

Modelling Interactions Between Flavanols and Amine Acids: Case of Catechin and Epicatechin with Alanine; NBO, AIM, NCI Analysis

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Abstract: The interactions between two flavanols (Catechin and Epicatechin) and (Ala) Alanine (aliphatic amino acid) are evaluated by theoretical chemistry methods. Calculations at the level DFT/B3LYP/6-31+G (d, p) determine their characteristics and those of the monomers. Geometric, energetic, and spectroscopic parameters in addition to QTAIM (Quantum Theory of Atoms In Molecules), NBO (Natural Bond Orbital) and NCI (Non-Covalent Interaction) topological analyses qualify the nature and type of these. The results indicate that the main interactions are O-H...O and O-H...N between the hydroxyl groups of Cat (Catechin) or Epicat (Epicatechin) and the heteroatoms of Ala. They mention the existence of a secondary one alongside the main. They classify them into proper, improper, moderate, and weak. The spectroscopic parameters prove that O-H...O, O-H...N and N-H...O are proper. They establish that the C-H...N and C-H...O are improper. QTAIM analysis presents O-H...O, O-H...N interactions as moderate and C-H...O and N-H...O as weak. Stabilization energies show that the most reactive sites of Ala N_{sp}^3 and O_{sp}^2 interact strongly with the $O_{28}-H_{29}$, $O_{32}-H_{33}$ and $O_{34}-H_{35}$ hydroxyl groups of EpiCat and Cat. These interactions lead to the most stable complexes. This research reveals the existence of the VDW (Van Der Waals) NCI type and repulsive (steric) interactions in these complexes.

Keywords: Flavanols, Catechin, Epicatechin, NBO, AIM, NCI, Hydrogen Bond

1. Introduction

Polyphenols are organic compounds well known to interact with organic and inorganic compounds. They are found in the vegetation kingdom. They are metabolized by plants to fight against all kinds of external aggression [1, 2]. Humans consume them primarily for their many antioxidants or anti-inflammatory properties [3–8]. It is also used to prevent and treat neurodegenerative diseases [9, 10]. Polyphenols are grouped into classes; these include phenolic acids, hydroxy-

cinnamic acids, phenylacetic acids, aceto-phenols, coumarin, xanones, stilbenes and flavonoids [11, 12]. Each is made up of several subfamilies. So, flavonoids contain that of flavanols.

When eating certain foods (fruits, vegetables, wine, and tea), polyphenols are released into the body. These encounter its compounds. They generate interactions. Indeed, a lot of theoretical and experimental research establish their existences between polyphenols and organic or inorganic molecules. Proteins, lipids, or sugars represent the first. Iron and magnesium are some examples of the second [13–17]. These interactions generally take place with the hydroxyl

groups of the polyphenols [18]. The theoretical study of Cat (Catechin) and EpiCat (Epicatechin) proves that the most reactive sites are the latter [19].

Moreover, many authors attest that the interactions of polyphenols with proteins are usually non-covalent, including those with HB (Hydrogen Bonds) [20–22]. These are important in biological processes; they influence the properties of water as a universal natural solvent [23]. They also contribute determining the structure and characteristics of several biotic molecules such as proteins [24]. However, in some cases, irreversible interactions may occur [25]. These interactions, both covalent and non-covalent, can modify those of polyphenols and proteins. This asset can justify the supposed virtues of polyphenols; these help to fight against certain illnesses. Polyphenols can reduce or prevent the appearance of many mechanisms involved in the genesis or in the cardiovascular or cancerous disease amplification [10, 26]. They associate with the proteins underlying these diseases. Nevertheless, few studies offer quantitative information that facilitates categorizing polyphenol-protein interactions. This research targets identifying and classifying those between two flavan-3-ols and an amino acid.

The first compounds correspond to a subclass of polyphenols. The second constitutes a basic component of proteins. In addition, this work aims to understand the stability of complexes correlated with their reactions. This would provide a more comprehensive database for the study of polyphenol-protein associations. Cat and EpiCat are the two flavan-3-ols used because of their high abundance in the human diet; Ala represents the amino acid [27, 28]. This will give us insight into the interactions of these two flavanols with the latter.

This research optimizes the geometry of the chemical structures studied. A frequency calculation performed in the gas phase follows this operation. Both estimates are made

with the DFT method at the theoretical level B3LYP 6–31 G+ (d, p). These calculations concern the monomers Cat, EpiCat and Ala. They also relate to the Cat-Ala and EpiCat-Ala complexes. They lead to their geometric, energetic, and vibratory parameters.

In addition, a NBO (Natural Bond Orbital) analysis was achieved on the complexes under study. It makes it possible to examine CT (LP transfers) and hybridization states during the interactions. A QTAIM investigation was carried out; It helps to identify their strength and the stability of complexes. Finally, this work implements a NCI (Non-Covalent Interaction) scrutiny. It aims to clarify their nature within these chemical structures.

2. Material and Computational Methods

2.1. Calculation Methods

The GAUSSIAN 09 software was used to optimize the geometries of the molecules [29, 30]. Figure 1 shows the structures obtained from Cat, Epicatechin (Epicat) or Alanine (Ala). Those of the Cat... Ala and EpiCat... Ala (Epicat... Ala) complexes appear respectively in figures 2 and 3. These complexes result from the interactions between the hydroxyl groups of the Cat or EpiCat and Ala heteroatoms. The first reactants designate O₂₈-H₂₉, O₃₂-H₃₃, O₃₄-H₃₅. The second ones refer to Nsp³, Osp³ and Osp² [19]. The AIMALL software exploits the "formatted checkpoint file" of geometrical optimizations for topological exams of electron density [31]. The NBO analysis is carried out based on a "single point" calculation from these structures obtained [30, 32]. Multiwfn and Chemcraft software were used to perform respectively the NCI assessment and the surface representation of this latter [33, 34].

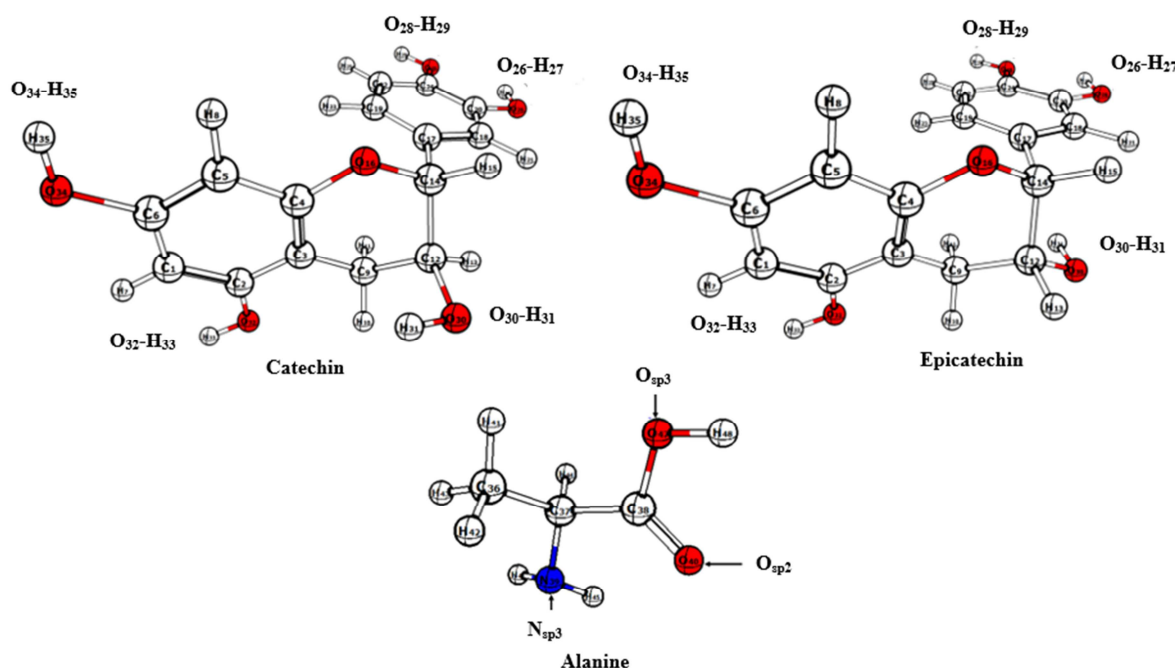


Figure 1. Structures of Cat, EpiCat and Ala.

