to be more effective than the parent quinoline because of their interesting bioactivity.Many studies have been performed on heterocyclic azo dyes and their metal chelators. Dyed azo metal chelates are of interest for use in molecular memory storage, nonlinear visualizations, and printing systems[8-10].This research aims to prepare novel complexes of the metal ions $(Cd^{+2}, Zn^{+2}, Fe^{+3}, Mn^{+2}, and V^{+4})$ using the azo ligand H2L. Thereafter, characterizationby spectroscopic analysis, thermal stability, and thermal decomposition will be studied using DSC and TGA curves, and the antioxidant activity of these compounds will be assessed against the DPPH radical and compared with D-ascorbic acidas a reference.

2.Experimental

2.1.*Material and Methods*

 All the materials used in this work were provided by SigmaAldrich, Merck, and other companies.The Urovector model EA/3000 singleV3O analyzer was used to record elemental microanalyses for carbon, hydrogen, nitrogen, and sulfur (C.H.N.S.).Through a gravimetric approach, mineralions were identified as M-O. Using DMSO as a solvent, a conductometer W-T-W estimated molar conductivities $(1\times10^{-3} \text{ M})$ at 25 °C. Mass spectrometry data were recorded using a Q-P-50-A-D-I Analysis Shimadzu QP(E170Ev)-2010-Pluss spectrometer. The UV-1800 Shimadzu Spectrophotometer was used to record the UV-visible absorption. The¹H and ¹³CNMR spectra were measured using aBruker (400MHz) spectrometer. The Fourier Transform Infrared (FT-IR) spectra were recorded using an IR Prestige-21, and the instruments

employed were the Braker $4000-500$ cm⁻¹ and the Shimadzu $4000-200$ cm⁻¹. Metals were identified using a Shimadzu (F.A.A.) 680 G atomic clock. Magnetic properties were used with the balancing susceptibility model MSR-MKI. All earlier types of thermal analysis employed Perkin-Elmer Pyris Diamond DSC/TGA.

2.2.*Synthesis of azo dye ligand*: *[*2*-[(3-hydroxyphenyl)diazenyl]-1,2-benzothiazol-3(2H)-one 1,1-dioxide]*

 To a solution of 3-aminophenol(1g,0.008mmol) inHCl (2 mL, xx N),ethanol (15mL), and distilled water (10 mL)at 0 to 5 °C,a solution of NaNO₂(1 g, 14.49mmol) in H₂O (10 mL)was added gradually. The reaction mixture was stirred for 45 minutes before adding a solution of 2 hydroxyquinoline (1.16 g, 0.008 mmol) in ethanol (15 mL). A change to a dark-colored solution was observed after 30 minutes of stirring. The solid crude material formed wasfiltrated and dried to give a brown precipitate with an 87% yield that melts at 253-255 °C.

2.3.*General approach for metal complexes synthesis*

A solution of metal salt $[VOSO₄.5H₂O (0.1 g, 1 mmol), FeCl₃ (0.058g, 1 mmol),$ MnCl₂.4H₂O (0.08g,1 mmol), ZnCl₂(0.05g, 1 mmol), or CdCl₂(0.07g, 1 mmol)] in water (10 mL). Azo ligand $H₂L$ (0.1g, 1mmol) was added drop by drop to the solution before heating at 40 °C for 2 hours. By briefly submerging the solid complexes in hot ethanol, any unreached components were separated from them and eliminated. The complexes were gathered, dried, and weighed.

3. Results and discussion

3.1.*Physical and analytical data for the ligand(H2L) and the complexes synthesized*

 The reaction of metal salts and ligandafforded the desired complexes as shown in Scheme 1. The findings of the elemental analysis show that all compounds have a 1:1 ratio of M:L.The results of the microelemental analysis were consistent with the theoretical calculations, as shown in Table 1.

