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Biomarkers charge-transfer complexes of melamine () CrossMark with quinol and picric acid: Synthesis, spectroscopic, thermal, kinetic and biological studies

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KEYWORDS

Charge transfer complexes; Melamine; TG; Biological evaluation **Abstract** Two new melamine (MA) charge transfer complexes with quinol and picric acid in aqua media have been synthesized and structurally characterized. The obtained complexes with the general formula [(MA)(acceptor)₂] with a 2:1 acceptor:donor molar ratio. Elemental analysis (CHN), electronic spectra, photometric titration, mid infrared spectra, ¹H NMR spectra and thermogravimetric analysis (TG) were used to predict the position of the charge transfer interaction between the donating and accepting sites. The MA CT-complexes were antimicrobial assessment against two kinds of bacterial and fungal species.

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1. Introduction

Melamine (MA; 1,3,5-triazine-2,4,6-triamine) is a kind of triazine analog with three amino groups and it can also be described as a trimer of cyanamide, three cyanamide units

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joined in a ring. Melamine has the chemical formula $C_3H_6N_6$ with molecular mass of just over 126, forms a white crystalline powder, and is only slightly soluble in water. It has been widely used as a raw material in the chemical industry. It is commonly used in the production of melamine resins, laminates, glues, adhesives, and combined with formaldehyde in the molding of plastics (Venkatasami and Sowa, 2010). Because of its high nitrogen content (66% nitrogen by mass), melamine shows the analytical characteristics of protein molecules when the Kjeldahl method is used for protein analysis in food (Xiang et al., 2011). Thus, melamine is unethically added to food products to have more protein content than they actually have. Melamine is a toxic compound to both animals and human beings (Lam et al., 2009). Unfortunately, it is possible that melamine accumulates in the body that causes toxicity prob-

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lems, which may lead to reproductive damages. It has been reported that the combination of melamine with cyanuric acid leads to the formation of insoluble melamine cyanurate crystals (Puschner et al., 2007), causing bladder or kidney stones, which can lead to renal failure (Brown et al., 2007) or bladder cancer (Heck and Tyl, 1985). Recently many young children have suffered from renal stone caused by melamine in the milk powder, and the contamination of milk with melamine in China likely caused 300 000 cases of renal complications in children in 2008, directly resulting from the consumption of tainted product (Mauer et al., 2009; Chan et al., 2008). As a result, governments around the world have enacted regulations that have set maximum acceptable limits for melamine in food products. Consequently, there is a demand for analytical techniques that can reliably monitor melamine in food products.

Currently, several analytical methods have been employed and developed for the determination of melamine in infant formula and milk-based products, including high-performance liquid chromatography (HPLC) (Kim et al., 2008; Sun et al., 2010; He et al., 2008), liquid chromatography/mass spectrometry (LC/MS) (Filigenzi et al., 2007; Ibanez et al., 2009; Sancho et al., 2005; Ehling et al., 2007), gas chromatography (GC) (Yokley et al., 2000), gas chromatography/mass spectrometry (GC/MS) (Li et al., 2009), thin layer chromatography (Rao et al., 2008), capillary electrophoresis (CE) (Tsai et al., 2009; Xia et al., 2010; Cook et al., 2005; Broszat et al., 2008), surface enhanced Raman spectroscopy (Lin et al., 2008), flow injection analysis based on chemiluminescence system (FI-CL) Zeng et al., 2011, electrochemical methods (Cao et al., 2009; Attia et al., 2011; Liu et al., 2011; Ma et al., 2011) and Matrix assisted laser desorption/ionization methods (MALDI) (Singh and Panchagnula, 2011).

Charge transfer (CT) interactions within a molecular complex forming an electron donor and electron acceptor involving a resonance with a transfer of charge from donor to acceptor were showed firstly by Mulliken (Mulliken, 1950, 1952; Mulliken and Pearson, 1969; Gutmann et al., 1997), and discussed widely by Foster (Foster, 1969). Formation of charge transfer complexes (CTC) is based on the interaction of energetically high lying HOMO of the donor with a low energy LUMO of the acceptor. In general, the charge transfer complexation occurs as an ionic band in a simple ion-radical pair interaction (Arslan et al., 2007). Molecular charge transfer materials become an attractive and realistic target for materials science to research for molecular CT systems accompanied with the changes in magnetic, transport optical, dielectrically properties and structural changes. CT complexes, which exhibit interesting optical, electrical and photoelectrical properties, play an important role in many electro physical and optical processes (Bortchagousky et al., 2004).