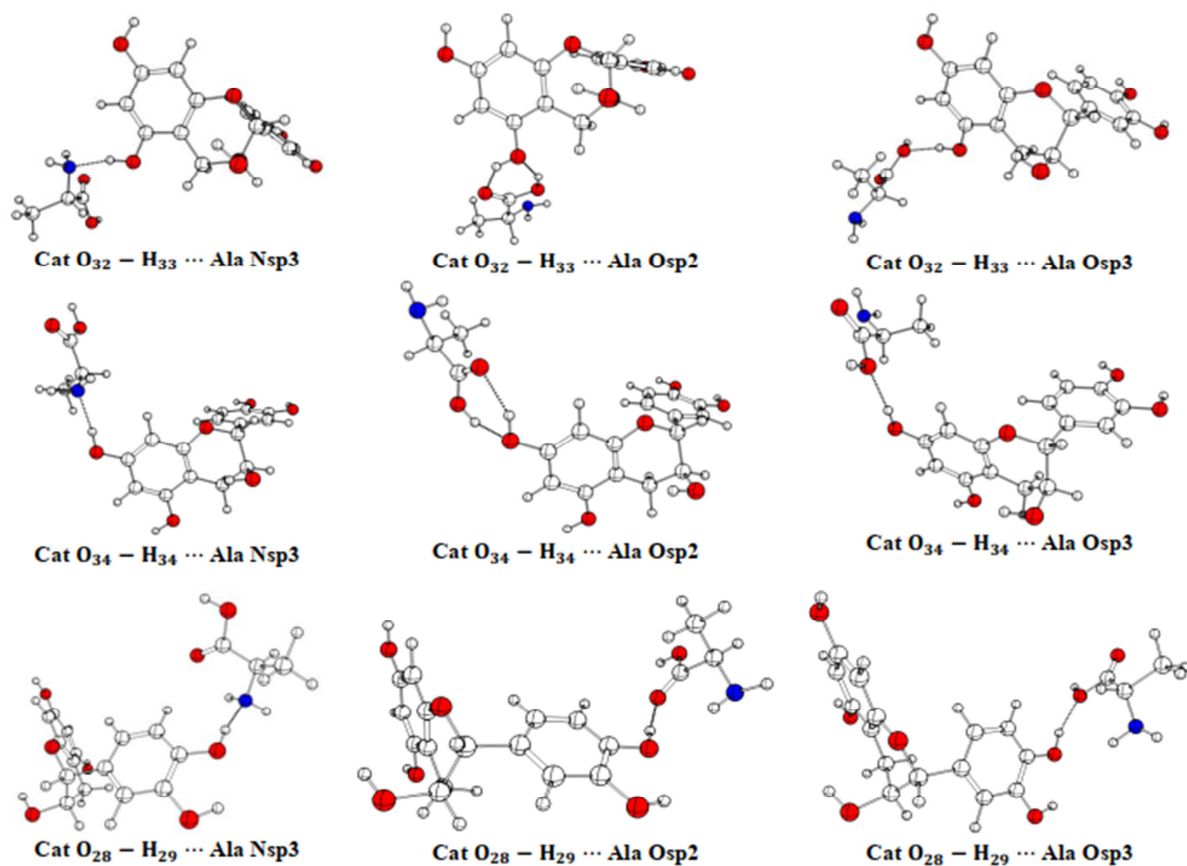


Figure 2. Cat... Ala complexes.

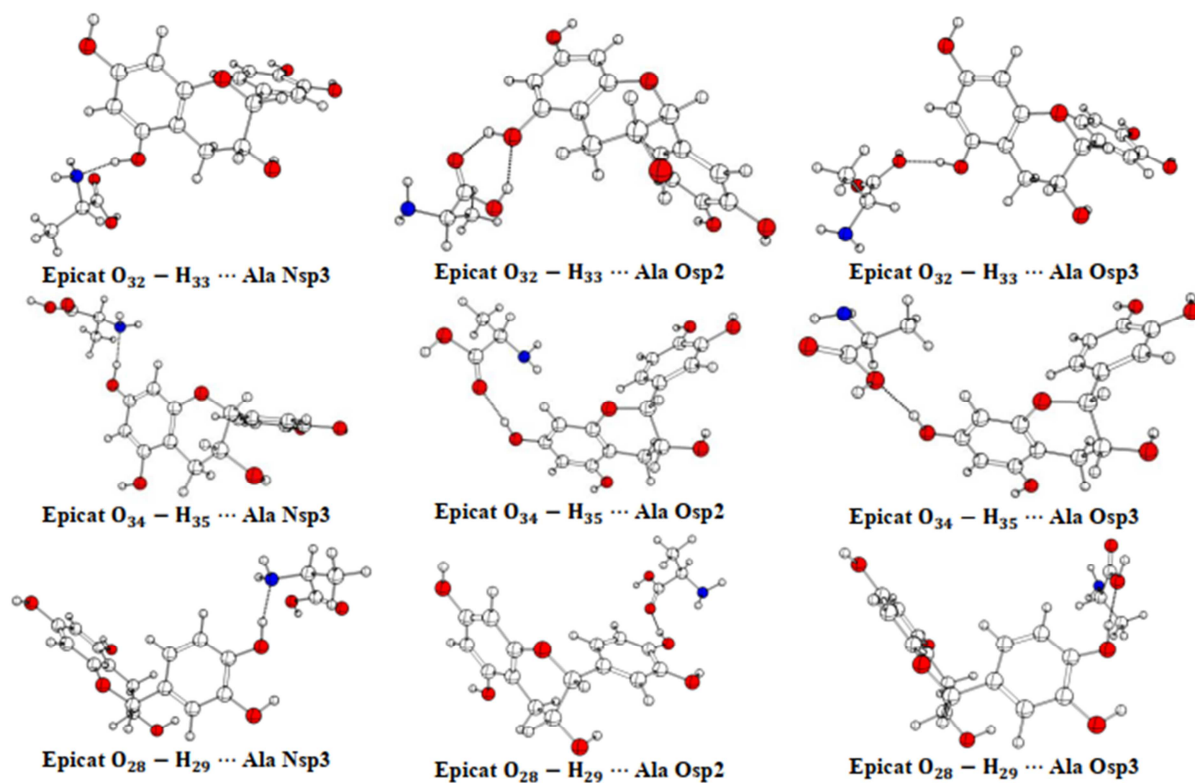


Figure 3. EpiCat... Ala complexes.

2.2. Interaction Energy

The interaction energy E_{int} is the difference between that of the entire complex and those of all its monomers; each molecular orbital is extended to a larger base than those of these. This results in excessive stabilization of this chemical structure. This energy is therefore revised using the BSSE (Basic Superposition Error) and the ZPE (zero-point energy) [35]. It is calculated from the relationship:

$$E_{\text{int}} = E_{\text{complex}} - \sum E_{\text{monomers}} + E_{\text{BSSE}} + \Delta \text{ZPE} \quad (1)$$

2.3. Nbo Analysis

The NBO method contributes to the study of intra- and intermolecular HB. It helps to clarify CT and interactions in molecular systems [36, 37]. The delocalization of electron density between the occupied bonds or lone pairs NBO orbitals and those unoccupied and antibonding corresponds to a donor-acceptor interaction. Its importance is determined using the stabilization energy $E^{(2)}$ whose expression is:

$$E^{(2)} = \Delta E_{ij} = q_i \frac{F^2(i, j)}{\epsilon_j - \epsilon_i} \quad (2)$$

$E^{(2)}$ measures the strength of the donor-acceptor interaction between orbitals ϕ_i and ϕ_j . The higher its value, the stronger the interaction. The second-order perturbation theory allows calculating the delocalization energy [38]. This study only takes into account $E^{(2)}$ greater than 0.15 kcal/mol.

2.4. AIM Analysis

Bader quantum theory or QTAIM is used for the topological analysis of chemical bonds [39]. In this theory, their presence between two atoms or interatomic interactions is conditioned by BCP (Bond Critical Points). The characteristics of the latter make it possible to describe the covalent or non-covalent quality of the HB. The electron density $\rho(r)$ and its Laplacian $\nabla^2 \rho(r)$, the total electron energy density $H(r)$, the electron potential energy $V(r)$ and the kinetic energy $G(r)$ are commonly employed. These parameters allow categorizing the HB by their strengths. The work of Rozas et al. [40], at distinct BCP, suggests the following classification: for strong HB $\nabla^2(\rho) < 0$, $H(r) < 0$; for medium ones $\nabla^2(\rho) > 0$, $H(r) < 0$; for weak ones $\nabla^2(\rho) > 0$, $H(r) > 0$. The ratio $-G(r)/V(r)$ is also used to determine the nature of the interaction at different BCP. If $-G(r)/V(r) > 1$, the bond is considered non-covalent [40, 41]. If $0.5 < -G(r)/V(r) < 1$, it becomes partial. If $-G(r)/V(r) < 0.5$, it is in a closed shell. The energy of the HB is evaluated with the relation $E_{\text{HB}} = -1/2 V(r)$ according to the work of Espinosa [42].

2.5. NCI Analysis

The NCI analysis is a theoretical method to specify them. This specification is a function of the $(\text{sign } \lambda_2) \times \rho(r)$. The factor λ_2 characterizes the variation of the electron density in a perpendicular plane to the internuclear axis.

$P(r)$ represents this quantity along the latter. It distinguishes HB from repulsive interactions (steric interactions) and Van Der Waal (VDW) ones through a 3D graphical visualization of "isosurfaces" [43–45]. NCI analysis associates non-covalent ones with the colours blue, green, and red. These correspond respectively to HB, VDW and repellent bonds. The 2D graphical visualization of the reduced density gradient as a function $(\text{sign } \lambda_2) \times \rho(r)$ allows us to correlate each type of interaction with the region where it appears [43–45]. Thus, the peaks linked to a negative sign of $\lambda_2 \times \rho(r)$ are HB. Those at its zero value corresponding to VDW interactions. Those connected to its positive sign match with their repulsive nature.

3. Results and Discussion

3.1. Thermodynamic Parameters and Interaction Energies

Table 1 shows the energy parameters of the Cat... Ala and EpiCat... Ala. The formations of the Cat... Ala and EpiCat... Ala are in all cases spontaneous and exothermic. The complexation ΔH (enthalpies) and ΔG (free enthalpies) are all negative. The complex $\text{CatO}_i\text{-H}_j\cdots\text{N}_{\text{sp}^3}$, $\text{CatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^2}$, $\text{EpicatO}_i\text{-H}_j\cdots\text{N}_{\text{sp}^3}$ and $\text{EpicatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^2}$ exhibit relatively lower ΔH and ΔG values than the $\text{CatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^3}$ and $\text{EpicatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^3}$ complexes. Thus the N_{sp^3} and O_{sp^2} heteroatoms of Ala give more stable complexes than the O_{sp^3} heteroatom.