Scheme1- Formation for ligand (H₂L) and their metal complexes

Compound no.	Formula Molecular weight	m.p. (°C)	Ratio	\mathcal{C} (%)	ο- H (%)	N (%)	M $(\%)$	Cl (%)	Conductivity In DMSO $\text{cm}^2 \Omega^{-1} \text{mol}^{-1}$
$\mathbf{1}$	$C_{15}H_{11}N_3O_2$ 265.27	253-255	found	67.92	4.18	15.84			
			Calculated	67.09	3.72	16.81			
$\overline{2}$	$C_{15}H_{11}N_3O_4V$ 348.21	289-291	found	51.74	3.18	12.07	14.63		12
			Calculated	50.62	3.83	12.28	14.88		
3	$C_{15}H_{11}N_3O_3Mn$ 336.20	276-278	found	53.59	3.35	12.50	16.34		16
			Calculated	52.69	3.09	13.16	13.02		
$\overline{4}$	$C_{15}H_{13}N_3O_4C$ IFe 390.58	292-294	found	46.13	3.35	10.76	14.30	9.08	19
			Calculated	46.89	3.97	11.89	13.54	9.57	
5	$C_{15}H_{11}N_3O_3Zn$ 346.66	277-280	found	51.97	3.20	12.12	18.86	-	11
			Calculated	50.99	4.11	13.10	18.55		
6	$C_{15}H_{11}N_3O_3Cd$ 393.68		found	45.76	2.82	10.67	28.55		10
		296-299	Calculated	44.89	3.55	11.89	28.22		

Table 1: Some elemental physical characteristics investigations of ligand and complexes

3.2. *¹H NMR and ¹³C NMR data of theazo ligandand Cd complex*

The ¹HNMR and ¹³CNMR spectra of azo ligand are displayed in Figure 1. The ¹HNMR data in DMSO-*d*⁶ (ppm) are 2.45 (2H, s,N-H),7.26(1H, s,Ar-H), 7.72-7.71(1H, d, Ar-H), 11.68- 11.66(1H, s, 2OH)[11]. ¹³CNMR data in DMSO-*d*⁶ (ppm) are 127.5 (C1) 149.0 (C2),118.2 (C3) 145.0 (C4), 157.2 (C5), 169.8 (C6), 132.2 (C7) 165.3 (C8), 137.3 (C9), 166.9 (C10) I55.2 (C11), 122.2 (C12), 178.1 (C13), 182.0 (C14), 189.75 (C15)[12].

The 1 H NMR and 13 C NMR spectra of Cd-complex are displayed in Figure 2. The 1 H NMR data of Cd complex in DMSO- d_6 (ppm) are 3.34 (DHO impurities), 8.46-7.36 (9H, m, Ar-H).¹³C NMR data in DMSO-*d*⁶ (ppm) are106.6 (C1), 189.8 (C2), 145.5 (C3) 181.7 (C4), 190.2 (C5), 118.6 (C6) 137.6 (C7), 127.8 (C8), 132.5 (C9),155.4 (C10), 172.6 (C11), 158.0 (C12),178.7 (C13), 165.8 (C14), 196.2 (C15)[11].

Figure1 : ¹H NMR and ¹³C NMR spectra of azo ligand (H₂L)