Charge transfer complexation is currently achieving the great importance in biochemical, bioelectrochemical energy transfer process (Roy et al., 2005), biological systems, and drug-receptor binding mechanism, for examples, drug action, enzyme catalysis, ion transfers through lipophilic membranes (Dozal et al., 2000), and certain π -acceptors have successfully been utilized in pharmaceutical analysis of some drugs in their pure form or in pharmaceutical preparations (Pandeeswaran et al., 2009; Pandeeswaran and Elango, 2010). Recently, many studies have been widely reported about the rapid interactions between different kinds of drugs and related compounds as donors with several types of σ and π -electron acceptors (Refat

et al., 2010a,b, 2011b,c; Refat, 2011; Refat and El-Zayat, 2010). On the other hand, electron donor–acceptor (EDA) interaction has a great importance in many application topics and fields, like in non-liner optical materials and electrical conductivities (Yakuphanoglu et al., 2005), second order non-liner optical activity (Krishnamurthy et al., 1988), micro emulsion, surface chemistry (Andrade et al., 2000), photo catalysts (Dabestani et al., 1998), dendrimers (Jakubiak et al., 2000), solar energy storage (Takahasi et al., 1993), organic semiconductors (Eychmuller and Rogach, 2000), as well as in studying redox processes (Brueggermann et al., 1992).

The charge transfer complexes of melamine have not been reported in the literature, therefore the aim of this paper is directed to investigate these complexes. For this purpose, two new CT-complexes formed between melamine (MA) as the donor with quinol (QL) and picric acid (PA) as π -acceptors have been studied spectrophotometrically. These complexes are readily prepared from the reaction of MA with QL and PA in aqua media. The synthesized MA CT-complexes were structurally characterized to interpret the behavior of interactions using IR. ¹H NMR. UV–Vis techniques and elemental analyses (CHN). Benesi-Hildebrand and its modification methods were applied for the determination of the spectroscopic physical data such as formation constant (K_{CT}) , molar extinction coefficient (ε_{CT}), standard free energy (ΔG°), oscillator strength (f), transition dipole moment (μ), resonance energy (R_N) and ionization potential (I_D) . The thermal behavior of the obtained complexes as well as kinetic thermodynamic parameters $(E^*,$ ΔS^* , ΔH^* and ΔG^*) have also been investigated. Finally, antimicrobial activity of MA complexes was tested against different Gram (+) and Gram (-)bacterial species such as, Staphylococcus aureus (S. aureus), Bacillus subtilis, Escherichia coli (E. coli) and Pseudomonas aeruginosa (P. aeruginosa), and antifungal screening was studied against two species, Aspergillus flavus and Candida albicans.

2. Experimental

2.1. Materials

Pure melamine (MA, MF = $C_3H_6N_6$), quinol (QL) and picric acid (PA) were obtained from Merck and were used without further purification.

2.2. Synthesis of MA CT-complexes

The solid CT-complexes of MA with QL and PA acceptors were prepared by adding 0.1 mmol of the MA (12.6 mg) to 50 ml boiled water, and the suspension was heated to 100 °C with stirring for 1 h until all MA content dissolved. 0.2 mmol of acceptor dissolved in 10 ml of water was added to the MA solution, and the mixture was then allowed to stir for 30 min. at room temperature. The suspended solid was collected by means of filtration, washed several times with hot water, and then dried under vacuum over anhydrous calcium chloride.

2.3. Photometric titration measurements

The photometric titration measurements were performed for the reactions of the MA donor with QL and PA acceptors at 295 and 354 nm, respectively. A 0.25, 0.50, 0.75, 1.00, 1.50, 2.0, 2.50, 3.00, 3.50 and 4.00 mL aliquots of a standard solution $(5.0 \times 10^{-4} \text{ M})$ of the appropriate acceptor in H₂O were added to 1.00 ml of 5.0×10^{-4} M MA also in H₂O. The total volume of the mixture was 5 mL. The concentration of MA (C_d^0) in the reaction mixture was kept fixed at 5.0×10^{-4} M while the concentration of the π -acceptors (C_a^0) changed over a wide range of concentrations from 0.25×10^{-4} M to 4.00×10^{-4} M, to produce solution in each case of donor:acceptor molar reaction varying from 4:1 to 1:4. The stoichiometry of the molecular CT-complexes was determined by the application of the conventional spectrophotometric molar ratio according to the known methods (Skoog, 1985) through the plot of the absorbance of each CT-complex as a function with the ratio of (C_d^0) : (C_a^0) . The modified Benesi-Hildebrand plots were obtained (Benesi and Hildebrand, 1949; Abu-Eittah and Al-Sugeir, 1976) in order to calculate the formation constant, K_{CT} , and the absorptivity, ε_{CT} , values for each CT-complex resulted from this study.

2.4. Instrumentation and physical measurements

The electronic absorption spectra of the donor, acceptors and the resulted CT-complexes were recorded in the region of (800-200 nm) by using a Perkin-Elmer Lambda 25 spectrophotometer with a 1 cm quartz cell. The mid infrared spectra (IR) measurements (KBr discs) of the solid CT-complexes were carried out on a Genesis II FT-IR spectrophotometer in the v =4000–400 cm⁻¹ range. ¹H NMR spectra were obtained on a Bruker DRX-250 spectrometer, operating at 250.13 MHz using a dual 5 mm probe head. The measurements were carried out at ambient temperature using DMSO-d₆ as the solvent and TMS as an internal reference. ¹H NMR data are expressed in parts per million (ppm), referenced internally to the residual proton impurity in DMSO solvent. Thermogravimetric measurements (TGA) were carried out in the temperature range 25-800 °C in air atmosphere using a Shimadzu TGA - 50H thermal analyzer. The elemental analyses of carbon, hydrogen and nitrogen contents were performed by the microanalysis unit at the Cairo University, Egypt, using a Perkin-Elmer CHN 2400 (USA). The data of the elemental analyses (CHN) of the reaction products, the CT-complexes, were matched with the molar ratio results from the photometric titrations.