Interaction energies E_{int} of the complexes are all negative. For those with EpiCat... Ala (Table 1), they range from -20.60 kcal/mole to -14.56 kcal/mole for the Cat... Ala and from -19.78 kcal/mole to -14.48 kcal/mole. Similar to the complexation ΔH and ΔG , interaction energies of the complex $\text{CatO}_i\text{-H}_j\cdots\text{N}_{\text{sp}^3}$, $\text{CatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^2}$, $\text{EpicatO}_i\text{-H}_j\cdots\text{N}_{\text{sp}^3}$ and $\text{EpicatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^2}$ are lower compared with those of the $\text{CatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^3}$ and $\text{EpicatO}_i\text{-H}_j\cdots\text{O}_{\text{sp}^3}$ complexes. Thus, the N_{sp^3} and O_{sp^2} sites of Ala form more stable complexes than O_{sp^3} . Among complexes formed with the N_{sp^3} site, order is $E_{\text{int}}(\text{CatO}_{34}\text{-H}_{35}\cdots\text{N}_{\text{sp}^3}) < E_{\text{int}}(\text{CatO}_{28}\text{-H}_{29}\cdots\text{N}_{\text{sp}^3}) < E_{\text{int}}(\text{CatO}_{32}\text{-H}_{33}\cdots\text{N}_{\text{sp}^3})$ for Cat and $E_{\text{int}}(\text{EpicatO}_{32}\text{-H}_{33}\cdots\text{N}_{\text{sp}^3}) < E_{\text{int}}(\text{EpicatO}_{28}\text{-H}_{29}\cdots\text{N}_{\text{sp}^3}) < E_{\text{int}}(\text{EpicatO}_{34}\text{-H}_{35}\cdots\text{N}_{\text{sp}^3})$ for EpiCat. With O_{sp^2} , ranking of interaction energies in ascending order gives: $E_{\text{int}}(\text{CatO}_{34}\text{-H}_{35}\cdots\text{O}_{\text{sp}^2}) < E_{\text{int}}(\text{CatO}_{32}\text{-H}_{33}\cdots\text{O}_{\text{sp}^2}) < E_{\text{int}}(\text{CatO}_{28}\text{-H}_{29}\cdots\text{O}_{\text{sp}^2})$ for Cat and $E_{\text{int}}(\text{EpicatO}_{32}\text{-H}_{33}\cdots\text{O}_{\text{sp}^2}) < E_{\text{int}}(\text{EpicatO}_{28}\text{-H}_{29}\cdots\text{O}_{\text{sp}^2}) < E_{\text{int}}(\text{EpicatO}_{34}\text{-H}_{35}\cdots\text{O}_{\text{sp}^2})$ for EpiCat are cited in the text.

From the above analysis, it appears that regarding Cat... Ala complexes, $\text{CatO}_{28}\text{-H}_{29}\cdots\text{N}_{\text{sp}^3}$, $\text{CatO}_{32}\text{-H}_{33}\cdots\text{N}_{\text{sp}^3}$, $\text{CatO}_{34}\text{-H}_{35}\cdots\text{N}_{\text{sp}^3}$, $\text{CatO}_{28}\text{-H}_{29}\cdots\text{O}_{\text{sp}^2}$, $\text{CatO}_{32}\text{-H}_{33}\cdots\text{O}_{\text{sp}^2}$ and $\text{CatO}_{34}\text{-H}_{35}\cdots\text{O}_{\text{sp}^2}$ are more stable. While for EpiCat... Ala complexes, $\text{EpicatO}_{28}\text{-H}_{29}\cdots\text{N}_{\text{sp}^3}$, $\text{EpicatO}_{32}\text{-H}_{33}\cdots\text{N}_{\text{sp}^3}$,

EpicatO₃₄-H₃₅...N_{sp}³, EpicatO₂₈-H₂₉...O_{sp}², EpicatO₃₂-H₃₃...O_{sp}² and EpicatO₃₄-H₃₅...O_{sp}² are the most

stable. NBO and AIM analyses are performed to identify the nature of the interactions that occur in these complexes.

Table 1. Energy parameters in kcal/mol of EpiCat... Ala and Cat... Ala complexes.

Complexes EpiCat ... Ala	ΔH (kcal/mol)	ΔG	E_{int}	Complex Cat... Ala	ΔH (kcal/mol)	ΔG	E_{int}
EpicatO ₂₈ -H ₂₉ ...N _{sp} ³	-20.12	-10.99	-19.30	CatO ₂₈ -H ₂₉ ...N _{sp} ³	-21.21	-11.08	-20.32
EpicatO ₂₈ -H ₂₉ ...O _{sp} ²	-18.73	-9.19	-18.36	CatO ₂₈ -H ₂₉ ...O _{sp} ²	-18.63	-8.94	-18.27
EpicatO ₂₈ -H ₂₉ ...O _{sp} ³	-14.88	-6.78	-14.84	CatO ₂₈ -H ₂₉ ...O _{sp} ³	-14.87	-5.92	-14.56
EpicatO ₃₂ -H ₃₃ ...N _{sp} ³	-20.22	-11.10	-19.62	CatO ₃₂ -H ₃₃ ...N _{sp} ³	-20.35	-11.31	-19.76
EpicatO ₃₂ -H ₃₃ ...O _{sp} ²	-20.08	-10.21	-19.67	CatO ₃₂ -H ₃₃ ...O _{sp} ²	-19.77	-9.90	-19.34
EpicatO ₃₂ -H ₃₃ ...O _{sp} ³	-16.22	-7.98	-16.25	CatO ₃₂ -H ₃₃ ...O _{sp} ³	-14.70	-6.22	-14.64
EpicatO ₃₄ -H ₃₅ ...N _{sp} ³	-19.53	-10.01	-18.85	CatO ₃₄ -H ₃₅ ...N _{sp} ³	-21.20	-11.98	-20.60
EpicatO ₃₄ -H ₃₅ ...O _{sp} ²	-16.65	-7.44	-16.48	CatO ₃₄ -H ₃₅ ...O _{sp} ²	-20.29	-9.97	-19.84
EpicatO ₃₄ -H ₃₅ ...O _{sp} ³	-14.49	-6.35	-14.48	CatO ₃₄ -H ₃₅ ...O _{sp} ³	-14.81	-6.27	-14.85

3.2. NBO Analysis

Tables 2 to 10 provide the electronic transitions between NBO donor and acceptor sites, stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$, and CT of associated with complexes resulting from the interactions between O₂₈-H₂₉, O₃₂-H₃₃, O₃₄-H₃₅ of Cat and EpiCat with N_{sp}³, O_{sp}² and O_{sp}³ of Ala NBO analysis allows us to specify, in the complexes, all interactions between Cat and Ala, on the one hand, and then between EpiCat and Ala on the other. X - H ... Y is characterized by electronic transitions $LP_Y \rightarrow \sigma_{X-H}^*$ in which σ_{X-H}^* acts as an electron acceptor and LP_Y (Y's lone pair) as an electron donor. These interactions are all defined by a second-order perturbation energy $E^{(2)}$ and a CT. In EpicatO₂₈-H₂₉...Ala N_{sp}³ complex (Table 2), the total stabilization energy is $\sum E^2 = 31.09$ kcal/mol. In this complex, we note the presence of the O₂₈-H₂₉...N₃₉ HB governed by $LP_{N39}^{(1)} \rightarrow \sigma_{O28-H29}^*$ interaction. This interaction results in electron transfer from the lone pair of the nitrogen atom ($LP_{N39}^{(1)}$) to the antibonding orbital $\sigma_{O28-H29}^*$ with a

stabilization energy $E^2 = 29.9$ kcal/mol and a CT of 56.64 me. O₂₈-H₂₉...N₃₉ HB contributes 96.2% of the Epicat's total stabilization energy O₂₈-H₂₉...Ala N_{sp}³ complex. Hence, the HB O₂₈ - H₂₉ ... N₃₉ is considered as the main interaction of the Epicat O₂₈-H₂₉...Ala N_{sp}³ complex. Figure 4 illustrates $LP_{N39}^{(1)} \rightarrow \sigma_{O28-H29}^*$ natural bonding orbital of this main interaction. Besides the latter, the NBO analysis reveals the presence of others with low stabilization energies, including C₃₆-H₄₂...O₂₈ ($E^2 = 0.09$ kcal/mol, CT = 0.16 me), O₂₈-H₂₉...H₄₅-N₃₉ ($E^2 = 0.62$ kcal/mol, CT = 0.86 me), O₂₈-H₂₉...N₃₉-C₃₇ ($E^2 = 0.36$ kcal/mol, CT = 0.45 me),

The total stabilization energy of the Cat O₂₈-H₂₉...Ala N_{sp}³ complex is $\sum E^2 = 30.53$ kcal/mol (Table 2). The significant intermolecular interactions that occur are: O₂₈-H₂₉...N₃₉, C₂₂-H₂₅...O₄₀, C₃₆-H₄₂...O₂₆, C₂₂-H₂₅...C₃₈=O₄₀, O₂₈-H₂₉...H₄₅-N₃₉ and O₂₈-H₂₉...H₄₆-N₃₉. That of O₂₈-H₂₉...N₃₉ is 27.94 kcal/mol. This energy stabilization energy of the Cat O₂₈-H₂₉...Ala N_{sp}³ complex accounts for 91.5% of its total value. O₂₈-H₂₉...N₃₉ is therefore the main interaction in this complex.

Table 2. Stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$ and CT of Epicat O₂₈-H₂₉...Ala N_{sp}³ and Cat O₂₈-H₂₉...Ala N_{sp}³.