Figure 2: ¹H NMR and ¹³C NMR spectra of the Cdcomplex

3.3*. LC-mass spectra for ligand and its complexes*

One of the most crucial methods for characterizing the ligand (H_2L) and some products is LC-Mass spectrum testing. This method is supplementary to the other methods that estimate the molecular weight of the chemical using the relationship $(m/z)[13]$. The fragmentation pattern and the extract mass for each pattern are shown in Scheme 2. The fragment's molecular ion peak $[M]^+$ of C₁₅H₁₀N₃O⁺ is easily visible, and its relative abundance is about 67% in Figure 3. Other peaks observed at 45, 32, and 88% belong to the $C_{15}H_{10}N_3^+$, $C_9H_6N^+$, and $C_6H_5N_2^+$ ions, respectively[8]. For the $\text{[Cd}(L)H_2O$] (Figure 4 and Scheme 2), we can observe the molecular ion peak (M^+) at 392.33m/z with a relative abundance of 10%, and the next patterns are $C_{15}H_7CdN_3O_2^{++}$, $C_{15}H_8CdN_3O^{++}$, $C_{15}H_8CdN_2O^{++}$, $C_9H_4CdN_2O^{++}$ and $C_6H_5^+$, which correspond to 374.58 m/z,358.78 m/z, 344.64 m/z,267.69m/z, and77.84m/z,respectively .Inthe $[Mn(L)H_2O]$ complex (Scheme 2), the fragments (M^+) at 336.42m/z with a relative abundance of 10% , and the next patterns are $C_{15}H_8N_3O_2Mn^+$, $C_9H_4N_3OMn^+$, $C_9H_4N_2OMn^+$, $C_9H_4NOMn^+$ and $C_6H_4O^{+}$, whichcorresponded to 317.42m/z, 255.02 m/z, 211.35m/z , 197.42m/z , and 92.33m/z . For [Fe(L)(H₂O)₂Cl] complex(Scheme5), the fragments (M⁺) at 390.44m/z with a relative abundance of 15% and the next patterns are $C_{15}H_9CIFeN_3O_2^+$, $C_{15}H_9FeN_3O_2^+$, $C_{15}H_9FeN_3O^+$, $C_9H_5FeNO^+$, $C_6H_5N_2^+$, $C_7H_5N_2^*$ ⁺ and C_2HFeO^+ corresponded to 354.64m/z,319.08 m/z, 303.32m/z, 198.14 m/z, 105.44 m/z,103.21m/z, and 96.89m/z. In the [VO(L)H₂O]complex (Scheme 2), the fragments(M^+) at 348.34m/z with relative abundance 20% and next pattern: $C_{15}H_8N_3O_3V^+$, $C_9H_4N_3O_2V^+$, $C_9H_4N_2O_2V^+$, $C_9H_4NO_2V$ and $C_6H_4O^+$, which corresponded to 329.22m/z, 237.01m/z, 223.22 m/z, 209.41m/z, and 92.36m/z.Forthe $[Zn(L)H_2O]$ complex in Scheme 2, the next fragments $(M+)$ are at 345.86m/z with a relative abundance of 25% and the next patterns areC₁₅H₈N₃O₂Zn⁺,C₁₅H₈ZnN₃O⁺,C₁₅H₈ZnN₂O⁺,C₉H₃ZnN₂O⁺and C₆H₅⁺which correspond to 327.63m/z, 311.65 m/z, 297.35m/z,and 220.54m/z,respectively[14] .

Figure 3: Mass spectrum of ligand H₂L

Exact Mass: 96.87

Scheme 2: Pattern of fragmentation of ligand and theircomplexes

3.4.*The ligand (H2L) and its complexes are studied by UV-Vis*

The electronic spectrum for ligand $(H₂L)$ in Figure 5 exhibits strong absorptions at 275 nm and 36363.64cm⁻¹ ascribed to the $\pi \rightarrow \pi^*$ transition and a peak at 350nm and 28571.43cm ¹ attributed to the n $\rightarrow \pi^*$ transition, a peak with a high intensity band formed withabsorption maxima[15]. The electronic transition of the V^{+4} complex depicts peaks of 280,345,390,620, and780 nm assigned to $\pi \rightarrow \pi^*$, $n \rightarrow \pi^*$, ${}^2B_2g \rightarrow {}^2E_g$, and ${}^2B_2g \rightarrow {}^2B_1g$, respectively, which is indicative of a square pyramidal geometry.The electronic spectrum of the divalent zinc compound was studied, and it was found that it does not give d-d transitions because it contains $(d¹⁰)$ in the valence shell. However, it gave two peaks, each belonging to the ligand spectrum: peaksat 290nm and 470 nm assigned to $\pi \rightarrow \pi^*$ and $n \rightarrow \pi^*$, respectively, which is indicative of a tetrahedral. The electronic spectrum of the Cd^{+2} compound was studied, and it was found that it does not give d-d transitions because it contains (d^{10}) in the valence shell, but it gave three peaks, each belonging to the ligand spectrum, peaks in 255, 340, and 390 nm assigned to $\pi \rightarrow \pi^*$, $n \rightarrow \pi^*$, and C.T. M \rightarrow L, respectively, which is indicative of a tetrahedral. The electronic absorption of Mn^{2+} complex exhibited peaks of 235, 295, 325, 595, and 785 nm ascribed to the $\pi \rightarrow \pi^*$, $n \rightarrow \pi^*$, C.TML, ${}^6A_1g \rightarrow {}^4Eg_{(G)}$, ${}^{6}A1g \rightarrow {}^{4}T_2g_{(G)}, {}^{6}A_1g \rightarrow {}^{4}T_1g_{(G)},$ and ${}^{6}A_1g \rightarrow {}^{4}T_2g_{(D)},$ respectively, which is indicative of a tetrahedral. The electronic transition of Fe^{+3} complex show in Figure 6 exhibited peaks of 270,310,545, and 720 nm assigned to $\pi \rightarrow \pi^*$, $n \rightarrow \pi^*$, C.T M $\rightarrow L$,

