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PH-metric and Theoretical Studies of The Complexation of $2-[\alpha-(o-hydroxyphenyl)ethylidenehydrazino]-4,6-dimethylquinoline$ $and <math>2-[\alpha-(o-methoxyphenyl)methylidenehydrazino]-4,6-dimethylquinoline$

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 $2^{-[\alpha-(o-Hydroxyphenyl)ethylidenehydrazino]-4,6-dimethylquinoline (AHQ) and 2-[\alpha-(o-methoxyphenyl)methylidenehydrazino]-4,6-dimethylquinoline (BHQ) have been synthesized and characterization is performed by elemental analysis and electronic, vibrational and mass spectra. The pK^H and log K were gritty in 75% solvent-water pH-metrically for BHQ ligand and at various temperatures for AHQ ligand. The dissociation and stability constants in the aqueous medium for BHQ ligand have been calculated by the relation of pK^H or log K with solvent parameters (D, ET, AN and DN). The isokinetic temperature was determined by using a linear regression analysis (LRA) of <math>\Delta$ H° vs. Δ S° for AHQ complexes. The thermodynamic parameters of AHQ compounds were analyzed into their electrostatic (el) and non-electrostatic (non) or cratic components. Full geometrical and structural optimizations of the ligands have been performed by a *DFT* study by using the hyperchem program.

Keywords: Quinolinyl hydrazones, Stability constants, Theoretical studies, Solvent and thermodynamic parameters

Introduction

Hydrazones are excellent chelating agents, which attracted special interest as well as their metal complexes because of their important applications including antimicrobial, antitumor and other biological applications[1-9]. The quinolone derivatives and their metal complexes play essential role in many fields[10]. The heterocycle compounds that have N atoms can carry a positive charge and act as a hydrogen bond acceptor or donor[11]. Many drugs in the market are prepared from quinolone derivatives[12]. They have pharmacological activities including antimicrobial, antioxidant, toxicity assessment[12], antiplasmodial, antimalarial[13], antituberculosis^[14], anti-inflammatory^[15] and anticancer properties[16]. This work is extension of studies on quinolone hydrazones[17-24]. The aim of the pH-metric studies is calculation of the dissociation constants of the ligands and the stability constants of their metal complexes in

solution. However, these studies were performed under different experimental conditions *viz*.

Various mixed solvents, 75% (v/v) solvent-water; solvent = dioxane, isopropanol, ethanol and methanol) at 303K for BHQ ligand with some divalent 3d transition metal ions (Mn^{II} , Co^{II} , Ni^{II} and Cu^{II}), non transition 3d and 4d metal ions (Zn^{II} and Cd^{II}), trivalent 4f lanthanide metal ions (La^{III} , Ce^{III} , Sm^{III} and Ho^{III}) and 5f actinide metal ions (UO_{2}^{II}).

Various temperatures (283, 293, 303 and 313K) for AHQ ligand with Mn^{II} , Co^{II} , Ni^{II} , Cu^{II} and Zn^{II} -ions.

Experimental

Materials

p-Toluidine, ethyl acetoacetate, phosphorus oxychloride, hydrazine hydrate, *o*-hydroxyacetophenone, *o*-anizaldehyde, hexamine, indicators, salts and solvents used

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in this investigation were of the highest purity available (Merck, BDH, Aldrich and Fluka).

Preparation of the hydrazones

2-Hydrazino-4,6-dimethylquinoline (HQ)is prepared as described in our previous publication[17]. To an ethanolic solution of 2-hydrazino-4,6-dimethylquinoline (HQ) (0.01 mol), o-anizaldehyde or o-hydroxyacetophenone (0.012 mol) was added. The mixture was heated under reflux for 1/2 hour. After cooling, the formed yellow compounds were filtered off, washed with ether and the ligands are crystallized by ethanol. The results of elemental analyses, % yield, colour and m.p °C are shown in Table 1. The resultant hydrazones are; 2-[a-(o-hydroxyphenyl) ethylidenehydrazino]-4,6-dimethylquinoline 2-[α-(*o*-methoxyphenyl) (AHQ) and methylidenehydrazino]-4,6-dimethylquinoline (BHQ). The results of elemental analyses (Table 1) are in best agreement with the proposed formulae.

Measurements

Physical measurements

IR spectra (4000-400 cm⁻¹) were verified on a BRUKER Vector 22 spectrometer (Germany) using KBr pellets. UV-Visible spectra were elucidated on a Jasco V-550 UV/VIS spectrophotometer. Mass spectra were examined at 70 eV on a gas chromatographic GCMSqp 1000-ex Shimadzu mass spectrometer.

Potentiometric measurements

WTW-D-8120 digital pH-meter fitted with a combined glass electrode was used to measure pH-reading of the titration of standard solutions of metal nitrates (0.001M) with ligands (0.003M). Solutions were adjusted to 0.05M ionic strength by addition of KNO₃, the total volume was made up to 30 mL and maintained at a constant temperature by circulated water through a sealed-jacketed cell.

Results and Discussion

Characterization of the ligands

The structures of the ligands have been determined by elemental analyses, IR, mass and UV-Vis spectra.