Contact	electronic transitions	$E^{(2)}$ (kcal/mol)	$\sum E^{(2)}$ (kcal/mol)	CT (me)
Epicat O ₂₈ -H ₂₉ ...Ala N _{sp} ³				
O ₂₈ -H ₂₉ ...N ₃₉	$LP_{N39}^{(1)} \rightarrow \sigma_{O28-H29}^*$	29.9		56.64
C ₃₈ -H ₄₁ ...O ₂₈	$LP_{O28}^{(1)} \rightarrow \sigma_{C38-H41}^*$	0.09		0.16
O ₂₈ -H ₂₉ ...H ₄₄ -C ₃₇	$\sigma_{C37-H44}^{(1)} \rightarrow \sigma_{O28-H29}^*$	0.12	31.09	0.20
O ₂₈ -H ₂₉ ...N ₃₉ -C ₃₇	$\sigma_{C37-N39}^{(1)} \rightarrow \sigma_{O28-H29}^*$	0.36		0.45
O ₂₈ -H ₂₉ ...H ₄₅ -N ₃₉	$\sigma_{N39-H45}^{(1)} \rightarrow \sigma_{O28-H29}^*$	0.62		0.86
Cat O ₂₈ -H ₂₉ ...Ala N _{sp} ³				
O ₂₈ - H ₂₉ ... N ₃₉	$LP_{N39}^{(1)} \rightarrow \sigma_{O28-H29}^*$	27.84		54.21
	$LP_{O40}^{(1)} \rightarrow \sigma_{C22-H25}^*$	0.44		0.57
C ₂₂ -H ₂₅ ...O ₄₀	$LP_{O40}^{(2)} \rightarrow \sigma_{C22-H25}^*$	0.29		0.61
C ₂₂ -H ₂₅ ...O ₄₀ -C ₃₈	$\pi_{C38-O40}^{(2)} \rightarrow \sigma_{C22-H25}^*$	0.35	30.53	0.60
O ₂₈ -H ₂₉ ...N ₃₉ -C ₃₇	$\sigma_{C37-N39}^{(1)} \rightarrow \sigma_{O28-H29}^*$	0.27		0.35
O ₂₈ -H ₂₉ ...H ₄₂ -C ₃₆	$\sigma_{C36-H42}^{(1)} \rightarrow \sigma_{O28-H29}^*$	0.1		0.17

Contact	electronic transitions	$E^{(2)}$ (kcal/mol)	$\sum E^{(2)}$ (kcal/mol)	CT (me)
$O_{28}-H_{29}\cdots H_{45}-N_{39}$	$\sigma_{N_{39}-H_{44}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.57		0.84
$O_{28}-H_{29}\cdots H_{46}-N_{39}$	$\sigma_{N_{39}-H_{45}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.67		0.92

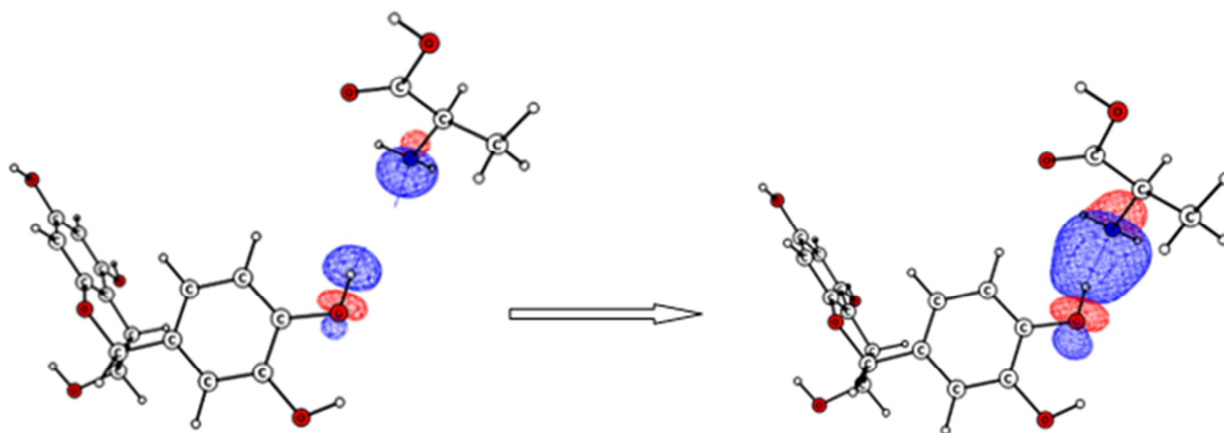


Figure 4. Natural bond orbital of $LP_{N_{39}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ in $O_{28}-H_{29}\cdots N_{39}$ interaction.

In the Epicat $O_{28}-H_{29}\cdots Ala O_{sp^2}$ complex ($\sum E^2 = 23.97$ kcal/mol) (Table 3), the interactions that occur are: $O_{28}-H_{29}\cdots O_{40}$, $N_{39}-H_{46}\cdots O_{28}$, $C_{38}-O_{40}\cdots O_{28}$, $N_{39}-H_{46}\cdots H_{29}-O_{28}$, $C_{22}-H_{25}\cdots O_{40}$, $O_{28}-H_{29}\cdots O_{40}-C_{38}$ and $C_{22}-H_{25}\cdots O_{40}-C_{38}$. $O_{28}-H_{29}\cdots O_{40}$ is governed by two electronic relocations. These are delocalization of electrons $LP_{O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ ($E^2 = 8.70$ kcal/mol, CT = 11.31 me) which result in their transfers from the first lone pair denoted $LP_{O_{40}}^{(1)}$ of the oxygen to the antibonding orbital $\sigma_{O_{28}-H_{29}}^*$. It contributes 66.3% of the Epicat's total stabilization energy $O_{28}-H_{29}\cdots Ala O_{sp^2}$ complex. The electronic transition $LP_{O_{40}}^{(2)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ ($E^2 = 11.51$ kcal/mol, CT = 23.11 me) results in the electron transfer from the second free lone pair denoted

($LP_{O_{40}}^{(2)}$) of the oxygen atom to the antibonding orbital $\sigma_{O_{28}-H_{29}}^*$. It contributes 48% of the total stabilization energy. Thus, $O_{28}-H_{29}\cdots O_{40}$ contributes 84.3% of the total stabilization energy. Figure 5 provides the natural bond of the electronic transitions $LP_{O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ and $LP_{O_{40}}^{(2)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ of $O_{28}-H_{29}\cdots O_{40}$. In Cat $O_{28}-H_{29}\cdots Ala O_{sp^2}$ complex whose total stabilization energy is $\sum E^2 = 23.09$ kcal/mol (Table 3), the interactions that occur are: $O_{28}-H_{29}\cdots O_{40}$, $N_{39}-H_{46}\cdots O_{28}$, $C_{22}-H_{25}\cdots O_{40}$, $O_{28}-H_{29}\cdots O_{40}-C_{38}$ et $C_{22}-H_{25}\cdots O_{40}-C_{38}$. The $O_{28}-H_{29}\cdots O_{40}$ is the main interaction in this complex; its stabilization energy equals 19.83 kcal/mol, represents 84.2% of the Cat $O_{28}-H_{29}\cdots Ala O_{sp^2}$ complex's total value.

Table 3. Stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$ and CT of Epicat $O_{28}-H_{29}\cdots Ala O_{sp^2}$ and Cat $O_{28}-H_{29}\cdots Ala O_{sp^2}$.

b	electronic transitions	$E^{(2)}$ (kcal/mol)	$\sum E^{(2)}$ (kcal/mol)	CT (me)
Epicat $O_{28}-H_{29}\cdots Ala O_{sp^2}$				
$O_{28}-H_{29}\cdots O_{40}$	$LP_{O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	8.7		11.31
	$LP_{O_{40}}^{(2)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	11.51		23.11
$N_{39}-H_{46}\cdots O_{28}$	$LP_{O_{28}}^{(1)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	0.64		0.92
	$LP_{O_{28}}^{(2)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	1.81		3.52
$C_{38}-O_{40}\cdots O_{28}$	$LP_{O_{28}}^{(1)} \rightarrow \pi_{C_{38}-O_{40}}^*$	0.1		0.16
$N_{39}-H_{46}\cdots H_{29}-O_{28}$	$\sigma_{O_{28}-H_{29}}^{(1)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	0.28	23.97	0.37
$C_{22}-H_{25}\cdots O_{40}$	$LP_{O_{40}}^{(1)} \rightarrow \sigma_{C_{22}-H_{25}}^*$	0.13		0.17
$O_{28}-H_{29}\cdots O_{40}-C_{38}$	$\sigma_{C_{38}-O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.38		0.38
$C_{22}-H_{25}\cdots O_{40}-C_{38}$	$\pi_{C_{38}-O_{40}}^{(2)} \rightarrow \sigma_{C_{22}-H_{25}}^*$	0.12		0.20
$O_{28}-H_{29}\cdots O_{47}-C_{38}$	$\sigma_{C_{38}-O_{47}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.18		0.19
$O_{28}-H_{29}\cdots H_{46}-N_{39}$	$\sigma_{N_{39}-H_{46}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.12		0.16
Cat $O_{28}-H_{29}\cdots Ala O_{sp^2}$				
$O_{28}-H_{29}\cdots O_{40}$	$LP_{O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	8.58		11.31
	$LP_{O_{40}}^{(2)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	11.25	23.54	22.58
$N_{39}-H_{45}\cdots O_{28}$	$LP_{O_{28}}^{(1)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	0.67		1.01
	$LP_{O_{28}}^{(2)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	1.88		3.73

b	electronic transitions	$E^{(2)}$(kcal/mol)	$\sum E^{(2)}$(kcal/mol)	CT (me)
$C_{38}-O_{40}\cdots O_{28}$	$LP_{O_{28}}^{(1)} \rightarrow \pi_{C_{38}-O_{40}}^*$	0.1		0.16
$N_{39}-H_{45}\cdots H_{29}-O_{28}$	$\sigma_{O_{28}-H_{29}}^{(1)} \rightarrow \sigma_{N_{39}-H_{45}}^*$	0.29		0.37
$C_{22}-H_{25}\cdots O_{40}$	$LP_{O_{40}}^{(1)} \rightarrow \sigma_{C_{22}-H_{25}}^*$	0.12		0.17
$O_{28}-H_{29}\cdots O_{40}-C_{38}$	$\sigma_{C_{38}-O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.37		0.38
$O_{28}-H_{29}\cdots O_{40}-C_{38}$	$\pi_{C_{38}-O_{40}}^{(2)} \rightarrow \sigma_{C_{22}-H_{25}}^*$	0.11		0.20
$O_{28}-H_{29}\cdots O_{47}-C_{38}$	$\sigma_{C_{38}-O_{47}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.17		0.19