 ${}^{6}A_{1}g \rightarrow {}^{4}T_{2}g_{(G)}, {}^{6}A_{1}g \rightarrow {}^{4}T_{1}g_{(G)},$ and ${}^{6}A_{1}g \rightarrow {}^{4}Eg_{(D)},$ respectively, which is indicative of anoctahedral geometry.Table2 displays the electronic assignment of metal complexes[16,17].All complexes are non-electrolytes.

Compound	λ nm	\acute{v} cm ⁻¹	Abs	$\epsilon_{\rm max}$ L mol -1 cm^{-1}	Assignment	μ_{eff} (B.M) found (calculate)	Hybridizati on	Distributio $\mathbf n$
Ligand H_2L	275 350	36363.6 4 28571.4 3	1.89 2.13	1890 2130	$\pi\mathop{\rightarrow}\!\pi^*$ $n \rightarrow \pi^*$			
[VO(L)H ₂] O ₁ Square pyramidal	280 345 390 620 780	35714.2 9 28985.5 1 25641.0 3 16129.0 3 12820.5 1	1.20 $\boldsymbol{0}$ 1.12 $\boldsymbol{0}$ 0.40 0 0.26 $\mathbf{0}$ 0.25 $\boldsymbol{0}$	1200 1120 400 260 250	$\pi\mathop{\rightarrow}\!\pi^*$ $\mathbf{n} \longrightarrow \!\! \pi^*$ C.T ${}^{2}B_{2}g \rightarrow {}^{2}Eg$ ${}^2B_2g \rightarrow {}^2B_1g$	1.81 (1.73)	\rm{dsp}^3	$t_2g^1eg^0$
[Mn(L)H ₂] O ₁ Tetrahedra $\mathbf{1}$	235 295 325 595 785	42553.1 9 33898.3 1 30769.2 3 16806.7 2 12738.8 5	2.12 5 1.89 5 0.89 0 0.62 5 0.59 5	2125 1895 890 625 595	$\pi \rightarrow \pi^*$ $n \rightarrow \pi^*$ $C.TM\rightarrow L$ ${}^6A_1g \rightarrow {}^4Eg$ (G \mathcal{L} ${}^6A_1g \rightarrow {}^4T_2g$ G)	5.44 (5.91)	SP ³	$e^2t_2^3$
[Fe(L)H ₂ O] Cl]Octahe dral	270 310 545 720	37037.0 4 32258.0 6 18348.6 $\overline{2}$ 13888.8 9	2.11 5 1.99 8 0.25 $\boldsymbol{0}$ 0.12 5	2115 1998 250 125	$\pi\mathop{\rightarrow}\!\pi^*$ $n \rightarrow \pi^*$ ${}^6A_1g \rightarrow {}^4Eg(G)$ \mathcal{L} 6 A1g \rightarrow ⁴ T ₂ g(G	5.17 (5.91)	SP ³ d ²	$T_2g^3eg^2$
$[Zn(L)H_2O]$ Tetrahedra 1	290 470	34482.7 6 2127.66	2.31 5 0.52 5	2315 525	$\pi \rightarrow \pi^*$ $n \rightarrow \pi^*$	Diamagneti \mathbf{C} (0)	SP ³	$e^4t_2^6$
[Cd(L)H ₂] O ₁ Tetrahedra 1	255 340 390	39215.6 9 29411.7 6 25641.0 3	2.12 $\boldsymbol{0}$ 1.98 $\boldsymbol{0}$ 0.89 $\boldsymbol{0}$	2120 1980 890	$\pi \rightarrow \pi^*$ $n \rightarrow \pi^*$ $C.T M\rightarrow L$	Diamagneti \mathbf{C} (0)	SP ³	$e^4 t^6$ ₂

