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Insights on the coordination mode of quercetin with Al(III) ion from a combined experimental and theoretical study

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Combining potentiometric, spectroscopic and theoretical DFT computations we have studied the formation of Al(III)-quercetin complex in ethanol solution. The possible complexation sites have been considered on the basis of all the used experimental and theoretical tools. Results agree in proposing a 1:1 neutral complex and the possibility to have different isomers in solution.

Introduction

Flavonoids, the most abundant polyphenolic natural antioxidant compounds, are found in most plants, concentrating in seeds, fruit skin or peel, bark and flowers, and in a variety of beverages (tea, coffee, wines and fruit drinks). Flavonoids have been shown to have a variety of beneficial biological effects, including protection of cells against oxidative stress and antibacterial, anti-inflammatory and vasodilator activities.²⁻⁴ Polyphenol-rich diets have been repeatedly correlated with a low risk of developing cardiovascular diseases and cancers, the two major causes of mortality in occidental countries.⁵ Quercetin (3,3',4',5,7-pentahydroxyflavone, H₅Que), one of the most common flavonols present in nature (i.e. in grapes, onions, berries, green veggies and legumes), has attracted the attention of many researchers due to its biological properties. ⁶⁻¹⁰ In addition, it is well known that quercetin may chelate metal ions preventing the metal-mediated generation of damaging oxidizing radicals and protecting the biological targets against the oxidative stress. 11-14 In quercetin structure three chelation sites are possible: the 3-hydroxycarbonyl, the 5-hydroxy-carbonyl and the 3',4'-dihydroxyl functions (Scheme 1). 15,16

Scheme 1 Chemical structure of quercetin (H₅Que)

Previous experimental and theoretical investigations have been focused on the complexation of Fe(II) and Fe(III)^{17,18}, Cu(II)¹⁹ and Pb(II)²⁰ with quercetin in solution and with different stoichiometric ratio. Al(III)-quercetin complex has been previously studied at theoretical and experimental levels in solid phase²¹ and in solution combining spectroscopic UV measurement and semiempirical AM1 theoretical method.²² Furthermore, other two joint experimental and density functional studies concerns the complex in which the

experimental conditions gives stoichiometric ratio of 1:2, metal: ligand.^{23,24} Finally, the ability of flavonoids to form chelate complexes with a series of metals has been recently briefly reviewed.²⁵

Although the presence of these studies, the composition, structure and complex formation features are not exhaustively investigated and sometimes contradictory. ¹⁷⁻²¹

In this work, we have studied the complexation of quercetin with Al(III) ion in solution by using a combination of experimental (potentiometric measurements, IR and UV spectra) and computational (density functional theory) tools in order to attain structural and electronic properties of the resulting complex.

With respect to previous work on this subject we underline that in our investigation we use potentiometric and spectrophotometric measurements in a wide range of pH and first principle computations on molar complex ratio of 1:1 (metal: ligand) as indicated by the experimental data and scarcely investigated previously.

The reason for the choice of aluminium cation is essentially due to the fact that it is third most abundant element in the Earth's crust and often enters in the biotic cycle in many different ways.²⁶ The human exposure to aluminium is not fully explained but it is well known that it does not serve any essential function in human biochemistry.²⁶ By contrary, Al(III) cation can enter in the brain where persists for long time and a small increase seems sufficient to produce neurotoxicity.²⁷ For this reason, the presence of aluminium in the human brain has been associated to the Alzheimer's and other neurodegenerative diseases.²⁸ Chelation therapy with some ligands has been demonstrated to reduce some causes of aluminium toxicity.^{26,28} In this context, it is interesting to explore the ability of a natural product as quercetin to coordinate the Al(III) ion.

Experimental and Theoretical details

Computational details

Full optimizations in solution without any symmetry restriction have been carried out at DFT level employing the M052x²⁹ exchange-correlation functional coupled with the 6-31+G(d) basis set for all atoms. Solvent effects have been described through a continuum approach by means of the SMD version of the polarizable continuum model (PCM).³⁰ The dielectric constant of ethanol has been fixed at 24.85. Vibrational frequencies have been computed on each

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optimized structure at the same level of theory of the optimizations, and zero point vibrational corrections have been added to the electronic energies. TD-DFT approach³¹ has been used to obtain the vertical excitation energies. In these computations different hybrid and meta-hybrid exchange-correlation functionals (M052x, 29 M0632 and wB97XD³³) have been employed.