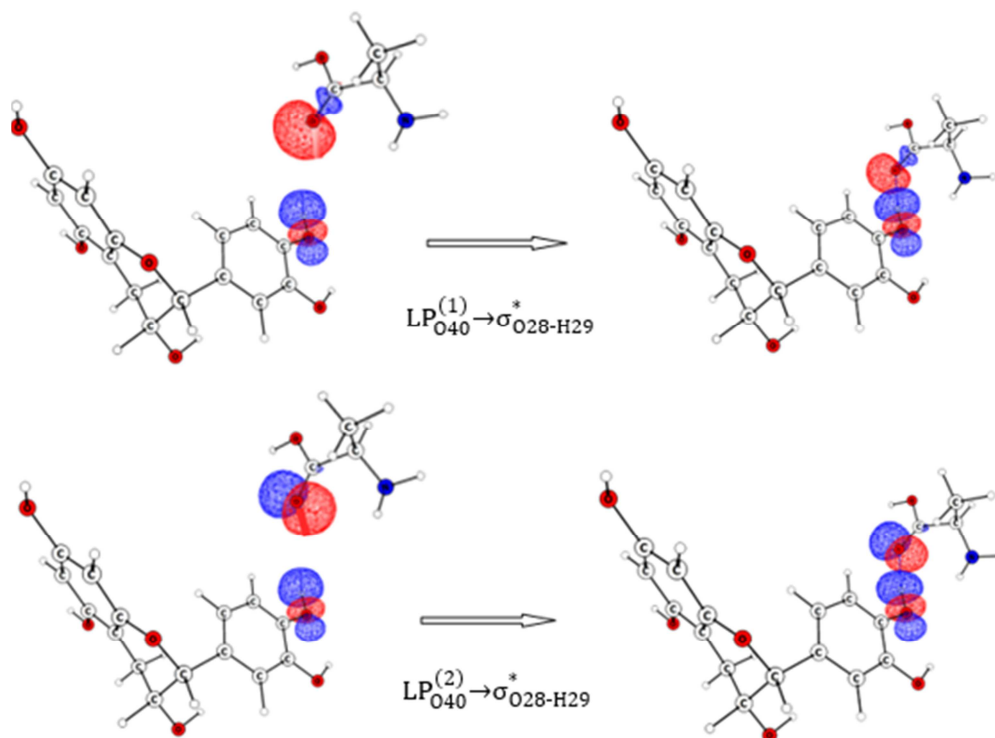


Figure 5. Natural bond orbital of $LP_{O_{40}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ and $LP_{O_{40}}^{(2)} \rightarrow \sigma_{O_{28}-H_{29}}^*$ in $O_{28}-H_{29}\cdots O_{40}$ interaction.

In Epicat $O_{28}-H_{29}\cdots Ala O_{sp^3}$ ($\sum E^2=11.86$ kcal/mol) and Cat $O_{28}-H_{29}\cdots Ala O_{sp^3}$ ($\sum E^2=14.81$ kcal/mol) complexes (Table 4), the stabilization energy values of $O_{28}-H_{29}\cdots O_{47}$ HB are 10.95 kcal/mol and 11.66 kcal/mol, respectively. $O_{28}-H_{29}\cdots O_{47}$ is the main interaction in Epicat $O_{28}-H_{29}\cdots Ala O_{sp^3}$ and Cat $O_{28}-H_{29}\cdots Ala O_{sp^3}$

complexes. Other interactions involved in these complexes are:

$C_{36}-H_{41}\cdots O_{28}$, $C_{22}-H_{25}\cdots O_{40}$, $C_{38}-O_{47}\cdots H_{29}-O_{28}$. $N_{39}-H_{46}\cdots O_{28}$ is the main interaction in this complex; its stabilization energy, estimated at 19.83 kcal/mol, represents 84.2% of the Cat total value.

Table 4. Stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$ and CT of Epicat $O_{28}-H_{29}\cdots Ala O_{sp^3}$ and Cat $O_{28}-H_{29}\cdots Ala O_{sp^3}$.

Contact	electronic transitions	$E^{(2)}$(kcal/mol)	$\sum E^{(2)}$(kcal/mol)	CT (me)
Epicat $O_{28}-H_{29}\cdots Ala O_{sp^3}$				
$O_{28}-H_{29}\cdots O_{47}$	$LP_{O_{47}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	10.31		14.43
	$LP_{O_{47}}^{(2)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.64		1.22
$C_{36}-H_{41}\cdots O_{28}$	$LP_{O_{28}}^{(2)} \rightarrow \sigma_{C_{36}-H_{41}}^*$	0.37		0.72
$C_{22}-H_{25}\cdots O_{40}$	$LP_{O_{40}}^{(2)} \rightarrow \sigma_{C_{22}-H_{25}}^*$	0.13	11.86	0.22
$C_{38}-H_{47}\cdots H_{29}-O_{28}$	$\sigma_{O_{28}-H_{29}}^{(1)} \rightarrow \sigma_{C_{38}-O_{47}}^*$	0.15		0.21
$O_{28}-H_{29}\cdots H_{41}-C_{36}$	$\sigma_{C_{36}-H_{41}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.14		0.20
$O_{28}-H_{29}\cdots H_{48}-O_{47}$	$\sigma_{O_{47}-H_{48}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	0.12		0.15
Cat $O_{28}-H_{29}\cdots Ala O_{sp^3}$				
$O_{28}-H_{29}\cdots O_{47}$	$LP_{O_{47}}^{(1)} \rightarrow \sigma_{O_{28}-H_{29}}^*$	11.66		16.59
	$LP_{O_{28}}^{(1)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	0.5	14.81	0.74
$N_{39}-H_{46}\cdots O_{28}$	$LP_{O_{28}}^{(2)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	2.18		4.09

Contact	electronic transitions	$E^{(2)}$ (kcal/mol)	$\sum E^{(2)}$ (kcal/mol)	CT (me)
$C_{38}-H_{47}\cdots H_{29}-O_{28}$	$\sigma_{O_{28}-H_{29}}^{(1)} \rightarrow \sigma_{C_{38}-O_{47}}^*$	0.11		0.17
$N_{39}-H_{46}\cdots H_{29}-O_{28}$	$\sigma_{O_{28}-H_{29}}^{(1)} \rightarrow \sigma_{N_{39}-H_{46}}^*$ $\sigma_{O_{28}-H_{29}}^{(1)} \rightarrow \sigma_{N_{39}-H_{46}}^*$	0.26		0.32
$C_{22}-H_{25}\cdots O_{47}$	$LP_{O_{47}}^{(2)} \rightarrow \sigma_{C_{22}-H_{25}}^*$	0.1		0.18

The total stabilization energy of Epicat $O_{32}-H_{33}\cdots Ala N_{sp^3}$ complex is ($\sum E^2=29.76$ kcal/mol) (Table 5). In this complex, the N_{sp^3} nitrogen of Ala establishes a HB with the $O_{32}-H_{33}$ hydroxyl groups of EpiCat. This interaction $O_{32}-H_{33}\cdots N_{39}$ results from electron transfer from the lone pair of the nitrogen atom ($LP_{N_{39}}^{(1)}$) to the $\sigma_{O_{32}-H_{33}}^*$ antibonding orbital which is electron depleted ($LP_{N_{39}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$). This interaction, having energy $E^2=27.32$ kcal/mol, with an estimated CT of 53.92 me, contributes 91.80% of the total stabilization energy. In addition to the $O_{32}-H_{33}\cdots N_{39}$, $C_1-H_7\cdots N_{39}$ ($E^2=0.83$ kcal/mol, CT=0.14 me), $O_{32}-H_{33}\cdots H_{45}-N_{39}$ ($E^2=0.41$ kcal/mol, CT=0.59 me), $O_{32}-H_{33}\cdots C_{37}-N_{39}$ ($E^2=0.45$ kcal/mol, CT=0.60 me), $O_{32}-H_{33}\cdots H_{46}-N_{39}$ ($E^2=0.61$ kcal/mol, CT=0.84 me).

($E^{(2)} = 0.45$ kcal/mol, CT=0.60 me), $O_{32}-H_{33}\cdots H_{46}-N_{39}$ ($E^2=0.61$ kcal/mol, CT=0.84 me) the interactions contribute to the stabilization of the complex with much lower energies. The essentials for Epicat $O_{32}-H_{33}\cdots Ala N_{sp^3}$ is then the $O_{32}-H_{33}\cdots N_{39}$. For Cat $O_{32}-H_{33}\cdots Ala N_{sp^3}$, it is the $O_{32}-H_{33}\cdots N_{39}$ HB (Table 5). Indeed, the stabilization energy of the $O_{32}-H_{33}\cdots N_{39}$ is equal to 27.61 kcal/mol. It represents 94.6% of the Cat total stabilization energy $O_{32}-H_{33}\cdots Ala N_{sp^3}$ complex ($\sum E^2=29.19$ kcal/mol). The other interactions that occur in this complex are $O_{32}-H_{33}\cdots H_{45}-N_{39}$, $O_{32}-H_{33}\cdots H_{46}-N_{39}$, $O_{32}-H_{33}\cdots C_{37}-N_{39}$ and $C_1-H_7\cdots N_{39}$. Their stabilization energies are weak.

Table 5. Stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$ and CT of Epicat $O_{32}-H_{33}\cdots Ala N_{sp^3}$ and Cat $O_{32}-H_{33}\cdots Ala N_{sp^3}$ complexes.