2.5. Antimicrobial investigation

Antimicrobial activity of the MA CT-complexes as well as the pure solvent was determined using a modified Bauer-Kirby disc diffusion method (Bauer et al., 1966), against different Gram (+) and Gram (-)bacterial species; S. aureus, B. subtilis, E. coli and P. aeruginosa, and antifungal screening was studied against two species, A. flavus and C. albicans. The microanalysis unit at the Cairo University, Egypt, performed the investigations. For these investigations, briefly, 100 µl of the test bacteria/fungi was grown in 10 ml of fresh media until they reached a count of approximately 108 cells/ml for bacteria or 105 cells/ml for fungi (Pfaller et al., 1988). 100 µl of microbial suspension was spread on agar plates. Isolated colonies of each organism that might be playing a pathogenic role were selected from primary agar plates and tested for susceptibility. After incubation for 48 h, inhibition (sterile) zone diameters (including disc) were measured and expressed in mm. Standard

discs of Tetracycline (Antibacterial agent) and Amphotericin B (Antifungal agent) served as positive controls for antimicrobial activity, where filter discs impregnated with 10 μ l of DMSO solvent were used as a negative control. The concentration of each solution was 1.0×10^{-3} mol dm⁻³. Commercial DMSO was employed to dissolve the tested samples.

3. Results and discussion

3.1. Elemental analysis

The elemental analysis data (C, H, and N) of the MA CT-complexes were performed and the obtained results are as follows;

- [MA (QL)₂]; C₁₅H₁₈O₄N₆; Mol. wt. = 346.22; Calc.: %C, 52.02; %H, 5.20; %N, 24.28, Found: %C, 52.17; %H, 5.12; %N, 24.21.
- (2) [MA (PA)₂]; C₁₅H₁₂O₁₄N₁₂; Mol. wt. = 584.20; Calc.: %C, 30.82; %H, 2.06; %N, 28.77, Found: %C, 31.06; %H, 2.11; %N, 28.69.

It can be seen that the resulted values agree quite well with the calculated values, and the suggested values are matched with the molar ratio introduced from the photometric titration curves.



Figure 1 Electronic absorption spectra of MA CT-complexes coupled with titration spectra of each system at the detectable peak.

3.2. Electronic absorption spectra of the MA/QL and MA/PA complexes

In the case of the MA/QL system, once beginning the addition of QL to a solution of MA in water, the color of the solution changes in four steps until it is constant: colorless \rightarrow light yellow \rightarrow pink \rightarrow red. On the other hand, in the case of the MA/ PA system, upon addition of PA to the MA donor dissolved in water, a yellow precipitate was formed immediately. The electronic absorption spectra of the MA, π -acceptors and the resultant charge transfer complexes are shown in Fig. 1. The spectra revealed new strong absorption bands assigned to the CT interaction at 295 and 354 nm for MA/QL and MA/PA complexes, respectively. The photometric titration measurements between MA and the defined acceptors QL and PA in water based on the characteristic absorption bands with reference to the CT-complexes, Fig. 2), proved that the molar ratio of the resultant complexes is 1:2 donor:acceptor. The formation of 1:2 complexes was supported by both elemental analysis and thermal measurements. It was necessary to calculate the values of the equilibrium constant (K) and the extinction coefficient (ɛ) of the MA complexes. For this purpose, the 1:2 modified Benesi-Hildebrand equation was used in the calculations;

$$(C_a^0)^2 C_d^0 / A = 1 / K \varepsilon + 1 / \varepsilon \ C_a^0 (4C_d^0 + C_a^0)$$
(1)

where C_a^0 and C_d^0 are the initial concentration of the π -acceptor (QL and PA) and donor (MA), respectively, and A is the absorbance of the detected CT-band. The data obtained C_d^0 , $(C_a^0)^2$, $C_a^0(4C_d^0 + C_a^0)$ and $((C_a^0)^2 C_d^0)/A$ in water were calculated. By plotting $((C_a^0)^2 C_d^0)/A$ values versus $C_a^0(4C_d^0 + C_a^0)$, straight lines are obtained with a slope of $1/\varepsilon$ and an intercept of $1/K\varepsilon$, as shown in Fig. 3.

The values of both K and ε associated with these complexes are given in Table 1. The data resulted (Table 1) show that the MA/QL and MA/PA systems give high values of both formation constant (K_{CT}) and molar absorptivity (ε_{CT}). The high values of both K and ε of the resultant CT-complexes caused high stabilities of the formed CT-complexes to be expected as a result of the expected high donation of the MA nucleus. The value of ε for [MA (QL)₂] is about twice higher than that of [MA (PA)₂].