All the computations have been performed with the Gaussian (G03) suite of programs.³⁴

Potentiometric measurements

The perchloric acid stock solution and the sodium hydroxide titrant solutions have been prepared and standardized as previously described. 35,36 A sodium perchlorate stock solution has been prepared and standardized according to Biedermann.³⁷ Aluminium(III) perchlorate has been prepared and standardized as reported by Ciavatta and Iuliano.³⁸ All solutions have been prepared with ethanol. The cell arrangement was similar to that described by Forsling and Hietanen.³⁹ Ag/AgCl electrodes have been prepared according to Brown.⁴⁰ Glass electrodes, manufactured by Metrohm, have been of the 6.0133.100 type. They acquired, after the addition of the reagents, a constant potential within 15 min that remained unchanged within \pm 0.1 mV for several hours. The titrations have been carried out with a programmable computer controlled data acquisition switch unit 34970 A supplied by Hewlett Packard. The EMF values have been measured with a precision of $\pm 10^{-5}$ V using an OPA 111 low-noise precision DIFET operational amplifier. A slow stream of nitrogen gas has been passed through three bottles (a-c) containing: a) 1 M NaOH, b) 1 M H₂SO₄ and c) 0.16 M NaClO₄, and then into the test solutions, stirred during titrations, through the gas inlet tube. During the EMF measurements, the cell assembly has been placed in a thermostat kept at (310.1 ± 0.1) K.

Vibrational spectra

The solid-state IR spectra (in KBr pellets) have been recorded on a Perkin-Elmer Spectrum One FT-IR spectrometer equipped for reflectance measurements.

Spectrophotometric measurements

The spectrophotometric measurements have been conducted with a Varian Cary 50 Scan UV visible Spectrophotometer. Absorbance values between (250 and 550) nm have been measured each 5 nm. The temperature of the cell holder has been kept at (310.1 ± 0.3) K by a Grant circulating water bath. Matched quartz cells of thickness 1 cm have been employed. The absorbance, A_{λ} , has been recorded to 0.001 units. The formulations of the parameters and the acquisition of the data have been managed with the aid of a computer connected to the tool.

Results and discussion

A sensitive tool to attain information on the formation and stoichiometry of metal complexes is potentiometric analysis. This methodology allows to measure potential difference as function of ion concentration in solution. The complex formation equilibria between Al(III) and quercetin has been studied, using ethanol as solvent at 310.15 K and in 0.16 M NaClO₄, by measuring with a glass electrode the competition of the ligand (H5Que) for the aluminium (III) and H^+ ions. The metal (C_M) and ligand (C_L) concentrations, ranged from 0.5 to 5 mM (the ligand-to-metal ratio varied between 1 and 10). The hydrogen ion concentration has been varied from 25 mM (pH 1.6) to incipient precipitation of basic salts which takes place in the range $[H^{+}] = 1.6-0.25$ mM (pH 2.8-3.6) depending on the specific ligand-to-metal ratio. A summary of the

relevant data taken in all titrations are reported in Table S1. The general equilibrium can be written as follows:

$$p \text{ Al}^{3+} + r \text{ H}_5 \text{Que} \rightleftarrows \text{Al}_p \text{H}_{-q} (\text{H}_5 \text{Que})_r^{(3p-q)} + q \text{ H}^+$$
 $\beta_{pqr} (1).$

This formulation takes into account the possible formation of simple (q=r), mixed $(q\neq r)$, mononuclear (p=1) and polynuclear (p>1)species. The most probable p, q and r values and the corresponding constants β_{par} have been obtained by a least squares fitting of the potentiometric data. 41 In the numerical treatments the first three acidic constants of quercetin, according to equilibria (2-4) and reported with the relative standard deviation, have been maintained invariant:

$$H_5Que \rightleftharpoons H_4Que^- + H^+ \qquad log K_{a1} = -8.40 \pm 0.01$$
 (2)

$$H_4Que^- \rightleftarrows H_3Que^{2-} + H^+$$
 $log K_{a2} = -7.83 \pm 0.06$ (3)
 $H_3Que^{2-} \rightleftarrows H_2Que^{3-} + H^+$ $log K_{a3} = -7.91 \pm 0.09$ (4)

$$H_3Que^{2-} \rightleftarrows H_2Que^{3-} + H^+ \qquad log K_{a3} = -7.91 \pm 0.09$$
 (4)

These constants have been determined by potentiometric measurements in the same experimental conditions used for the stability constants evaluation between metal and ligand ions (i.e. in ethanol as solvent at 310.15 K and in 0.16 M NaClO₄). The last two acidic constants of quercetin have not been considered due to the experimental pH range (i.e. 1.6-3.6).