Epicat $O_{32}-H_{33}\cdots Ala N_{sp^3}$				
Contact	electronic transitions	$E^{(2)}$ (kcal/mol)	$\sum E^{(2)}$ (kcal/mol)	CT (me)
$O_{32}-H_{33}\cdots N_{39}$	$LP_{N_{39}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	27.32		53.92
$C_1-H_7\cdots N_{39}$	$LP_{N_{39}}^{(1)} \rightarrow \sigma_{C_1-H_7}^*$	0.83		0.14
$O_{32}-H_{33}\cdots H_{45}-N_{39}$	$\sigma_{N_{39}-H_{45}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	0.41	29.76	0.59
$O_{32}-H_{33}\cdots C_{37}-N_{39}$	$\sigma_{N_{39}-C_{37}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	0.45		0.60
$O_{32}-H_{33}\cdots H_{46}-N_{39}$	$\sigma_{N_{39}-H_{46}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	0.61		0.84

Cat $O_{32}-H_{33}\cdots Ala N_{sp^3}$				
Contact	electronic transitions	$E^{(2)}$ (kcal/mol)	$\sum E^{(2)}$ (kcal/mol)	CT (me)
$O_{32}-H_{33}\cdots N_{39}$	$LP_{N_{39}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	27.61		53.92
$C_1-H_7\cdots N_{39}$	$LP_{N_{39}}^{(1)} \rightarrow \sigma_{C_1-H_7}^*$	0.07		0.14
$O_{32}-H_{33}\cdots H_{45}-N_{39}$	$\sigma_{N_{39}-H_{45}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	0.42	29.19	0.59
$O_{32}-H_{33}\cdots C_{37}-N_{39}$	$\sigma_{N_{39}-C_{37}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	0.48		0.60
$O_{32}-H_{33}\cdots H_{46}-N_{39}$	$\sigma_{N_{39}-H_{46}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$	0.61		0.84

In Epicat $O_{32}-H_{33}\cdots Ala O_{sp^2}$ complex ($\sum E^2=29.73$ kcal/mol) appears two HB $O_{32}-H_{33}\cdots O_{40}$ and $O_{47}-H_{48}\cdots O_{32}$ with high stabilization energies (Table 6). $O_{32}-H_{33}\cdots O_{40}$ HB ($E^2=13.73$ kcal/mol) results in two specific electronic transitions namely $LP_{O_{40}}^{(1)} \rightarrow \sigma_{O_{32}-H_{33}}^*$ ($E^2=4.45$ kcal/mol, CT= 5.98 me) and $LP_{O_{40}}^{(2)} \rightarrow \sigma_{O_{32}-H_{33}}^*$ ($E^2=9.28$ kcal/mol, CT= 20.00 me), contributing 46.2% of the total stabilization energy. Whereas, $O_{47}-H_{48}\cdots O_{32}$ ($E^{(2)} = 14.23$ kcal/mol), contributes 47.9% to the total stabilization energy and is governed by electron delocalization $LP_{O_{32}}^{(1)} \rightarrow \sigma_{O_{47}-H_{48}}^*$ ($E^2=6.10$ kcal/mol, CT= 9.61 me) and $LP_{O_{32}}^{(2)} \rightarrow \sigma_{O_{47}-H_{48}}^*$ ($E^2=8.13$ kcal/mol, CT= 17.00 me). These two HB $O_{32}-H_{33}\cdots O_{40}$ and $O_{47}-H_{48}\cdots O_{32}$ constitute the main interactions of Epicat $O_{32}-H_{33}\cdots Ala O_{sp^2}$ complex. However,

it is important to note the presence of other interactions including $O_{47}-H_{48}\cdots C_2-C_1$, $O_{47}-H_{48}\cdots H_{33}-O_{32}$, $O_{32}-H_{33}\cdots O_{40}-C_{38}$. The total stabilization energy of the Cat $O_{32}-H_{33}\cdots Ala O_{sp^2}$ complex is $\sum E^2=30.63$ kcal/mol (Table 6). The intermolecular interactions that most stabilize this complex are: $O_{32}-H_{33}\cdots O_{40}$ and $O_{47}-H_{48}\cdots O_{32}$. Their stabilization energies are 14.43 kcal/mol and 14.24 kcal/mol, respectively.

The total stabilization energies of Epicat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ and Cat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ complexes are 10.71 kcal/mol and 10.93 kcal/mol respectively (Table 7). The interactions that occur in these complexes are $O_{32}-H_{33}\cdots O_{47}$, $C_{37}-H_{44}\cdots O_{32}$, $O_{36}-H_{43}\cdots C_2-C_1$, $C_1-H_7\cdots O_{47}$ and $O_{32}-H_{33}\cdots H_{48}-O_{47}$. In Epicat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ complex, the stabilization energy

of $O_{32}-H_{33}\cdots O_{47}$ is 10.2 kcal/mol and represents 95.3% of $\sum E^{(2)}$. This energy is equal to 10.52 kcal/mol and represents 96.2% of $\sum E^{(2)}$ in Cat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ complex. Thus, the main interaction of Epicat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ and Cat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ complexes is $O_{32}-H_{33}\cdots O_{47}$.

In Epicat $O_{34}-H_{35}\cdots Ala N_{sp^3}$ complex ($\sum E^2=27.31$ kcal/mol) (Table 8), we note the presence of the $O_{34}-H_{35}\cdots N_{39}$, $C_{36}-H_{42}\cdots O_{34}$, $C_5-H_8\cdots N_{39}$ HB and other interactions with low stabilization energies. $O_{34}-H_{35}\cdots N_{39}$

HB is resulted in the electronic transition $LP_{N_{39}}^{(1)} \rightarrow \sigma_{O_{34}-H_{35}}^*$ ($E^{(2)}=25.64$ kcal/mol, CT= 50.73 me). It accounts for 93.9% of the total stabilization energy. $O_{34}-H_{35}\cdots N_{39}$ is considered the main interaction of the Epicat $O_{34}-H_{35}\cdots Ala N_{sp^3}$ complex. The total stabilization energy of the Cat $O_{34}-H_{35}\cdots Ala N_{sp^3}$ complex is $\sum E^2=28.93$ kcal/mol. The intermolecular interactions that stabilize this complex most strongly is $O_{34}-H_{35}\cdots N_{39}$. His stabilization energy is worth 27.14 kcal/mol.

Table 6. Stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$ and CT of Epicat $O_{32}-H_{33}\cdots Ala O_{sp^2}$ and Cat $O_{32}-H_{33}\cdots Ala O_{sp^2}$.

Epicat O ₃₂ -H ₃₃ ⋯Ala O _{sp} ²				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₄₇ -H ₄₈ ⋯O ₃₂	LP _{O₃₂} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	6.1	29.73	9.61
	LP _{O₃₂} ⁽²⁾ →σ _{O₄₇-H₄₈} [*]	8.13		17.00
O ₃₂ -H ₃₃ ⋯O ₄₀	LP _{O₄₀} ⁽¹⁾ →σ _{O₃₂-H₃₃} [*]	4.45		5.98
	LP _{O₄₀} ⁽²⁾ →σ _{O₃₂-H₃₃} [*]	9.28		20.00
O ₄₇ -H ₄₈ ⋯C ₂ -C ₁	σ _{C₁-C₂} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	0.11		0.15
O ₄₇ -H ₄₈ ⋯H ₃₃ -O ₃₂	σ _{O₃₂-H₃₃} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	1.06		1.42
O ₃₂ -H ₃₃ ⋯C ₃₈ -C ₃₇	σ _{C₃₇-C₃₈} ⁽¹⁾ →σ _{O₃₂-H₃₃} [*]	0.19		0.28
O ₃₂ -H ₃₃ ⋯O ₄₀ -C ₃₈	σ _{C₃₈-O₄₀} ⁽¹⁾ →σ _{O₃₂-H₃₃} [*]	0.23		0.25
C ₂ -O ₃₂ ⋯H ₄₈ -O ₄₇	σ _{O₄₇-H₄₈} ⁽¹⁾ →σ _{C₂-O₃₂} [*]	0.18		0.27
Cat O ₃₂ -H ₃₃ ⋯Ala O _{sp} ²				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₄₇ -H ₄₈ ⋯O ₃₂	LP _{O₃₂} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	6.63	30.63	10.24
	LP _{O₃₂} ⁽²⁾ →σ _{O₄₇-H₄₈} [*]	7.61		15.60
O ₃₂ -H ₃₃ ⋯O ₄₀	LP _{O₄₀} ⁽¹⁾ →σ _{O₃₂-H₃₃} [*]	4.56		6.28
	LP _{O₄₀} ⁽²⁾ →σ _{O₃₂-H₃₃} [*]	9.87		21.10
O ₄₇ -H ₄₈ ⋯C ₂ -C ₁	σ _{C₁-C₂} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	0.11		0.15
O ₄₇ -H ₄₈ ⋯H ₃₃ -O ₃₂	σ _{O₃₂-H₃₃} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	1.1		1.42
C ₂ -O ₃₂ ⋯O ₄₀	LP _{O₄₀} ⁽²⁾ →σ _{C₂-O₃₂} [*]	0.11		0.24
O ₃₂ -H ₃₃ ⋯C ₃₈ -O ₄₀	σ _{C₃₈-O₄₀} ⁽¹⁾ →σ _{O₃₂-H₃₃} [*]	0.25		0.25
O ₃₂ -H ₃₃ ⋯C ₃₇ -C ₃₈	σ _{C₃₈-C₃₇} ⁽¹⁾ →σ _{O₃₂-H₃₃} [*]	0.19		0.28
C ₂ -O ₃₂ ⋯H ₄₈ -O ₄₇	σ _{O₄₇-H₄₈} ⁽¹⁾ →σ _{C₂-O₃₂} [*]	0.2		0.27

Table 7. Stabilization energies $E^{(2)}$, total stabilization energies $\sum E^{(2)}$ and CT of Epicat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ and Cat $O_{32}-H_{33}\cdots Ala O_{sp^3}$ complexes.