IR & Mass spectra of the ligands

The IR spectra of the hydrazones (AHQ & BHQ) (Table 1) showed very strong bands at *ca.* 1616 - 1614 and 3440 – 3175 cm⁻¹ which are attributed to v(C=N) and v(NH), respectively[25-34]. On the other hand, the broad band centered at 2868 cm⁻¹ is assignable to v(OH--N) of the phenolic group of AHQ, ligand[35-37]. The mass spectrum of the BHQ ligand showed molecular ion and base peaks at m/z 305 and 172, respectively, which coincide with its formula weight.

Electronic spectra of the ligands

The electronic absorption spectra of the investigated ligands as well as the hydrazino compound; HQ (5x10⁻⁵ mol. dm⁻³) in dioxane involve three or two sets of bands in the region 239 -418 nm (Table 2). On the basis of the high molar absorptivity ϵ_{max} , the highest energy bands at λ_{1} (239 -247 nm) are assigned to π - π^* transitions of the quinoline ring[7], phenyl rings and/or azomethine groups. Also, the moderate energy bands at λ_{2} (344 -351 nm) may be composite bands due to charge transfer (CT) interactions [25-29]. Finally, the lower energy bands at λ_2 (329 - 418 nm) can be ascribed to the low intensity n - π^* forbidden transitions. The π - π * transition (band at 247 nm) of the non condensed HQ compound is more intense by $\approx 2 - 2$ 4 times than the corresponding hydrazones. Also, this band shows a red shift (Table 2). The etheric hydrazone BHQ has the highest λ_{max} and ε_{max} . Therefore, the weak n - π^* transition was hidden under the strong π - π^* transitions.

Molecular orbital calculations

The calculated structural parameters are

TABLE 1. Analytical, physical data and selected IR absorption bands (cm⁻¹) of the ligands.

Compound			Elem % F	ental Ana ound / (C	ılysis alcd)	m.p.ºC	IR spec	ctral bands	(cm ⁻¹)
(F.Wt) M.F.	Colour	% Yield	С	Н	Ν	-	v(C=N)	v(OH -N)	v(NH)
AHQ (305) C ₁₉ H ₁₉ N ₃ O	Yellowish- Orange	75	74.38 (74.75)	6.55 (6.20)	13.75 (13.77)	165	1616	2868	3440
BHQ (305) C ₁₉ H ₁₉ N ₃ O	Yellow	60	75.10 (74.75)	6.40 (6.20)	13.76 (13.77)	167	1614		3175

presented in Table 3. The following remarks can be pointed out:

- a) The dipole moment are higher for the ligands than that of HQ, which may be due to the rigid aromatic structure of BHQ and AHQ.
- b) The shortening of the (N-N) bonds indicates electron delocalization as a result of condensation.
- c) The length of the (C=NQ) bonds increases,

TABLE 2. Electronic spectral data* of the ligands in dioxane.

indicating the weakens of the bonds as a result of condensation.

- d) The formation of hydrazones is endothermic. The stability of ligands is higher than that of HQ.
- e) The ligands are favorite to coordinate to metal ion, due to the small value of ΔE of ligands[38].

Ligand	λ_1	ε	λ_2	ε2	λ,	ε,
HQ	247	59634			329	13401
BHQ	241	28697	351	38076		
AHQ	239	23901	344	22063	418	3929
Assignment	π ·	- π*	π - π'	* + CT	n ·	- π*

* λ_{max} in (nm) and ϵ in (cm mol / L)⁻¹.



Scheme 1. H-bonding and tautomeric forms of the hydrazone ligands.







HQ (NN-donor) (one 5-membered chelate ring)

BHQ (NNO-donor) AHQ (NNO-donor) (one 5- and one 6- membered chelate rings) Scheme 2. Chelating ability of the ligands.

pH-metric studies

Choice of the ligands

The ligands (phenolic: AHQ and etheric: BHQ) were selected to examine and investigate the effect of the following: (i) nature of the donor atoms, (ii) steric effects (BHQ), and (iii) rigidity of the structure produced by the presence of the benzene ring (AHQ and BHQ). This could be illustrated as shown in Schemes 1, 2.

Dissociation constants

Representative pH-metric titration curves of the free and complexed BHQ ligand at 303K in 75% (v/v) dioxane-water are presented in Fig. 1. All ligands dissociate just one proton, which is weakly ionized. The dissociated proton is that either of the phenolic group (AHQ) or belongs to



Fig. 1. pH-Metric titration curves of BHQ (3 x 10⁻³ M) in presence and absence of metal ions (1.5 x 10⁻³ M) in 75% (v/v) dioxane-water at 303K and Vo = 30 ml.



ofStability constantsinThe Irving-Rossotti[43] relations have beeng.used to calculate the parameter ñ (average numberchof ligand ions attached / one metal ion) and pLat(the free ligand exponent). The following Irving-toRossotti relations were used;

observed in our previous studies[40-42].

the-NH at the quinoline ring or in the side chain

(BHQ). Also, representative calculations of the

dissociation constants (pKH's) of the investigated

BHQ ligand according to Albert- Serjeant

method[39] are plotted graphically in Fig. 2. The

inhibition of the second dissociation in AHQ is

due to ionization of the phenolic proton firstly,

which increases the electron density and results in a more stable structure. Such behavior was



Fig. 2. Evaluation of the pK^H's of BHQ in 75% (v/v) solvent-water at 303K.