The equilibrium constants for AlOH²⁺, Al(OH)₂⁺ and Al(OH)₄⁻ have been kept fixed during the numerical treatment because they are well known from the literature. 42 Various models have been tested by adding a single species during the elaboration; the best agreement has been obtained with the neutral complex Al(H)-3(H5Que), in which the metal-ligand stoichiometry is 1:1 and whose stability constant, $\log \beta_{1.-3.1}$, is -5.79 ± 0.06 (the uncertainty represents 3σ) according to the general equilibrium (1). As no other species lowered the minimum, this model has been assumed as the best data fitting, also in consideration that the standard deviation (σ =0.4) is comparable with the experimental uncertainty (σ =0.2).

The refined equilibrium constant as previously determined has been used to represent the distribution of the metal in the different species (see Fig. 1).

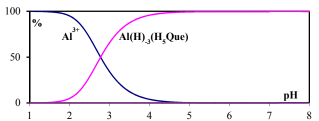


Fig. 1 Distribution of Al(III) species for $C_L = 5 \cdot 10^{-3} \text{ mol/dm}^3$, $C_M =$ $5.10^{-3} \text{ mol/dm}^3$

As can be seen from Fig. 1, the proposed complex is formed in appreciable amounts and none of the hydrolytic species reach a significant percentage. Information regarding the structure and the coordination sites of the dissolved metal ion complex cannot be obtained using the employed experimental method but can be supplied by high level theoretical investigation.

Following the indications coming from the potentiometric measurements we have considered the complexes with zero total charge and with the aluminium ion that retains the esa-coordination (see Scheme 2).

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Scheme 2 Formation of the investigated Al-complexes starting from the considered deprotonated forms of quercetin

The considered complexes include structures in which quercetin is—deprotonated in 3 or 5 positions and with Al ion bi-coordinated with quercetin and surrounded by two water molecules and two OH groups (see a3-4 and a4-5 species in Scheme 2), or deprotonated in—both 3 and 5 positions and linked to the Al(H₂O)₃(OH) moiety (see b3-4 and b4-5 species in Scheme 2). Since the experimental solution pH, the complex resulting from neutral ligand has not been considered. In all cases both the equatorial (a3-4eq, a4-5eq and b4-5eq, b4-5eq) and axial (a3-4ax, a4-5ax and b4-5ax, b4-5ax)—topologies for the OH groups have been considered. The energetic data are collected in Table 1.

Table 1 Total (hartree) and relative (kcal/mol) energies for the neutral species

Species	E _{ZPE}	ΔE
H ₄ QueA	$I(H_2O)_2(OH)_2$	
a3-4eq	-1650.390618	2.9
a3-4ax	-1650.389227	3.6
a4-5eq	-1650.393320	1.2
a4-5ax	-1650.395214	0.0
H ₃ QueA	l(H ₂ O) ₃ (OH)	
b3-4eq	-1650.387088	5.1
b3-4ax	-1650.383435	7.4
b4-5eq	-1650.385786	5.9
b4-5ax	-1650.383394	7.4

Results show that the complex in which the metal is coordinated to the positions 4 and 5 between the rings A and C (a4-5ax) is the most stable.

In the next lying minimum, at 2.9 kcal/mol, the Al ion coordinates to the C=O moieties in 3 and 4 positions inside of the ring C (a3-4eq). The zero charge complexes derived by the deprotonation of both the 3 and 5 O-H groups are higher in energy by 5.1 kcal/mol (b3-4eq) and 5.9 kcal/mol (b4-5eq), respectively (see Table 1). The structures of the two complexes have been reported in Fig. 2.

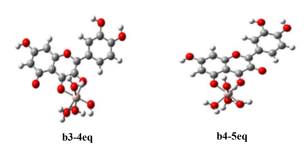


Fig. 2 M052x optimized structures of b3-4eq and b4-5eq complexes

The greater stability is reached when a six membered ring is formed upon the coordination process. From the obtained structures emerges that the coordination occurs in a planar arrangement in all the studied species. In Table 2 are reported the geometrical parameters involving the coordination of metal ion for all the optimized structures.