Table 2: The ligand (H₂L) and its complexes are studied by UV-Vis

Figure 5 UV-Vis spectrum of ligand (H2L) **Figure6**- UV-Vis spectrum of Fe-complex

3.*5*.*Thermal analysis*

 The results of the thermal analysis can be seen in Tables 4 and 5 and in Figures 7 and 8 for the ligand (H2L) and the complexes that were made [18]. Scheme 8 outlines the potential breakdown response of metal complexes. Based on the thermograms, computations of the decomposition phases, temperature ranges, decomposition products, and weight loss complex percentages indicated consistency. That verifies the elemental analysis findings and recommended equations between their thermal decomposition results and computed values[10].In this work, it was noted that in the ligand and metal complexes of V^{+4} , Fe^{+3} , Zn^{+2} , Cd^{+2} , and Mn^{+2} , the remaining ligand was carbon and the remaining metal oxide. Based on the results of the thermogravimetric studies, it can be shown that the complexes and the ligand form one to three distinct phases.Using the DCS curve, the thermodynamic parameters enthalpy ∆H, entropy ∆S and Gibbs free energy (∆G)were calculated[19].

Figure 8: TGA&DSC curve of MnComplex

Scheme 8: Tentative decomposition reaction of ligand and metal complexes

Compound	$T_i \underset{C}{\circ} C$	$T_{f\!/\,{}^\circ\!C}$	Maximum temperature point $(^{\circ}C)$	ΔH (J/g)	ΔS (J)	ΔG (J)	Type
H_2L	67.21	128.67	97.27	-89.78	-1.4608	52.31	endothermic
[Fe(L)(H ₂ O)Cl]	105.94 156.16 314.05	112.21 159.31 319.44	316.57	-1.19 -0.79 -2.78	-0.178 -0.251 -0.515	55.159 78.669 160.256	endothermic endothermic endothermic
$[Zn(L)H_2O]$	263.32 278.01	276.00 287.47	257.72 270.71 281.09	-36.90 -11.55	-2.910 -1.220	713.065 331.379	endothermic endothermic
$[Mn(L)H_2O]$	43.76 218.42	108.80 357.93	74.42 288.99	182.62 249.94	2.807 1.791	49.512 -267.641	endothermic exothermic
[Cd(L)H ₂ O]	387.65	395.13	392.40	-3.03	-0.405	155.892	endothermic
[VO(L)H ₂ O]	46.17 60.34 172.41 324.45	63.49 98.63 257.74 334.99	56.51 76.54 223.54 332.57	-1.63 -24.35 54.84 -1.75	-0.094 -0.635 0.642 -0.166	5.2179 24.252 -88.673 53.456	endothermic endothermic exothermic endothermic