Fig. 3. Evaluation of the stability constants of the BHQ-complexes in 75% (v/v) dioxane-water at 303K.

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No.	Heat of formation	Dipole moment	HOMO Energy, [EV]	LUMO Energy, [EV]	ΔE_{gap}	global electro- philicity index (ω)	Electro- negativity (X)	global softness (S)	global hardness (ŋ)	C=N(Q)	N-N
HQ	-313808.8	0.9842	11.86395	13.84353	1.97958	83.4615	-12.8537	0.50516	0.98979	1.596	1.826
AHQ	-518082.3	3.633	12.09375	13.97758	1.88383	90.2038	-13.0357	0.53083	0.94192	1.664	1.733
BHQ	-518101.3	2.63	12.22082	14.04869	1.82787	94.3841	-13.1348	0.54709	0.91394	1.657	1.672

TABLE 3. Structural parameters and bond length of the compounds by DFT calculation.

TABLE 4. Stability constants and (K₁/K₂) for the complexes in 75% (v/v) dioxane-water at 303K.

Consta		AI	łQ			BHQ					
Complex	logK ₁	logK ₂	$log\beta_2$	K_1/K_2	logK ₁	logK ₂	$\log \beta_2$	K_1/K_2			
рКн	12.16				13.12						
Mn(II)	9.35	8.77	18.12	3.80	9.83	8.79	18.62	10.96			
Co(II)	10.98	10.00	20.98	9.55	8.53	8.55	17.08	0.96			
Ni(II)	10.01	9.33	19.34	4.79	8.60	8.50	17.10	1.26			
Cu(II)	11.43	9.06	20.49	234.42	10.70						
Zn(II)	9.99	9.68	19.67	2.04	9.15	9.25	18.40	0.79			
Cd(II)	9.50	9.17	18.67	2.14	8.51	8.21	16.72	2.00			
UO ₂ (II)	11.09	11.15	22.24	0.87	10.97	11.13	22.10	0.69			
La(III)	9.04	9.09	18.13	0.89	8.34	8.35	16.69	0.98			
Ce(III)	9.27	9.30	18.57	0.93	8.95	8.23	17.18	5.25			
Sm(III)	9.51	9.62	19.13	0.78	9.18	9.29	18.47	0.78			
Ho(III)	9.73	9.84	19.57	0.78	9.49	9.62	19.11	0.74			

 $Log (\tilde{n}/1 - \tilde{n}) = log K_1 - pL$ $\tilde{n} < 1.0$

 $Log (\tilde{n} - 1/2 - \tilde{n}) = logK_2 - pL \quad 1.0 < \tilde{n} < 2.0$

Also, representative graph of calculations of \tilde{n} and pL are given in Fig. 3. The data of stability constants were collected and summarized in Table 4. The degree of metal-ligand formation \tilde{n} is at the rang 0.1 – 1.9, suggesting that the higher species in solution is 1: 2; M: L.

Potentiometric titration curves

The titration curves (Fig. 1) give the following remarks:

- (i) All ligands dissociate just one proton between a = 0.00 and a = 1.0 in presence and absence of different metal ions (a = number of moles of KOH / mole of ligand). This suggests that all ligands act as monoprotic (monobasic) species; HL towards the metal ions in solution.
- (ii) For most transition metal ions (3d and 4d), two inflections (weak→sharp) at m = 1.0 and m =



Fig. 4. LogK vs. pK^H for the AHQ–complexes at different temperatures in 75% (v/v) dioxane–water.

2.0 were detected (m = number of moles of KOH / mole of metal ion). The two inflections correspond to the stepwise formation of monoand bis- chelates, respectively as shown by the following equilibria:

 $M^{+n} + HL + OH^{-} \qquad ML^{+n-1} + H_2O \quad (m = 1.0)$ $ML^{+n-1} + HL + OH^{-} \qquad ML_2^{+n-2} + H_2O \quad (m = 2.0)$ $M^{+n} + 2 HL + 2 OH^{-} \qquad ML_2^{+n-2} + 2 H_2O$ In general, this behaviour is consistent with

the values of K_1 / K_2 (Table 4).

(iii) The above feature was not observed with most 4f lanthanides and UO2^{II}, where only one overlapping buffer region between m = 0.0 to m = 3.0 (Fig. 1) was obtained, due to the formation of bis chelates directly as well as hydroxo complexes. This could be represented by the following equilibra:

Also, this is consistent with the lower values of K_1 / K_2 (Table 4).