Epicat O ₃₂ -H ₃₃ ⋯Ala O _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₃₂ -H ₃₃ ⋯O ₄₇	LP _{O47} ⁽¹⁾ →σ _{O32-H33} [*]	9	10.71	12.69
	LP _{O47} ⁽²⁾ →σ _{O32-H33} [*]	1.2		2.17
C ₃₇ -H ₄₄ ⋯O ₃₂	LP _{O32} ⁽²⁾ →σ _{C37-H44} [*]	0.22		0.44
C ₃₆ -H ₄₃ ⋯C ₂ -C ₁	π _{C1-C2} ⁽²⁾ →σ _{C36-H43} [*]	0.07		0.15
C ₁ -H ₇ ⋯O ₄₇	LP _{O47} ⁽¹⁾ →σ _{C1-H7} [*]	0.13		0.19
	LP _{O47} ⁽²⁾ →σ _{C1-H7} [*]	0.09		0.18
Cat O ₃₂ -H ₃₃ ⋯Ala O _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₃₂ -H ₃₃ ⋯O ₄₇	LP _{O47} ⁽¹⁾ →σ _{O32-H33} [*]	7.64	10.93	10.79
	LP _{O47} ⁽²⁾ →σ _{O32-H33} [*]	2.88		5.36
C ₁ -H ₇ ⋯O ₄₇	LP _{O47} ⁽¹⁾ →σ _{C1-H7} [*]	0.13		0.19

Cat O ₃₂ -H ₃₃ ...Ala O _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
C ₃₆ -H ₄₃ ...C ₂ -C ₁	$\pi_{C_1-C_2}^{(2)} \rightarrow \sigma_{C_{36}-H_{43}}^*$	0.17		0.40
O ₃₂ -H ₃₃ ...H ₄₈ -O ₄₇	$\sigma_{O_{47}-H_{48}}^{(1)} \rightarrow \sigma_{O_{32}-H_{38}}^*$	0.11		0.13

In the case of Epicat O₃₄-H₃₅...Ala O_{sp}² complex (Σ E²=23.84 kcal/mol) (Table 9), appear the interactions O₃₄-H₃₅...O₄₀, O₄₇-H₄₈...C₅-C₆ as well as other interactions with low stabilization energies such as O₄₇-H₄₈...H₈-C₅, O₃₄-H₃₅...O₄₀-C₃₈. O₃₄-H₃₅...O₄₀ is characterized by the electronic transitions LP₄₀⁽¹⁾→σ_{O₃₄-H₃₅}^{*} (E⁽²⁾= 7.73 kcal/mol, CT= 10.27 me) et LP₄₀⁽²⁾→σ_{O₃₄-H₃₅}^{*} (E⁽²⁾= 7.74 kcal/mol, CT= 16.06 me). It contributes 62.10% to the total stabilization energy of the complex. Similarly, O₄₇-H₄₈...C₅-C₆ interaction is governed by two electronic

transitions that are σ_{C₅-C₆}⁽¹⁾→σ_{O₄₄-H₄₅}^{*} (E⁽²⁾= 0.19 kcal/mol, CT= 0.26 me) et π_{C₅-C₆}⁽²⁾→σ_{O₄₄-H₄₅}^{*} (E⁽²⁾= 6.13 kcal/mol, CT= 14.55 me). In Cat O₃₄-H₃₅...Ala O_{sp}² complex, the total stabilization energy is 30.14 kcal/mol. The HB O₃₄-H₃₅...O₄₀ (E⁽²⁾= 14.06 kcal/mol, CT= 26.28 me) et O₄₇-H₄₈...O₃₄ (E⁽²⁾= 14.18 kcal/mol, CT= 26.41 me) together account for 98.7% of the Cat total stabilization energy O₃₄-H₃₅...Ala O_{sp}² complex and are its main interactions.

Table 8. Stabilization energies E⁽²⁾, total stabilization energies Σ E⁽²⁾ and CT of Epicat O₃₄-H₃₅...Ala N_{sp}³ and Cat O₃₄-H₃₅...Ala N_{sp}³ complexes.

Epicat O ₃₄ -H ₃₅ ...Ala N _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₃₄ -H ₃₅ ...N ₃₉	LP _{N₃₉} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	25.64		50.73
C ₃₆ -H ₄₂ ...O ₃₄	LP _{O₃₄} ⁽¹⁾ →σ _{C₃₆-H₄₂} [*]	0.22		0.34
C ₅ -H ₈ ...N ₃₉	LP _{N₃₉} ⁽¹⁾ →σ _{C₅-H₈} [*]	0.06		0.10
O ₃₄ -H ₃₅ ...H ₄₄ -C ₃₇	σ _{C₃₇-H₄₄} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.1	27.31	0.16
O ₃₄ -H ₃₅ ...N ₃₉ -C ₃₇	σ _{C₃₇-N₃₉} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.26		0.36
O ₃₄ -H ₃₅ ...H ₄₅ -N ₃₉	σ _{N₃₉-H₄₅} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.5		0.72
O ₃₄ -H ₃₅ ...H ₄₆ -N ₃₉	σ _{N₃₉-H₄₆} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.53		0.79

Cat O ₃₄ -H ₃₅ ...Ala N _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₃₄ -H ₃₅ ...N ₃₉	LP _{N₃₉} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	27.14		53.92
C ₃₆ -H ₄₃ ...O ₃₄	LP _{O₃₄} ⁽¹⁾ →σ _{C₃₆-H₄₃} [*]	0.27		0.60
C ₅ -H ₈ ...N ₃₉	LP _{N₃₉} ⁽¹⁾ →σ _{C₅-H₈} [*]	0.08	28.93	0.14
O ₃₄ -H ₃₅ ...N ₃₉ -C ₃₇	σ _{C₃₇-N₃₉} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.39		0.49
O ₃₄ -H ₃₅ ...H ₄₅ -N ₃₉	σ _{N₃₉-H₄₅} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.58		0.86
O ₃₄ -H ₃₅ ...H ₄₆ -N ₃₉	σ _{N₃₉-H₄₆} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.47		0.65

Table 9. Stabilization energies E⁽²⁾, total stabilization energies Σ E⁽²⁾ and CT of Epicat O₃₄-H₃₅...Ala O_{sp}² and Cat O₃₄-H₃₅...Ala O_{sp}².

Epicat O ₃₄ -H ₃₅ ...Ala O _{sp} ²				
Contact	electronic transitions	E ⁽²⁾	Σ E ⁽²⁾	CT
O ₃₄ -H ₃₅ ...O ₄₀	LP _{O₄₀} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	7.73		10.27
	LP _{O₄₀} ⁽²⁾ →σ _{O₃₄-H₃₅} [*]	7.74		16.06
C ₅ -H ₈ ...O ₄₀	LP _{O₄₀} ⁽²⁾ →σ _{C₅-H₈} [*]	0.1		0.22
O ₄₇ -H ₄₈ ...C ₅ -C ₄	σ _{C₄-C₅} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	0.12		0.16
O ₄₇ -H ₄₈ ...C ₆ -C ₅	σ _{C₅-C₆} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	0.19		0.26
	π _{C₅-C₆} ⁽²⁾ →σ _{O₄₇-H₄₈} [*]	6.13	23.84	14.55
O ₄₇ -H ₄₈ ...H ₈ -C ₅	σ _{C₅-H₈} ⁽¹⁾ →σ _{O₄₇-H₄₈} [*]	1.2		1.99
C ₃₈ -O ₄₀ ...H ₃₅ -O ₃₄	σ _{O₃₄-H₃₅} ⁽¹⁾ →π _{C₃₈-O₄₀} ^{(2)*}	0.14		0.31
O ₃₄ -H ₃₅ ...C ₃₈ -C ₃₇	σ _{C₃₇-C₃₈} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.2		0.26
O ₃₄ -H ₃₅ ...O ₄₀ -C ₃₈	σ _{C₃₈-O₄₀} ⁽¹⁾ →σ _{O₃₄-H₃₅} [*]	0.29		0.29

Cat O ₃₄ -H ₃₅ ...Ala O _{sp} ²				
Contact	electronic transitions	E ⁽²⁾	Σ E ⁽²⁾	CT
O ₃₄ -H ₃₅ ...O ₄₀	LP _{O40} ⁽¹⁾ → σ _{O34-H35} *	4.58	30.14	6.28
	LP _{O40} ⁽²⁾ → σ _{O34-H35} *	9.48		20.00
O ₄₇ -H ₄₈ ...O ₃₄	LP _{O34} ⁽¹⁾ → σ _{O47-H48} *	6.22		9.88
	LP _{O34} ⁽²⁾ → σ _{O47-H48} *	7.96		16.53
O ₄₇ -H ₄₈ ...C ₆ -C ₅	σ _{C5-C6} ⁽¹⁾ → σ _{O47-H48} *	0.09		0.12
O ₄₇ -H ₄₈ ...H ₃₅ -O ₃₄	σ _{O34-H35} ⁽¹⁾ → σ _{O47-H48} *	1.05		1.45
C ₆ -O ₃₄ ...O ₄₀	LP _{O40} ⁽²⁾ → σ _{C6-O34} *	0.11		0.24
O ₃₄ -H ₃₅ ...C ₃₈ -C ₃₇	σ _{C37-C38} ⁽¹⁾ → σ _{O34-H35} *	0.2		0.28
O ₃₄ -H ₃₅ ...O ₄₀ -C ₃₈	σ _{C38-O40} ⁽¹⁾ → σ _{O34-H35} *	0.25		0.25
C ₆ -O ₃₄ ...H ₄₈ -O ₄₇	σ _{O47-H48} ⁽¹⁾ → σ _{C6-O34} *	0.2		0.27

The stabilization energy of the O₃₄-H₃₅...O₄₇ HB is 9.96 kcal/mol in the Epicat O₃₄-H₃₅...Ala O_{sp}³ complex (Σ E²=10.40 kcal/mol) and 9.50 kcal/mol in the Cat O₃₄-H₃₅...Ala O_{sp}² complex (Σ E²=10.03 kcal/mol) (Table 10). These values account for approximately 95% of these two complexes total stabilization energy. The O₃₄-H₃₅...O₄₇ is thus their main interaction. The others are weak. They comprise C₅-H₈...O₄₇, C₃₆-H₄₁...O₃₄, O₃₄-H₃₅...H₄₄-C₃₇...

The NBO analysis makes it possible to specify all the interactions between Epicat or Cat and Ala in all the

complexes. In the latter, in addition to the bonds X - H ... Y (X = O, N, C; Y = O, N), several other interactions appear. These include O - H ... C = O, O - H ... C = C, C - H ... C = C, O - H ... H - O, C - H ... H - C, N - H ... H - O, O - H ... H - C. X - H ... Y is characterized by CT LP_Y → σ_{X-H}* in which σ_{X-H}* acts as electron acceptor and LP_Y as an electron donor. The O - H ... O, O - H ... N and N - H ... O HB are the main ones for EpiCat... Ala and Cat... Ala complexes. For this latter, interactions with the N_{sp}³ and O_{sp}² heteroatoms are stronger than those with O_{sp}³. They have the most intense stabilization energies E⁽²⁾.