Table3: Thermal decomposition DSC of Ligand and some complexes

3.6.Infrared spectra

 The infrared spectra of ligand H2L and its metal complexes were recorded with Cd^{+2} , Zn^{+2} , Fe^{+3} , Mn^{+2} , and V^{+4} , and the data has been organized in Table 5. The ligand showed bands at 3477, 3167, 1546-1600, and 1257cm⁻¹ that were ascribed to the stretching vibrations of O-H, C-H aromatic,C=C aromatic,and C-O [20].The infrared spectrum of the ligand showed a medium-intensity stretch band at frequency 1352cm⁻¹, which was attributed to the vibration frequencies of the doublebond N=N (Figure 9). The FT-IR spectra of all the prepared compounds showed that the azo-dye ligand was coupled to metal ions through two sites: the oxygen site *via* deprotonation of the phenolic group and the nitrogen site of the azo group[21]. New bands belonging to M-Nappeared at 611,617, 621, 615, and 619 cm⁻¹ for the V^{4+} , Mn²⁺, Cd^{+2} , Zn^{+2} , and Fe⁺³complexes, respectively.M-O at 457, 462, 460, and 450 cm⁻¹ for the complexes V^{4+} , Mn²⁺, Cd⁺², Zn⁺², and Fe⁺³, respectively. M-Cl at 385 cm⁻¹ for the Fe⁺³complex [22].

Compound $v(H2O)$ aqua		$v(C-H)$ $v(C-O)$ aromatic		$v(N=N)$	Other bands
H_2L		3167	1257	1352	
[VO(L)H ₂ O]	3410 1643 835	3169	1213	1419	$611(V-N)$ $457(V-O)$ $979(V=0)$
$[Mn(L)H_2O]$	3383 1641 837	3015	1217	1409	$462(Mn-O)$ $617(Mn-N)$
$[Zn(L)H_2O]$	3400 1640 834	3100	1220	1400	$615(Zn-N)$ $460(Zn-O)$
[Fe(L)(H ₂ O)Cl]	3385 1645 830	3050	1215	1415	619 (Fe-N) 450 (Fe-O) 385 (Fe-CI)
[Cd(L)H ₂ O]	3388 1642 828	3160	1218	1412	$621(Cd-N)$ $458(Cd-O)$

Table 5: The FT-IR spectra bands (cm⁻¹) of the ligand and its complexes

3.7.Investigation of antioxidant activity

The antioxidant activity of H_2L and its mineral complexes was measured by ascorbic acid, and the DPPH assay was used to scavenge free radicals.Initially, each sample was diluted with the same volume of methyl alcohol, and then it was mixed with the same volume of a concentrated 0.135 mM DPPH solution. After adding the DPPH solution[23], the samples were kept at room temperature in the dark for 30 minutes.The absorbance of each sample was then determined to be 517 nm. The lower the IC_{50} value of the complexes compared to ascorbic acid, the greater their ability to suppress free radicals.As the ligand was the most effective free radical inhibitor, followed by zinc, which has the same value as ascorbic acid, followed by iron, vanadium, cadmium, and manganese[24,25], the values of IC₅₀are as follows: $H_2L > D$ -ascorbic $acid > Zn$ -complex $\geq Fe$ -complex $\geq V$ -complex $\geq Cd$ -complex $\geq Mn$ -complex (Figure 11).

Figure 11: Variations of IC_{50} values for H_2L ligand and its complexes

Table 6: The antioxidant results (P1%, RSA% and IC_{50}) of H_2L and their metal complexes

4. Conclusion

 A novel azo ligand was prepared *via* the reaction ofthe diazonium salt of 3-aminophenol with 2-hydroxyquinoline. This ligand was then employed to access new complexes with different metals.These complexes were identified using a number of analytical techniques, such as elemental microanalysis, metal chloride-containing, electrical conductivity measurement, magnetic susceptibility, ${}^{1}H$ and ${}^{13}C$ -NMR, FT-IR, and UV-Vis spectroscopy. Calculations of the thermodynamic parameters ΔH , ΔS , and ΔG were made using the DCS curve, and the atomic N,O, and O tridentate coordination sites in the ligand were identified by comparing their FT-IR spectra to those of the metal complexes. The M:L ratio in every compound was 1:1. The dye used the complexes prepared from it to determine their ability to inhibit free radicals by measuring their ability as antioxidants using DPPH as a free radical and D-ascorbic acid as a standard substance and determining the value of IC_{50} . The ligand exhibited a significant capacity to suppress free radicals, and its ability to inhibit the complexes varied depending on the IC_{50} value. The results are as follows: $H_2L > D$ -ascorbic acid $\geq Zn$ -complex $\geq F$ e-complex $> V$ $complex > Cd$ -complex $> Mn$ -complex.

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