The dissociation constants of the ligands and their stepwise formation constants with metal ions at 303 K in 75% dioxane-water are summarized in Table 4. Inspection of the data discloses the following:

- a) $(K_1/K_2) > 1.0$ for most complexes particularly 3d and 4d-complexes. This is due to the coulombic attractions between M²⁺ and L⁻, which are higher than those between ML⁺ and L⁻, in addition to some steric hindrance for the second coordination.
- b) $(K_1/K_2) \le 1.0$ for UO₂^{II} and most lanthanide(III) complexes, suggesting their higher favorite for the second coordination, which is regular with their higher coordination numbers (8 - 10).
- c) (K_1/K_2) has unusual high value for Cu^{II}-AHQ chelates, suggesting either a high steric hindrance for the second coordination or a

change in the dentate character from tridentate in 1:1 to bidentate in 1:2; M:L chelates. The latter is dependable with the greater tendency of Cu^{II} ions to form tetra- coordinated square planar species.

- d) The unexpected high stability constants of MnII- chelates (Table 4), suggest either a high selectivity of such class of hydrazones (containing the quinoline moiety) for MnII ions or an oxidation took place. However, such behavior is not observed with other hydrazones[42, 44].
- e) The high stability of LnIII-AHQ chelates; (Ln = La, Ce, Sm and Ho) is consistent with the high affinity of lanthanides for O- donors especially phenolic[45] and carboxylate[46, 47].
- f) For all complexes, $\log\beta_{2}$ for AHO is greater than that for BHO. This is due to the presence of a bulky Me group on the donor O-atom (BHQ), which causes the steric hindrance. In case of Co^{II} and Ni^{II}- complexes, BHQ shows reverse order. On the other hand, AHQ and BHQ exchange their positions in case of Mn^{II}-chelates.
- g) The higher stability of Cu^{II}-complexes compared to that of Ni^{II}-complexes is due to Jahn-Teller distortion in addition to the shorter Cu-N bonds than Ni-N bonds which increases the antibonding overlap[48]. On the other hand, the small stability of Cd^{II}-complexes is consistent with their CFSE = 0; Cd^{II} is d^{10} system.
- h) The unusual high stability of Co^{II}- AHQ chelate (Table 4) may be attributed to the oxidation of $Co^{2+} \rightarrow Co^{3+}$.
- i) For lanthanide(III) and UO2^{II}-complexes, $\log\beta_2$ has the order;

TABLE 5. Comparison of logK₁ for some *o*-hydroxyacetophenone hydrazones.

		logK ₁ in 75%(v/v) diox	kane-water at 303K	
Complex	AHQ (Quinoline)	AHP _m (Pyrimidine)	AHP _z (Pyridazine)	AHT (Triazine)
	Our Study	[18]	[19]	[25]
Co(II)	10.98	9.37	11.92	12.26
Ni(II)	10.01	9.21	11.85	12.32
Cu(II)	11.43	11.87	12.35	12.44
Zn(II)	9.99	8.85	10.19	11.03



Fig. 5. LogK vs. pK^H for the BHQ-complexes in 75% (v/v) solvent-water.

TABLE 6. LogK₁(str)* for some hydrazone–complexes in 75 % (v/v) dioxane-water at 303K.

		logV		logK ₁ (str) = logK ₁ (hydrazone chelates) - logK ₁ (HQ chelates).					
Complex		log K ₁							
	HQ	AHQ	BHQ	AHQ	BHQ				
Co(II)	7.65	10.98	8.53	3.33	0.88				
Ni(II)	7.13	10.01	8.60	2.88	1.47				
Cu(II)	11.01	11.43	10.70	0.42	-0.31				
Zn(II)	7.53	9.99	9.15	2.46	1.62				

 $UO_2^{II} > Ho^{III} > Sm^{III} > Ce^{III} > La^{III}$

The relatively high stability of UO_2^{II} complexes is due to the bonded O-atoms which increase the electrostatic attraction between the U^{VI} and the coordinated ligands, which in turn compensate the steric hindrance offered by oxygen of UO_2^{II} cation.

The order of Ln^{III}-complexes is consistent with the increase of ionic size Ho^{III}; (0.89 A°) La^{III}; (1.06 A°) i.e. this order is linearly correlated with either 1/r or z^2/r (ionic potential of the Ln^{III}-ions), where r is the ionic radii of Ln^{III}-ions.

Hardness / softness

According to the previous work in our laboratory, a significant comparison of the formation constant $(\log K_1)$ must be made for some similar hydrazones bearing the triazine[49], pyridazine[41], pyrimidine[42] and quinoline rings (our study) under the same experimental conditions as shown in Table 5. Inspection of the data gives that the order of stability for most metal

ions is as follows;

Triazine >Pyridazine > Quinoline > Pyrimidine

This order confirm that the N-atom of the ring shares in chelation, where the electron density on the that ring increases in this order.

Chelate effect

The investigated ligands; BHQ and AHQ act as tridentate (NNO- donors) and form two chelate rings and as shown in Scheme 2. In general, the formation of two chelate rings and also the resonance within chelate rings lead to increasing stability of the formed chelates (Scheme 2). The effect of such chelate ring size can be illustrated by comparing logK₁ for HQ-chelates, which can form one 5- membered chelate ring with logK₁ for the hydrazone chelates, which can procedure two chelate rings (Scheme 2 and Table 6). The data showed that logK₁(str) is positive and ranges from 0.42 - 3.33, (an exception Cu^{II}-BHQ).