3.3. Determination of the spectroscopic and physical data of the MA CT-complexes

The spectroscopic and physical data such as standard free energy (ΔG°), oscillator strength (*f*), transition dipole moment



Figure 2 Photometric titration curves for MA/QL and MA/PA systems at detectable peaks of 295 and 354 nm, respectively.



Figure 3 The modified Benesi–Hildebrand plots of MA/QL and MA/PA systems at detectable peaks of 295 and 354 nm, respectively.

Table 1	1 Spectrophotometric results of the MA CT-complexes.									
Complex	$\lambda_{\max} (nm)$	$E_{CT} (\mathrm{eV})$	K (L mol ⁻¹)	$\varepsilon_{max} (\mathrm{L} \ \mathrm{mol}^{-1} \ \mathrm{cm}^{-1})$	f	μ	I_p	D	R_N	ΔG° (25 °C) (kJ mol ⁻¹)
MA/QL	295	4.22	5.18×10^{7}	4.17×10^{4}	60.04	61.35	10.95	32.70	0.79	44,013
MA/PA	354	3.52	5.57×10^{7}	8.65×10^{4}	37.34	53.00	10.08	32.70	0.80	44,192

MA	QL	Complex [MA (QL) ₂]	Assignments
3470 s	3262 br	3261 s, br	v(N–H); MA, complex
3420 s			v(O-H); QL, complex
3334 s			
3134 vs			
-	3031 m	3031 s	v(C-H); aromatic rings
2831 w	2857 m	2720 m	$v_s(C-H) + v_{as}(C-H)$; QL, complex
2685 w	2836 m	2607 m	
	2716 m		
2203 w	2590 w	2362 ms	v(C==N); MA
	2467 vw	2337 m, sh	Hydrogen bonding
-	1866 m	1859 w	Overtone of $\delta(C-H)$
	1855 m		
1666 vs	1628 w	1515 vs	$\delta(N-H)$; MA, complex
1641 sh, s	1609 w		$\delta(NH_3^+)_{def}$, complex
1532 vs, br			v(C=C) (in-ring), aromatic; QL
-	1518 vs	-	v(C=O) + v(C=C); QL
1470 vs	1477 vs	1472 vs	v(C==N); MA
1442 vs			v(C=C) (in-ring), aromatic; QL
-	1366 ms	1352 ms	$\delta(NH_3^+)_{sym}$, complex $\nu(C-C) + \nu(C-O)$
1200 m	1244 vs	1210 s	v(C–N); MA, complex
1031 ms	1222 vs	1097 m	v(C-O); QL
	1210 vs		
-	1164 ms	-	CH, in plane bend; QL
	1097 m		
814 s	827 s	829 s	$\rho(\mathrm{NH_3}^+)$, complex
773 m			N—H wag
736 m			
-	759 vs	761 s	C-H out of plane bending
		693 m	
616 ms	616 m	613 m	N—H wag
580 ms	525 ms	519 m	CH out of plane bend
494 ms, br			Skeletal vibrations

s, strong; w, weak; m, medium; sh, shoulder; v, very; vs very strong; br, broad.

v, stretching; v_s , symmetrical stretching; v_{as} , asymmetrical stretching; δ , bending.

(μ), resonance energy (R_N), and ionization potential (I_P), were estimated in water at 25 °C. The calculations can be summarized as follows;

The oscillator strength (f), which is a dimensionless quantity used to express the transition probability of the CT band, can be estimated by using the approximate formula (Tsubomura and Lang, 1964);

$$f = 4.319 \times 10^{-9} \int \varepsilon_{CT} dv \tag{2}$$

where $\int \varepsilon_{CT} dv$ is the area under the curve of the extinction coefficient of the absorption band in question vs. frequency. To a first approximation;

$$f = 4.319 \times 10^{-9} \varepsilon_{CT} v_{1/2} \tag{3}$$

where ε_{CT} is the maximum extinction coefficient of the CTband and $v_{1/2}$ is the half bandwidth in cm⁻¹, i.e., the width of the band at half of the maximum extinction. The observed oscillator strength values of the CT bands are given in Table 1. The transition dipole moment (μ) of the MA CT-complexes, Table 1, have been calculated from Eq. (4) (Rathone et al., 1997);

$$\mu = 0.0958 [\varepsilon_{CT} v_{1/2} / v_{\text{max}}]^{1/2} \tag{4}$$

The transition dipole moment is useful for determining if transitions are allowed, that the transition from a bonding π orbital to an antibonding π^* orbital is allowed because the integral defining the transition dipole moment is nonzero. The ionization potential (I_P) of the MA donor in their charge transfer complexes were calculated by using an empirical equation derived by Aloisi and Pignataro (1972) Eq. (5);

$$I_p(eV) = 5.76 + 1.53 \times 10^{-4} v_{CT}$$
(5)

where v_{CT} is the wave number in cm⁻¹ corresponding to the CT band formed from the interaction between donor and