Table 2 Main geometrical parameters of the investigated structures by using two different protonation states of quercetin. Distances are in Å and angles are in degrees (°)

Species			H ₄ Qu	eAl(H ₂ O	$)_2(OH)_2$		
	Al-	Al-	Al-	Al-	Al-	O3-	O4-Al-
	O3	O4	O5	Ow	OH	Al-	O5
						O4	
a3-4eq	1.896	1.982	-	2.045	1.811	81.7	-
a3-4ax	1.902	1.957	-	1.999	1.836	83.3	-
a4-5eq	-	1.957	1.891	2.026	1.820	-	88.5
a4-5ax	-	1.952	1.882	1.998	1.827	-	91.0
			H ₃ Qu	eAl(H ₂ C	O)3(OH)		
b3-4eq	1.857	1.910	-	1.987	1.787	84.4	-
b3-4ax	1.860	1.903	-	1.966	1.797	85.2	-
b4-5eq	-	1.865	1.848	1.987	1.799	-	93.2
b4-5ax	-	1.868	1.838	1.982	1.801	-	93.8

In the most stable structure (a4-5ax) the Al-O5 distance is 1.882 Å while the Al-O4 one is 1.952 Å reflecting the fact that the C-O5 bond is 1.309 Å versus the 1.273 Å of C=O4 carbonyl bond. This behaviour is similar in all other studied structures. The O4-Al-O5 angle is 91° in a4-5ax while assumes a smaller value when the coordination occurs with the oxygen atoms that lie in the same ring (e.g. 81.7° in a3-4eq). Since the pH of the experimental solution and according to the equilibria (3) and (4), we can reasonably hypothesize that both the O3-H and O5-H groups, that have very similar p K_a (7.83±0.06 and 7.91±0.09), undergo deprotonation. In this situation it is interesting to further consider the corresponding complexes b3-4eq and b4-5eq even if exhibit a slightly reduced energetic stability. These two complexes differ in energy by only 0.8 kcal/mol and can be both populated in the experimental conditions. In order to have further insights on this possibility we have considered the interconversion process that can occur throughout the rotation of the Al(H₂O)₃(OH) fragment around the C-O bond (Ψ angle in Fig. 3). Along this path we have located a transition state that lies at 5.8 kcal/mol above the lowest minimum (Fig. 3). This result suggests that the interconversion between to the two b3-4eq and b4-5eq isomers is kinetically possible and both species should be present in solution.

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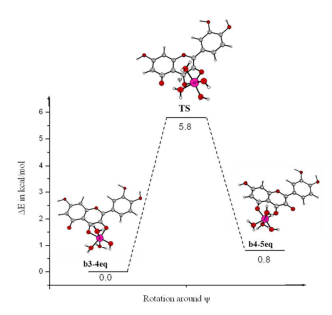


Fig. 3 Potential energy profile for the interconversion process as a function of the rotation around the C-O bond (Ψ angle) between b3-4eq and b4-5eq isomers

The computed and experimental vibrational spectra of metallated complexes have been reported in Figures S1 and S2 respectively, while the main stretching modes that can be affected by the complex formation, are given in Table 3. We underline that the obtained experimental spectra concern the solid phase while the simulated ones are referred to the liquid phase. Since our work is mainly devoted to the complexation process in solution we think that the comparisons can be regarded with cautions.

Table 3 Selected calculated and experimental vibrational frequencies (cm⁻¹) of Al(III)-quercetin complex

Mode	b3-4eq	b4-5eq	
C ₃ -O ₃	1634	1576	
C_4 - O_4	1582	1540	
C_5 - O_5	1390	1371	
	Exp		
1638, 1:	596, 1579, 1545	, 1514, 1374, 1328	

Table 3 clearly show that the considered C-O vibrational modes in the **b4-5eq** assume values lower than that in the corresponding **b3-4eq** complex. Looking at the experimental values in the frequency range that include the C-O vibrational modes we note that the 1579, 1545 and 1374 cm⁻¹ well fits with the three C-O stretching computed for the **b4-5eq** structure.

To gain additional insight on the complex formation, we have compared the UV-vis spectra of the free and bound quercetin. Absorption spectrum of quercetin 0.1 mM in ethanol solution is presented in Fig. 4 (line 1).