Effect of the reaction medium

In order to provide an explanation of the dependence of pK^{H} and logK (or ΔG°) on the solvent parameters, the $pK^{H's}$ of the BHQ ligand and logK of its chelates were resoluted pH-metrically in 75% (v/v) solvent (dioxane, isopropanol, ethanol and methanol) -water at 303K. The results are listed in Table 7. Inspection of the data tells that:

- (i) As the polarity of the medium increases, the pK^H and logK values decrease.
- (ii) The following order showed the decreasing of stability of the chelates in 75% solvent-water.



Dioxane > Isopropanol > Ethanol > Methanol

This order means that the basicity of the ligand (pK^{H}) decreases by increasing the polarity of the medium and the coordinating ability of the solvent.

According to Gergely and Kiss[50], dioxane molecules progressively break-down the H–bonded of H_2O , while alcohols can form H–bonded associations with H_2O . Thus, it is predictable that the extent of H–bonding in alcohols is greater than that in dioxane.

Since the ligands are insoluble in water and



Fig. 6. LogK $_{\rm 1}$ vs. ${\rm E}_{_{\rm Tm}}\left(A \right)$ and AN $_{_{\rm m}}\left(B \right)$ for BHQ-complexes in 75% (v/v) solvent–water.

TABLE 7. Stability constants of the BHQ-complexes in 75% (v/v) solvent-water at 303K and calculation of logK inaqueous medium by using the LRA.

	Dio	Dioxane Isopropanol Ethanol Methanol		nanol	Н	20	Н	2 0				
	E _{Tm} (48.8)	E _{Tm}	()	E _{Tm} (55.05)	E _{Tm} (56.9)	Method	A ^a	Method	I B ^b
Complex	AN _m (37.30)	AN _m (44.16)	AN _m (44.60)	AN _m (47.70)	logV	logV	lagV	lagV
	logK ₁	logK ₂	logK ₁	logK ₂	logK ₁	logK ₂	logK ₁	logK ₂	- log ₁	log ₂	logic	log ₂
Mn(II)	9.83	8.79	8.17	7.47	7.42	6.30	6.51	6.09	4.06	3.73	4.42	4.13
Co(II)	8.53	8.55	7.25	7.31	6.49	6.54	6.03	6.31	4.05	4.39	5.12	4.68
Ni(II)	8.60	8.50	7.35	7.33	6.65	6.45	6.09	5.88	4.19	3.85	4.45	4.22
Cu(II)	10.70		9.93		8.84		$\downarrow \downarrow \downarrow \downarrow$				7.36	
Zn(II)	9.15	9.25	7.92	8.08	7.16	7.30	6.55	6.42	4.59	4.42	4.90	4.76
Cd(II)	8.51	8.21	7.33	6.95	6.33	6.15	5.92	5.94	3.82	3.97	4.17	4.24
UO2(II)	10.97	11.13	9.83	9.50	8.91	8.59	7.92	7.99	5.77	5.50	6.20	5.87
La(III)	8.34	8.35	7.19	7.23	6.41	6.52	6.23	6.37	4.38	4.62	4.66	4.87
Ce(III)	8.95	8.23	7.59	7.76	7.07	7.03	6.64	6.68	4.81	4.46	5.02	5.78
Sm(III)	9.18	9.29	7.82	8.08	7.25	7.51	6.76	7.03	4.89	5.24	4.31	5.49
Ho(III)	9.49	9.62	8.23	8.46	7.70	7.88	7.13	7.36	5.35	5.63	5.57	5.89
$\mathbf{p}\mathbf{K}^{\mathrm{H}}$	13.12=	±0.088	11.39	±0.085	10.88	±0.135	10.33	±0.099	8.	13	8.	33

^aThese values were obtained (at 63.1; E_{Tm}) by LRA of logK₁ logK₂ or pK^H vs. E_{Tm} of the 75% (v/v) solvent-water (r = 0.977 - 1.00). ^bThese values were obtained (at 54.8; AN_m) by LRA of logK₁ logK₂ or pK^H vs. AN_m of the 75% (v/v) solvent-water (r = 0.7099 - 0.970).

in an attempt to decide the dissociation constants $(pK^{H}s)$ of the BHQ ligand and its stability constants $(logK_1 and logK_2)$ in pure water, a LRA of pK^{H} , $logK_1$ or $logK_2$ vs. Reichardt (E_{Tm}) , (Fig. 6; method A) and acceptor number (AN_m) (Fig. 6; method B) was constructed and analyzed (Table 7). In conclusion, the stability of the chelates increases by decreasing both the E_{Tm} and AN_m of the solvent.

Effect of temperature

In an attempt to calculate the thermodynamic parameters, the formation constants of the phenolic AHQ ligand with Mn^{II}, Co^{II}, Ni^{II}, Cu^{II} and Zn^{II}-ions were calculated pH-metrically in 75% (v/v) dioxane-water at various temperatures *viz.* 283, 293, 303 and 313 K (Table 8).

The thermodynamic functions (Table 8) are calculated by using the well-known relations:

(i) $\Delta G^{\circ} = -19.12 \text{ T logK}$ where ΔG° is in J mol⁻¹.