MA	PA	Complex [MA (PA) ₂]	Assignments
3470 s	3416 br	3574 m	v(N–H); MA, complex
3420 s		3419 s, sh	v(O-H); PA, complex
3334 s		3375 s	
3134 vs		3255 m s, br	
		3157 ms, br	
-	3103 ms	_	v(C-H); aromatic rings; PA
2831 w	2980 sh	_	$v_s(C-H) + v_{as}(C-H); PA$
2685 w	2872 w	_	
2203 w	-	2363 m	v(C==N); MA
		2338 m, sh	Hydrogen bonding
-	1861 ms	_	$v_{as}(NO_2); PA$
1666 vs	1632 vs	1646 vs	$\delta(N-H)$; MA, complex
1641 sh, s	1608 vs	1605 s, sh	$\delta(\mathrm{NH_3}^+)_{\mathrm{def}}$, complex
1532 vs br		1564 vs	v(C = 0) + v(C = N)
		1546 s, sh	
-	1529 vs	1495 ms	v(C==C), (in-ring) aromatic
			v(C = O) + v(C = C)
1470 vs	1432 s	1428 m	v(C==N); MA
1442 vs			v(C-C) (in-ting), aromatic
-	1343 ms	1365 ms	v(C-C) + v(C-O)
	1312 w	1332 vs	$\delta(\mathrm{NH_3}^+)_{\mathrm{sym}}$, complex v
		1271 s	_s NO ₂
			$v_{as}(C-N)$
1200 m	1263 w	1194 w	v(C-N); MA, complex
1031 ms	1150 ms	1162 ms	CH, in plane bend
-	1086 s	1081 ms	v(C-O)
			$v_s(C-N)$
814 s	917 vs	980 mw	N—H wag
773 m		936 w	$\delta(C-H)$ in plane bending
736 m		912 mw	
-	829 w	818 w	δrock, CH ₂ rock skeletal vibrations
			C-H out of plane bending
-	781 s	786 ms	C-H out of plane bending
	732 s	744 m	$\rho(\mathrm{NH_3}^+)$, complex
	703 s	711 ms	Skeletal vibration
616 ms	652 sh	615 mw	N—H wag
580 ms	522 ms	546 w	CH out of plane bend
494 ms, br		522 w	Skeletal vibrations
			δ(ONO): ΡΑ

s, strong; w, weak; m, medium; sh, shoulder; v, very; vs very strong; br, broad.

 ν_{s} stretching; $\nu_{s},$ symmetrical stretching; $\nu_{as},$ asymmetrical stretching; $\delta,$ bending.

acceptor. The electron donating power of a donor molecule is measured by its ionization potential which is the energy required to remove an electron from the highest occupied molecular orbital. Briegleb and Czekalla (1960) theoretically derived the relation to obtain the resonance energy (R_N) given as below;

$$\varepsilon_{CT} = 7.7 \times 10^{-4} / [h v_{CT} / R_N] - 3.5]$$
(6)

where ε_{CT} is the molar absorptivity coefficient of the CT-complex at the maximum of the CT absorption, v_{CT} is the frequency of the CT peak and R_N is the resonance energy of the complex in the ground state, which, obviously is a contributing factor to the stability constant of the complex (a ground state property). The values of R_N for the complexes under study are given in Table 1. The energy (E_{CT}) of the π - π^* interaction between donor (PA) and acceptors, was calculated by using the equation derived by Briegleb (1960, 1964);



Formula 1 Suggested structure of [MA (QL)2] complex.



Formula 2 Suggested structure of [MA (PA)2] complex.

$$E_{CT} = (hv_{CT}) = 1243.667 / \lambda_{CT} (\text{nm})$$
(7)

where λ_{CT} is the wavelength of the CT band. The calculated values of E_{CT} are given in Table 1. The standard free energy

changes of complexation (ΔG°) were calculated from the formation constant by using the equation derived by Martin et al. (1969);

$$\Delta G^{\circ} = -2.303 RT \log K_{CT} \tag{8}$$

where ΔG° is the free energy change of the CT-complexes (kJ mol⁻¹), *R* is the gas constant (8.314 J mol⁻¹ K), *T* is the temperature in Kelvin degrees (273 + °C) and K_{CT} is the formation constant of the complex (L mol⁻¹) at room temperature.

3.4. IR and ¹H NMR spectra

Infrared spectral studies shed light on the place of donation in donor species and the differences occur in the spectra of both QL and PA charge transfer complexes. The full infrared

Table 4 Thermo analytical results for the MA CT-complexes.						
Samples	Stage	TG results temp. peak (°C)	Max. temp. (°C)	Weight loss (%)	Evolved moiety	
MA/QL	Ι	63–134	95	27.218	2QL + MA	
$(C_{15}H_{18}N_6O_4)$	II	191–275	233	38.579		
	III	346–571	476	13.138		
MA/PA	Ι	247–349	326	23.201	2PA + MA	
$(C_{15}H_{12}N_{12}O_{14})$	II	349–393	358	38.254		
	III	393–707	576	33.951		