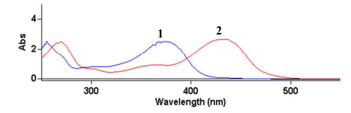


Fig. 4 UV-vis spectra of the quercetin 0.1 mM (line 1) and of the complex formed between quercetin and Al(ClO₄)₃ 0.1 mM (line 2)

Ouercetin exhibits a strong absorption band at 368 nm. Upon addition of Al(ClO₄)₃ to quercetin in solution (1:1 ratio of Al(III) salt:H₅Que) the UV-vis data showed a 60 nm bathochromic (red) shift in absorbance (of the corresponding peak for quercetin) indicating that the complexation occurs (Fig. 4, line 2). Moreover an isobestic point has been observed at 392 nm, confirming the presence of two species in equilibrium and therefore the formation of the complex

In order to obtain further information on the origin of the UV-vis spectra we have computed the excitation energies for quercetin and the two more probable complexes (a3-4eq and b4-5eq) by using different exchange-correlation functionals in the framework of timedependent density functional theory.³¹ Results are reported in Table

Table 4 Excitation energies (nm) and oscillator strength in parenthesis for a3-4eq, b4-5eq and guercetin in ethanol computed by using different exchange-correlation functionals

a3-4eq	b4-5eq	quercetin
	M052x	
396 (0.50)	416 (0.63)	312 (0.59)
245 (0.54)	251 (0.48)	232 (0.49)
	M06	
438 (0.37)	460 (0.55)	350 (0.48)
270 (0.37)	261 (0.24)	258 (0.21)
	wB97D	
400 (0.50)	426 (0.58)	317 (0.56)
247 (0.59)	251 (0.32)	234 (0.51)
	Exp	
429,269	429,269	368,255

For quercetin, although all the employed exchange-correlation functionals well reproduce the experimental data, better agreement has been found in the M06 values. For this system, the main bands at 255 and 368 nm are due to a HOMO→LUMO+1 (42%) and HOMO→LUMO (69%) orbital transition, respectively. In the a3-4eq and b4-5eq species, the two main transitions are originated by the HOMO-4→LUMO (about 70%) and HOMO→LUMO (about 60%) excitations. The M06 excitation energies for the considered complexes give results closer to the experimental absorption for the a3-4eq species, while the other two functionals seem to privilege the agreement with the **b4-5eq** complex. In any case the differences between the experimental and theoretical absorption spectra support the possibility that both the a3-4eq and b4-5eq isomers can contribute to the experimental UV-vis spectrum.

Since the presence of the complex (see Fig. 1) at physiological pH and the presence of Al(OH)₄ hydrolytical species in absence of the ligand (see Figure S3) that indicate as the ligand is able to quantitatively sequester the metal, our results can be related to the aluminium toxicity in living organism since highlight the properties and the formation process of Al(III)-quercetin complex that can bind to the DNA affecting its transcription and, consequently, inhibiting growth of cancer cells.²⁵ In addition, our results can be related also to the reduction of toxic metal bioavailability, since the formation of Al(III)-quercetin complex reduces the excess aluminium in diet thereby reducing the role of the aluminium in neurological disorders. 20,21,25

Conclusions

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In this work we have investigated the formation of Al(III)-quercetin complex in solution by using a combination of experimental (potentiometric measurements, IR and UV spectra) and computational (density functional theory and time-dependent density functional theory) tools. From our results the following conclusions can be drawn:

- in the considered pH range only complexes with 1:1 stoichiometric ratio between metal and ligand are possible;
- the best fit of the potentiometric data indicates that the formed species should be neutral;
- DFT computations on the possible hydrated complexes indicate that the preferred complexation site of quercetin should be the deprotonated 3 and 5 positions of the A and C ring, respectively. Moreover, the transition between the a4-5ax and a3-4eq requires a low barrier indicating that both species can be present in solution;
- the UV-vis experimental and theoretical spectra indicate the possible coexistence of the two isomers in the ethanol solution;
- at physiological pH quercetin is able to efficiently sequestrate the aluminium ion.

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Notes and references

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Electronic Supplementary Information (ESI) available: The primary data comprise 4 titrations with 65 data points, and summary of the relevant data taken in all titrations have been reported in Table S1. The computed and experimental vibrational spectra of metallated complexes have been reported in Figures S1 and S2, respectively. The distribution diagram of Al(III) ion in the hydrolytical species has been reported in Figure S3.

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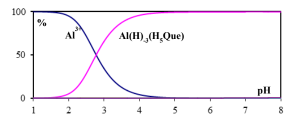
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TABLE OF CONTENTS

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Taking into account the importance of aluminium it is interesting to explore the ability of quercetin to coordinate it.