(ii) LogK = $(-\Delta H^{\circ} / 19.12 \text{ T}) + (\Delta S^{\circ} / 19.12).$

Thus, on plotting logK vs. 1/T (Fig. 8), one gets a straight line whose slope = $-\Delta H^{\circ} / 19.12$ and its intercept = $\Delta S^{\circ} / 19.12$. Inspection of the data (Table 8) gives the following:

- (i) High positive values of ΔG° of the dissociation process indicate that it is non-spontaneous process. The large negative values of ΔG° of the formed complexes indicate that the complexation proceeds spontaneously[51].
- (ii) Large negative value of ΔS° of the dissociation process refers that the ionization



of the ligand is entropically unfavourable. Large positive values of ΔS° of the formed metal complexes[51] (an exception Cu^{II}-complex) indicate that the complexation process is entropically favourable and the mechanism of complexation is based upon replacement of H₂O by L⁻ according the following interaction:

$$[Ni (OH_2)_6]^{++} + 2HL \longrightarrow NiL_2 + 6H_2O + 2H^+$$

3 particles 7 particles

So that the above reaction proceeds with an increase of the entropy.

- (iii) Large negative values of ΔH° for all metal chelates indicate that the complexation process is enthalpically favourable and it is exothermic process[51].
- (iv) $\Delta S_{1}^{\circ} \ll \Delta S_{2}^{\circ}$ i.e. more orderless upon the second coordination. This is due to a large number of water molecules is released upon the second coordination than the first coordination. Furthermore, this is caused by the transformation of the dentate character of AHQ ligand from tridentate in 1:1 to bidentate in 1:2; M:L chelates to relieve steric hindrances (K/K₂ > 1.0; Table 4).

$$(v)$$
 $-\Delta H^{\circ}_{1} >> -\Delta H^{\circ}_{2}$

i.e. the bond strength from the ligand to the metal ion is stronger in the first coordination than that in the second coordination. Also, this provides another strong evidence of the



Fig. 8. ΔH° vs. ΔS° for AHQ-complexes in 75% (v/v) dioxane–water (Evaluation of Tiso).

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Complex	283K		293K		303K	K 313K			$-\Delta G^{\circ}$		-ΔH°		ΔS°	
Complex	logK ₁	logK2	logK ₁	logK ₂	logK ₁	logK ₂	logK ₁	logK ₂	$-\Delta G^{\circ}_{1}$	$-\Delta G^{\circ}_{2}$	$-\Delta H^{\circ}_{1}$	$-\Delta H^{\circ}_{2}$	ΔS°_{1}	ΔS_{2}°
Mn(II)	9.88	9.14	9.54	8.95	9.35	8.77	9.03	8.60	53.78	50.47	46.40	30.49	24.76	67.05
Co(II)	11.57	10.55	11.36	10.23	10.98	10.00	10.55	9.89	63.25	57.88	57.97	32.26	17.71	67.91
Ni(II)	10.46	9.69	10.29	9.50	10.01	9.33	9.72	9.17	57.60	53.65	42.17	29.31	51.77	81.65
Cu(II)	12.24	9.77	11.95	9.39	11.43	9.06	10.97	8.80	66.26	52.66	73.09	54.97	-22.91	-7.78
Zn(II)	10.56	10.07	10.23	9.84	9.99	9.68	9.80	9.49	57.74	55.62	42.81	32.20	50.11	78.58
pK^{H}	12.67		12.32		12.16		12.00		- 69.96		- 36.95		- 110.79	

TABLE 8. Stability constants and thermodynamic functions* of the AHQ-complexes in 75% (v/v) dioxane-water.

* ΔG° and ΔH° are in k J mol⁻¹ while ΔS° is in J mol⁻¹ K⁻¹ (r = 0.9831 - 0.9999).

alteration of the dentate character of the AHQ ligand.

(vi) The values of $\Delta H^{\circ} = \Delta H^{\circ}_{1} + \Delta H^{\circ}_{2}$ (Table 8) indicates that a tetrahedral geometry around Mn^{II}, Zn^{II} and Ni^{II}-ions, an octahedral environment around Co^{II}-ions and a square planar geometry around Cu^{II}-ions. The exothermicity has order:

$$D_{4h}$$
 >> O_{h} >> T_{d}

(Square planar) (Octahedron) (Tetrahedron)

Isokinetic ΔH° - ΔS° relationships

In an attempt to evaluate the isokinetic temperature (T_{iso}) i.e. the temperature at which the complexation reactions proceed at the same rate[52]. Using the LRA, the following relations for AHQ chelates in 75% (v/v) dioxane-water (Fig. 8) were obtained:

$\Delta H^{\circ}_{1} = 0.418 \Delta$	$S_{1}^{\circ} - 62.65$	$r^2 = 0.934$
$T_{1(iso)} = 418K,$	$\Delta G^{o}_{1(iso)} = -6$	2.65 kJ/mol.
$\Delta \mathrm{H^{o}}_{2} = 0.286 \Delta$	$S_{2}^{\circ} - 52.31$	$r^2 = 0.971$
$T_{2(iso)} = 286K,$	$\Delta G^{o}_{2(iso)} = -5$	52.31kJ/mol.