Figure 4 Plots of Coats-Redfern (CR) and Horowitz-Metzger (HM) relations for [MA (QL)₂] and [MA (PA)₂] charge transfer complexes.

absorption bands of MA charge transfer complexes are recorded in Tables 2 and 3. The band assignments between 4000 and 400 cm^{-1} show the main characteristic absorption bands of the NH_3^+ group. For example the v(NH), $v_{def}(NH_3^+$), $\delta_{svm}(NH_3^+)$ and $\rho(NH_3^+)$ vibration of [MA (QL)₂] complex occurs at about 3260, 1500, 1350 and 830 cm^{-1} , respectively. The corresponding values for [MA (PA)₂] complex are 3150, 1560, 1330 and 740 cm^{-1} , respectively. The presence of these bands confirmed the proton transfer phenomena from the acidic center of each acceptor to the lone pair of electron of the $-NH_2$ group to form NH_3^+ ammonium ion upon complexation (Refat et al., 2011a). Also, the values characteristic of both -OH and the -NH2 groups happened to shift to lower values and also decreased for the intensities of vibration motions, which indicates the interactions placed among the -OH group of each of QL and PA and the -NH₂ group of MA through the hydrogen bonding (region 3000- 2000 cm^{-1}). There is some small changes in comparison between the spectra of the CT-complexes, those of the reactants, could also be notable from the expected changes in the symmetry, and electronic configurations of both reactants upon charge transfer complexation. According to these observations, the suggested structures of MA/QL and MA/PA complexes are presented in Formula 1 and 2, respectively.

¹H NMR (DMSO-d₆, δ ppm) data for symmetrical 6-amino-1,3,5-triazine-2,4-diaminium-4-hydroxyphenolate is $\delta = 4.70-4.76$ (s, 6H, 2NH₃⁺), 7.70 (s, 2H, NH₂), 7.83 (d, 4H, 2(Ar-H, C-2, C-6)), 8.25 (d, 4H, 2(Ar-H, C-3, C-5)), 8.75 (s, 2H, Ar–OH, phenolic protons). The ¹H NMR of melamine with quinol revealed 5 different bands at; $\delta = 4.70-4.76$ which attributed to the ammonium ion protons which are more deshielded, $\delta = 7.70$ for the NH₂ protons which becomes more deshielded due to the presence of two ammonium groups attached to the ring, $\delta = 7.83$ attributed for the quinol aromatic protons at C-2 and C-6 are more shielded due to their location ortho to the negative oxygen, while the band at $\delta = 8.25$ attributed to aromatic protons at C-3, and C-5 are ortho to the phenolic groups and considered more deshielded than the previous one, and the band at $\delta = 8.75$ is attributed to the phenolic protons of quinol. The reaction was expected to carry out through the 1:3 melamine to quinol, ratio but experimentally it was found that the reaction was carried out through the 1:2 melamine to quinol, ratio and this may be attributed to the fact that the formation of two ammonium ions attached to the ring decreases the ionic strength of the



Figure 5 SEM image of [MA (QL)₂] complex.



Figure 6 SEM image of [MA (PA)₂] complex.



Figure 7 TEM image of [MA (QL)₂] complex in aqueous media.

lone pair of electron of the third amino group and let them more delocalized and tends to be resonating to the triazine ring.

The reaction of MA as donor with PA as acceptor gave a new charge transfer complex named by 6-amino-1,3,5-triazine-2,4-diaminium-2,4,6-trinitrophenolate (Formula 2). ¹H NMR (DMSO-d₆, δ ppm): δ = 5.16 (s, 6H, 2NH₃⁺), 7.65 (s, 2H, NH₂), 8.10 (s, 2H, 2Ar—H, picrate protons). The presence of three nitro groups in the CT-complex led to the down field shift for the NH₃⁺ band that appeared in the ¹H NMR if compared to the quinol complex and this attributed to the withdrawing effect of these nitro groups. In addition, the ratio of the complex formation 1:2 melamine to picric acid was attributed to the same reason mentioned above.

3.5. Thermal analysis and kinetic measurements

The thermo analytical results of two MA complexes are listed in Table 4. The kinetic thermodynamic diagrams of the released MA CT-complexes of both QL and PA acceptors during the non-isothermal process ($\Delta T \neq 0$) heating in air atmospheres are shown in Fig. 4. The TG-DTG curve of MA-QL complex shows mass loss in three weak steps between 63 and 134 °C, 191 and 275 °C, 346 and 571 °C corresponding to endothermic peaks at 95 °C, 233 °C, and 476 °C, attributed to the fusion and evaporation of two molecules of 1,4-dihydroxybenzene (QL) then melamine moiety, respectively. The total mass loss found is 78.935% with few carbon atoms remaining as a final residual. For the [MA (PA)₂] charge transfer complex, the TG-DTG curve in O₂ atmosphere shows mass loss in three steps one of them has sharp event and thermal events corresponding to these losses or due to physical nature. The first mass loss between 247 and 349 °C, corresponding to an endothermic peak at 326 °C with mass loss 23.201% is due to decomposition of first picric acid molecule. The second stage of endothermic peak at 358 °C with mass loss 38.254% in TG curve is due to fusion of second picric acid molecule. In case of MA-PA charge transfer complex, it is thermally stable up to 247 °C and above this temperature, the mass loss exhibited between 349-393 °C and 393-707 °C with weight loss of 38.254% and 33.951%, corresponding to endothermic peaks at 358 °C and 576 °C attributed to the decomposition of picric acid and melamine molecules accompanied by carbonaceous residue.

The thermodynamic parameters of decomposition processes of both QL and PA melamine charge transfer complexes, called, activation energy (E_a), enthalpy (ΔH^*), entropy (ΔS^*), and Gibbs free energy change of (ΔG^*) were evaluated graphically by employing the Coats–Redfern and Horowitz–Metzger methods (Coats and Redfern, 1964; Horowitz and Metzger, 1963). These methods are integral methods assuming various orders of reaction and comparing the linearity in each case to select the correct order by using;

3.5.1. Coats-Redfern equation

$$\ln\left[\frac{1-(1-\alpha)^{1-n}}{(1-n)T^2}\right] = \frac{M}{T} + B \quad \text{for } n \neq 1$$
(9)

$$\ln\left[\frac{-\ln(1-\alpha)}{T^2}\right] = \frac{M}{T} + B \quad \text{for } n = 1$$
(10)

where M = -E/R and $B = \ln AR/\Phi E$; each of E, R, A, and Φ is the heat of activation, the universal gas constant, pre-exponential factor and heating rate, respectively. The correlation coefficient, r, was computed using the least square method for different values of n, by plotting the left-hand side of Eq. (9) or (10) versus 1000/T (Fig. 4).

3.5.2. Horowitz-Metzger equation

The relations derived are as follows; $\ln[-\ln(1-\alpha)] = \frac{E}{RT_m}\Theta$ (11)

where α , is the fraction of the sample decomposed at time *t* and $\Theta = T - T_m$.

A plot of $\ln[-\ln(1-\alpha)]$ against Θ , was found to be linear, from the slope of which *E*, was calculated and *Z* can be deduced from the relation:

$$Z = \frac{E\phi}{RT_m^2} \exp\left(\frac{E}{RT_m}\right)$$
(12)

where φ is the linear heating rate, from the intercept and linear slope of such stage, the *A* (Arrhenius factor) and *E_a* values were determined. The other kinetic parameters, ΔH , ΔS and ΔG were computed using the relationships;

$$\Delta H = E - RT \tag{13}$$

$$\Delta S = R \ln(Ah/kT) \tag{14}$$

$$\Delta G = \Delta H - T \Delta S \tag{15}$$

where k is the Boltzmann's constant and h is the Planck's constant.

The kinetic data obtained from the non-isothermal decomposition of the complexes are given in Table 5. The activation energy of the complexes is expected to increase with increasing

Table 5 Kinetic parameters determined using the Coats-Redfern (CR) and Horowitz-Metzger ((HM).

Complexes	Stage	Method	Parameters ^a					
			$\overline{E^*}$	A	ΔS^{*}	ΔH^{*}	ΔG^*	
[MA (QL) ₂]	1st	CR	4.81E + 04	1.89E + 02	-2.06E + 02	4.37E + 04	1.53E + 05	0.9932
		HM	5.50E + 04	1.62E + 03	-1.88E + 02	5.06E + 04	1.50E + 05	0.9889
$[MA (PA)_2]$	1st	CR	1.49E + 05	1.60E + 10	-5.57E + 01	1.43E + 05	1.78E + 05	0.9976
		HM	6.02E + 05	7.66E + 48	-6.85E + 02	5.97E + 05	1.68E + 05	0.9935
9 4 4 5 6		· · · · · · · · · · · · · · · · · · ·	-1 + c : T	1-1 77-1 4 77	1 4 6 1 4 4 1-1			

^a Units of parameters: E in kJ mol⁻¹, A in s⁻¹, ΔS in J mol⁻¹ K⁻¹, ΔH and ΔG in kJ mol⁻¹.