Inspection of the relations reveals the following:

- (i) $T_{1(iso)} >> T_{2(iso)}$ suggesting a change in the dentate character of AHQ ligand from tridentate (1 : 1) to bidentate (1 : 2; M : L), to relieve steric hindrances.
- (ii) $T_{1(iso)} >> T_{exp}$ i.e. the 1^{st} coordination is enthalpically controlled.
- (iii) $T_{2(iso)} \approx T_{exp}$ i.e. the 2nd coordination is entropically and enthalpically controlled.

Electrostatic and cratic thermodynamic functions

The thermodynamic parameters (Table 8) were analyzed into their electrostatic (el) and non-electrostatic (non) or cratic components (Table 9) to get an insight into the extent of ionic and covalent nature of the formed complexes. Inspection of the data (Table 9) reveals the following;

(i) ΔH°_{el} has positive values, this is consistent with $T = 298K > \theta = 219K$ and results in an endothermic contribution to the total change of enthalpy;

$$\Delta \mathrm{H}^{\circ}_{\mathrm{el}} = (\mathrm{T} - \theta) \, \Delta \mathrm{S}^{\circ}_{\mathrm{el}}.$$

TABLE 9. Electrostatic and cratic thermodynamic functions^{a,b,c} of AHQ-chelates.

			1stcoord	ination		2 nd coordination						
Complex	$-\Delta G^{o}_{el}$	- ΔG^{o}_{cratic}	$\Delta \mathbf{H}^{o}_{el}$	- ΔH^{o}_{cratic}	ΔS^{o}_{el}	T*	$-\Delta G^{o}_{el}$	- ΔG^{o}_{cratic}	$\Delta \mathbf{H}^{o}_{el}$	- ΔH^{o}_{cratic}	ΔS^{o}_{el}	Τ*
Mn(II)	12.75	41.03	4.60	51.00	58.20	362	22.01	28.46	7.94	38.43	100.49	338
Co(II)	11.20	52.05	4.04	62.01	51.15	401	22.20	35.68	8.01	45.66	101.35	368
Ni(II)	18.66	38.94	6.73	48.90	85.21	355	25.20	28.45	9.09	38.40	115.09	338
Cu(II)	2.31	63.95	0.83	73.92	10.53	443	5.62	47.04	2.03	57.00	25.66	416
Zn(II)	18.30	39.44	6.60	49.41	83.55	357	24.53	31.09	8.85	41.05	112.02	349

° T* is in K.

^a $\Delta G^{\circ}_{x} \Delta H^{\circ}_{x}$ are in k J mol⁻¹ while ΔS°_{el} is in J mol⁻¹ K⁻¹ (x = el or cratic).

^b $\Delta S^{\circ}_{cratic} = constant = -33.44 \text{ J mol}^{-1} \text{ K}^{-1}$ (not included in the Table).

(ii)
$$-\Delta G_{1 \text{ non}}^{\circ}, -\Delta H_{1 \text{ non}}^{\circ} >> -\Delta G_{2 \text{ non}}^{\circ}, -\Delta H_{2 \text{ non}}^{\circ}$$

(1st coordination) (2nd coordination)

This order suggests a higher degree of covalency in the 1:1 compared to the 1:2 complexes. Hence, the higher negative values of ΔH°_{non} would reflect the bond strength from ligand to metal ion which is stronger in the 1st coordination than the 2nd coordination.

(iii) ΔG°_{el} has negative comparable values and suggests that the complexation process is affected by the temperature and environment (Table 9).

Ligand field stabilization energy (LFSE)

N.B.: Calculation of the LFSE (Table 10) is based on the assumption that an O_h complex is formed in the solution.

The negative heat of the reaction;

$$Zn^{2+}_{(g)} + [MnL(H_2O)_n]^{2+} \longrightarrow Mn^{2+}_{(g)} + [ZnL(H_2O)_n]^{2+}$$

is defined as the transition series contraction energy, E_{r} , and this can be calculated from the following equation;

$$E_{r}^{}=(\Delta H^{\circ}_{h}+\Delta H^{\circ}_{C})^{Mn(II)}-(\Delta H^{\circ}_{h}+\Delta H^{\circ}_{C})^{Zn(II)}$$

For convenience, $E_r = \Delta H^{\circ}_r (Mn^{2+}) - \Delta H^{\circ}_r (Zn^{2+})$

where $\Delta H_r^{\circ} = \Delta H_h^{\circ} + \Delta H_C^{\circ}$; ΔH_C° is the heat of complex formation and ΔH_h° is the heat of hydration of the bivalent transition metal ions. The ligand field stabilization energy, ∂H , can be calculated by the following equation: $\partial H = \Delta H_r^{\circ}$ (Mn²⁺) - {(n - 5)/5}E_r - ΔH_r° (M²⁺)

Inspection of the data given in Fig. 9 and Table 10, reveals that;

A. There is some agreement between E_r and ∂H values for each metal complex indicating the identical coordination as was supported by Irving–Rossotti relationship[43].

B. The LFSE, ∂H , followed the order;

 $Cu^{II} > Ni^{II} > Co^{II}$

which is in good accordance to the Irving-Williams order for stability of the complexes.