Table 6	The inhibition	diameter zone	values (n	nm) for	· MA	CT-complex	es
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Inhibition zone diameter (mm/mg sample)							
Bacteria	Fungi						
Bacillus subtilis, (G ⁺)	Escherichia coli, (G ⁻)	Pseudomonas aeruginosa, (G ⁻)	Staphylococcus aureus, (G ⁺)	Aspergillus flavus	Candida albicans		
0.0	0.0	0.0	0.0	0.0	0.0		
34.0	32.0	34.0	30.0	-	_		
-	_	-	-	18.0	19.0		
22.0	11.0	8.0	28.0	25.0	12.0		
9.0	14.0	5.0	19.0	11.0	7.0		
	Inhibition zone Bacteria Bacillus subtilis, (G ⁺) 0.0 34.0 - 22.0 9.0	Inhibition zone diameter (mm/ BacteriaBacillus subtilis, (G ⁺)Escherichia coli, (G ⁻) 0.0 0.0 0.0 34.0 32.0 $ 22.0$ 11.0 9.0 14.0	Inhibition zone diameter (mm/mg sample)BacteriaBacillus subtilis, (G ⁺)Escherichia coli, (G ⁻)Pseudomonas aeruginosa, (G ⁻) 0.0 0.0 0.0 0.0 34.0 32.0 34.0 $ 22.0$ 11.0 8.0 9.0 14.0 5.0	Inhibition zone diameter (mm/mg sample) Bacteria Bacteria Bacillus Escherichia Pseudomonas Staphylococcus aureus, (G ⁺) 0.0 0.0 0.0 34.0 30.0 - - - - - - 22.0 11.0 8.0 28.0 9.0 14.0 5.0 19.0	Inhibition zone diameter (mm/mg sample)BacteriaFungiBacteriaFseudomonas aeruginosa, (G ⁻)FungiBacillus subtilis, (G ⁺)Escherichia coli, (G ⁻)Pseudomonas aeruginosa, (G ⁻)Staphylococcus aureus, (G ⁺)Fungi Aspergillus flavus0.00.00.00.00.00.00.034.032.034.030.018.022.011.08.028.025.09.014.05.019.011.0		



Figure 8 Inhabitation zone diameter of melamine CT-complexes toward some kind of bacteria and fungi; 1, *B. subtilis*; 2, *E. coli*; 3, *P. aeruginosa*; 4, *S. aureus*; 5, *Aspergillus flavus* and 6, *Candida albicans*.

thermal stability of complexes. Hence, the ΔE^* value in the main stages for the [MA (PA)₂] complex is higher than [MA (QL)₂] complex. The calculated ΔE^* values using Coats–Redfern and Horowitz–Metzger methods for the main decomposition stage of the complexes are found to be E_a [MA (PA)₂] = 375.5 kJ mol⁻¹ > E_a [MA (QL)₂] = 51.55 kJ mol⁻¹. The negative values of ΔS^* seen in Table 5, indicate that the reaction rates are slower than normal (Frost and Pearson, 1961). Furthermore, these data indicate that the reactants.

3.6. Scanning (SEM) and transmission (TEM) electron microscopy

Scanning and transmission electron microscopy techniques give a general perception about microstructure, surface morphology, particle size and chemical composition of the respective picric acid and 1,4-dihydroxybenzene melamine charge transfer, [MA (PA)₂] and [MA (QL)₂] complexes. Figs. 5–7 designed the SEM and TEM photographs of the QL and PA CT-complexes. The uniformity and similarity between the particle forms of synthesized melamine complexes indicate that the existence of morphological phases of QL and PA complexes have a homogeneous matrix. A homogeneous phase formation of [MA (QL)₂] complex having ice, crack morphologies in the form of a dispersed with particle size 0.6 µm is exhibited in Fig. 5. The picric acid melamine CTcomplex has a single-phase formation of straight bundle of sticks with particle size 2.5 μ m, Fig. 6. The [MA (QL)₂] charge transfer complex nanoparticles were characterized with TEM technique. Transmission electron microscopy studies of [MA (QL)₂] nanoparticles were carried out to understand the shape and size of particles. The transmission electron photography for the nanoparticles of QL complex obtained in a neutral aqueous media with about 100 nm in diameter with spherical shape is shown in Fig. 7.

3.7. Biological assessments

Biological assessments were observed in terms of antimicrobial activities of target MA charge transfer complexes against Gram-positive (*B. subtilis* and *S. aureus*) and Gram-negative (*E. coli* and *P. aeruginosa*) bacterial species and two strains of fungus (*A. flavus* and *C. albicans*). Results from the agar disc diffusion tests for antimicrobial activities of target complexes are presented in Table 6, and illustrated in Fig. 8.

Both [MA (QL)₂] and [MA (PA)₂] tested charge transfer complexes displayed a different degree of antimicrobial effect that [MA (OL)₂] in nano spherical shape displayed a degree of antimicrobial activities greater than [MA (PA)₂] against all organisms tested with bacteria of gram (G+) and gram (G-) types and A. flavus fungus strain. Regarding the inhibition zone diameter, [MA (QL)₂] complex in nano type has the highest antimicrobial activity against all target organisms compared to other [MA (PA)₂] tested complex. In addition, [MA (OL)₂] complex in nano type gained approximately 65% of activity of antibacterial agent (Tetracycline) and Antifungal agent (Amphotericin B). The biomarker activity of [MA (QL)₂] complex in nano type may be due to the outer membrane of all target organisms (bacteria or fungus) more permeable for $[MA (QL)_2]$ complex in nano type structure. The most reasons for lethal action of tested compounds may be due to their interactions with critical interacellular sites causing the death of cells. The variety of antimicrobial activities of tested compounds may be due to a different degree of tested complexes penetration through cell membrane structure of target organism. In conclusion, the interactions between the melamine and 1,4-dihydroxybenzene (QL) acceptor in nano-structural form result in the development of the effectiveness of biological characters of melamine antibiotics.

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