Conclusions

In conclusion, the following remarks can be pointed out:

- (i) The hydrazones act as monobasic (monoprotic) ligands (HL) towards the metal ions in solution as evidenced from the titration curves.
- (ii) The maximum \tilde{n} values were found to be \approx 2, revealing that both ML and ML₂ species are formed in solution.
- (iii) For most complexes; $logK_1 > logK_2$ indicating that the binding of a first ligand with the vacant sites of the metal ions are more available than for a second one.
- (iv) Stable complexes will be formed with;
 - Higher charges and small sizes of the metal ions i.e. higher polarizing power.
 - Higher basicity of the ligands.
 - Lower temperatures, this is consistent with the exothermic complexation.
 - Less polar media. The strong coordinating ability of the solvents (polar solvents) retards the metal-ligand interaction.
- (v) The dissociation of AHQ is endothermic, nonspontaneous and entropically unfavourable while its complexation is exothermic, spontaneous and entropically favourable.
- (vi) The LFSE follows the order:
 - Co^{II} < Ni^{II} < Cu^{II}

TABLE 10. Ligand field stabilization energy (LFSE) of the AHQ-chelates in 75% (v/v) dioxane-water.

Metal ion	$\Delta H_{h}^{o}(k J)$	$\Delta \mathbf{H}^{0}_{\mathbf{c}}$	$\Delta \mathbf{H}^{0}_{\mathbf{r}}$	{(n-5)/5}E _r	$\partial H(k J mol^{-1})$	%(-T∆S° ₁ /	%(-T∆S° ₂ /
3d ⁵ (Mn ²⁺)	2733.72	76.89	2810.61	0.00		$\frac{\Delta \mathbf{G}^{\mathbf{o}}_{\mathbf{f}}}{13.72}$	<u>ΔG°</u>) 39.59
$3d^{7}(Co^{2+})$	2913.46	90.23	3003.69	78	115.08	8.34	34.96
3d8(Ni ²⁺)	2992.88	71 48	3064 36	117	136 75	26 78	45 35
54 (141)	2772.00	/1.40	5004.50	117	150.75	20.70	-5.55
$3d^9(Cu^{2+})$	2996.64	128.06	3124.70	156	158.09		
$3d^{10}(Zn^{2+})$	2930.60	75.01	3005.61	195 (E _r)		25.86	42.10

in accordance to Irving-Williams order.

(vii) The stability of the chelates in different media follows the order:

Methanol < Ethanol < Isopropanol < Dioxane

- (Viii) For M^{II}-AHQ chelates: ΔH° is the main driving force for 1:1 chelates, whereas both ΔH° and ΔS° are the driving forces for 1:2 chelates.
- (ix) The structural features of the hydrazones have a considerable effect on the stability of their chelates.

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در اسات جهديه و نظرية لمتراكبات ٢- الفا- (ارثو هيدروكسي فنيل)ايثيليدين هيدرازينو [-4و6-ثنائي ميثيل كينولين و2- الفا- (ارثو ميثوكسي فنيل)ميثليدين هيدرازينو [-4و6-ثنائي ميثيل كينولين

> فاطمة سامي و حسين صقر سليم و علي طه و مجدي شبل و فاتن حنفي قسم الكيمياء - كلية التربية - جامعة عين شمس - روكسي - القاهرة 11341- مصر.

تم تخليق عوامل التراكب (-2]الفا-(ارثو هيدروكسي فنيل) ايثيليدين هيدر ازينو [4-و-6تنائي ميثيل كينولين و-2]الفا-(ارثو ميثوكسي فنيل) ميثليدين هيدر ازينو [4-و-6تنائي ميثيل كينولين) و تم التعرف عليهم بالتحليل العنصري وأطياف الأشعة تحت الحمراء و المرئية و الكتلة. و لقد أجريت " المعايرات " الجهدية و تم تعيين ثوابت التفكك لهذه الهيدر ازونات و كذلك ثوابت الإستقرار لمتر اكباتها في 75% (ح/ح) مذيب-ماء لعامل التراكب الثاني و عند درجات حرارة مختلفة لعامل التراكب الأول. و لقد تم حساب ثابت التفكك لعامل التر اكب الثاني و ثوابت الثاني و عند درجات حرارة مختلفة لعامل التراكب الأول. و لقد تم حساب ثابت التفكك لعامل التر اكب الثاني و ثوابت الاستقرار لمتر اكباته في الماء عن طريق العلاقة بين ثابت التفكك و ثوابت الاستقرار مع عوامل المذيب. كما تم تعيين درجة الحرارة الحركية باستخدام التحليل الانحدار الخطي بين التغير في الانثالبي و التغير في الانتروبي لمتر اكبات عامل التراكب الألول. و تم تحليل دوال الديناميكا الاحدار الخطي بين الكهر وساتاتيكية و عوامل المذيب. كما تم تعيين درجة الحرارة الحركية باستخدام التحليل الاحدار الخطي بين التعير في الانثالبي و التغير في الانتروبي لمتر اكبات عامل التراكب الول. و تم تحليل دوال الديناميكا الاحدار الخطي بين