Table 10. Stabilization energies E⁽²⁾, total stabilization energies Σ E⁽²⁾ and CT of Epicat O₃₄-H₃₅...Ala O_{sp}³ and Cat O₃₄-H₃₅...Ala O_{sp}².

Epicat O ₃₄ -H ₃₅ ... Ala O _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₃₄ -H ₃₅ ...O ₄₇	LP _{O47} ⁽¹⁾ →σ _{O34-H35} [*]	7.51	10.40	10.53
	LP _{O47} ⁽²⁾ →σ _{O34-H35} [*]	2.44		4.65
C ₃₆ -H ₄₁ ...O ₃₄	LP _{O34} ⁽²⁾ →σ _{C36-H41} [*]	0.06		0.12
C ₃₆ -H ₄₁ ...C ₆ -C ₅	π _{C5-C6} ⁽²⁾ →σ _{C36-H41} [*]	0.08		0.19
C ₅ -H ₈ ...O ₄₇	LP _{O47} ⁽¹⁾ →σ _{C5-H8} [*]	0.15		0.19
	LP _{O47} ⁽²⁾ →σ _{C5-H8} [*]	0.05		0.10
O ₃₄ -H ₃₅ ...H ₄₈ -O ₄₇	σ _{O47-H48} ⁽¹⁾ →σ _{O34-H35} [*]	0.11		0.15
Cat O ₃₄ -H ₃₅ ... Ala O _{sp} ³				
Contact	electronic transitions	E ⁽²⁾ (kcal/mol)	Σ E ⁽²⁾ (kcal/mol)	CT (me)
O ₃₄ -H ₃₅ ...O ₄₇	LP _{O47} ⁽¹⁾ →σ _{O34-H35} [*]	9.14	10.03	12.97
	LP _{O47} ⁽²⁾ →σ _{O34-H35} [*]	0.36		0.71
C ₃₇ -H ₄₄ ...O ₃₄	LP _{O34} ⁽²⁾ →σ _{C37-H44} [*]	0.18		0.37
C ₅ -H ₈ ...O ₄₇	LP _{O47} ⁽¹⁾ →σ _{C5-H8} [*]	0.12		0.16
	LP _{O47} ⁽²⁾ →σ _{C5-H8} [*]	0.14		0.28
O ₃₄ -H ₃₅ ...H ₄₄ -C ₃₇	σ _{C37-H44} ⁽¹⁾ →σ _{O34-H35} [*]	0.09		0.16

3.3. Geometric and Spectroscopic Parameters

Tables 11 and 12 summarize some geometric and spectroscopic parameters of X - H ... Y in EpiCat ... Ala and Cat ... Ala complexes, respectively. They show the bond distances and angles between interacting atoms, the stretching frequencies of O-H, N-H and C-H and their variations. The results indicate that for O - H ... O, O - H ... N and N - H ... O

HB, the fluctuations in the lengths of the hydroxyl groups (Δd_{O-H}) and those of the N - H group (Δd_{N-H}) are positive (Tables 11 and 12). These O - H and N - H bond elongations result from the CT of the LP_Y the electron-rich proton acceptor Y (Y = O, N) to the antibonding orbital σ_{X-H}* of X - H (X = O, N, C). This is the LP_Y → σ_{X-H}* electronic transition. This result is consistent with the conclusions of Albugin and al. [46]. These CT are explained by the reduction of the elongation

(redshift) frequencies of O-H and N-H ($\Delta\nu_{\text{O-H}} < 0$ and $\Delta\nu_{\text{N-H}} < 0$). From this fact, all O - H \cdots O, O - H \cdots N and N - H \cdots O interactions are proper HB [47.48]. O - H \cdots O and O - H \cdots N exhibit intermolecular distance $d(\text{H}\cdots\text{O})$ and $d(\text{H}\cdots\text{N})$ between 1.79761 Å and 1.97709 Å and then linearity angles $\angle \text{XHY}$ ranging from 139.4° to 176.6°. These data are in keeping with those of moderate HB ($1.5 \text{ Å} < d(\text{H}\cdots\text{O}) < 2.2 \text{ Å}$ and $130^\circ < \angle \text{XHY} < 180^\circ$) [49.50]. In contrast, for N - H \cdots O, $d(\text{H}\cdots\text{O})$ is between 2.35220 Å and 3.36209 Å with $\angle \text{XHY}$ between 147.3° and 154.9°. These values suggest that the N -

H \cdots O is weak in these complexes. The last column of Table 11 or 12 shows the strengths of HB in these due to both the contraction of the C - H bonds and the increase in its elongation frequency (blue shift) ($\Delta d_{\text{C-H}} < 0$ and $\Delta\nu_{\text{C-H}} > 0$). C - H \cdots Y interactions are improper or unconventional HB [47.48]. For these, the intermolecular distance $d(\text{H}\cdots\text{O})$ and $d(\text{H}\cdots\text{N})$ and the linearity angles $\angle \text{XHY}$ are between 2.64825 and 3.58144 Å and then between 119.2° and 158.8° respectively. These values indicate that these improper HB are weak.

Table 11. Geometric parameters of the X - H \cdots Y interactions (distance in Å, angle in degree) and the elongation frequencies ν (cm^{-1}) of the X - H bonds in Cat \cdots Ala complexes and their variation $\Delta\nu$ (cm^{-1}).

Complexes	Contact	$d(\text{H}\cdots\text{Y})$ (Å)	$\angle \text{XHY}^\circ$	$\Delta d(\text{X-H})$ (Å)	$\Delta\nu_{\text{X-H}}$ (cm^{-1})
Cat O ₂₈ -H ₂₉ \cdots Ala N _{sp} ³	O ₂₈ -H ₂₉ \cdots N ₃₉	1.81368	169.5	0.02911	-598.3
	C ₂₂ -H ₂₅ \cdots O ₄₀	2.64825	158.8	-0.00183	24.1
	C ₃₆ -H ₄₂ \cdots O ₂₈	2.98252	119.2	-0.00171	19.9
Cat O ₂₈ -H ₂₉ \cdots Ala O _{sp} ²	O ₂₈ -H ₂₉ \cdots O ₄₀	1.80941	167.9	0.01485	-291.3
	N ₃₉ -H ₄₅ \cdots O ₂₈	2.35220	147.7	0.00107	-9.1
	O ₂₂ -H ₂₅ \cdots O ₄₀	2.87216	123.1	-0.00158	20.6
Cat O ₂₈ -H ₂₉ \cdots Ala O _{sp} ³	O ₂₈ -H ₂₉ \cdots O ₄₇	1.91027	162.8	0.00827	-160.2
	N ₃₉ -H ₄₆ \cdots O ₂₈	2.36209	154.8	0.00174	0.3
	C ₂₂ -H ₂₅ \cdots O ₄₇	3.58144	130.0	-0.00101	12.5
Cat O ₃₂ -H ₃₃ \cdots Ala N _{sp} ³	O ₃₂ -H ₃₃ \cdots N ₃₉	1.81460	166.0	0.02773	-570.3
	C ₁ -H ₇ \cdots N ₃₉	3.05293	120.9	-0.00168	19.1
	O ₃₂ -H ₃₃ \cdots O ₄₀	1.89987	139.5	0.01517	-268.7
Cat O ₃₂ -H ₃₃ \cdots Ala O _{sp} ²	O ₄₇ -H ₄₈ \cdots O ₃₂	1.85602	153.2	0.01305	-276.1
	O ₃₂ -H ₃₃ \cdots O ₄₇	1.96426	169.3	0.00591	-118.9
	C ₁ -H ₇ \cdots O ₄₇	3.01039	124.9	-0.00091	12.6
Cat O ₃₄ -H ₃₅ \cdots Ala N _{sp} ³	O ₃₄ -H ₃₅ \cdots N ₃₉	1.82916	170.9	0.02664	-547.2
	C ₃₆ -H ₄₃ \cdots O ₃₄	2.92218	126.6	-0.00237	12.0
	C ₅ -H ₈ \cdots N ₃₉	2.97692	121.9	-0.00189	20.6
Cat O ₃₄ -H ₃₅ \cdots Ala O _{sp} ²	O ₃₄ -H ₃₅ \cdots O ₄₀	1.90542	139.8	0.01469	-259.6
	O ₄₇ -H ₄₈ \cdots O ₃₄	1.86270	153.3	0.01354	-280.4
	O ₃₄ -H ₃₅ \cdots O ₄₇	1.97331	171.9	0.00552	-106.2
Cat O ₃₄ -H ₃₅ \cdots Ala O _{sp} ³	C ₅ -H ₈ \cdots O ₄₇	2.95605	125.4	-0.00105	13.3

Table 12. Geometric parameters of the X - H \cdots Y interactions (distance in Å, angle in degree) and the elongation frequencies ν (cm^{-1}) of the X - H bonds in Epicat \cdots Ala complexes and their variation $\Delta\nu$ (cm^{-1}).

Complexes	Contact	$d(\text{H}\cdots\text{Y})$ (Å)	$\angle \text{XHY}^\circ$	$\Delta d(\text{X-H})$ (Å)	$\Delta\nu_{\text{X-H}}$ (cm^{-1})
Epicat O ₂₈ -H ₂₉ \cdots Ala N _{sp} ³	O ₂₈ -H ₂₉ \cdots N ₃₉	1.79761	171.1	0.03092	-634.8
	O ₃₆ -H ₄₁ \cdots O ₂₈	2.96005	123.7	-0.002	92.8
	O ₂₈ -H ₂₉ \cdots O ₄₀	1.80454	168.2	0.01523	-299.1
Epicat O ₂₈ -H ₂₉ \cdots Ala O _{sp} ²	N ₃₉ -H ₄₅ \cdots O ₂₈	2.36187	147.3	0.00096	-8.8
	C ₂₂ -H ₂₅ \cdots O ₄₃	2.85709	123.2	-0.00157	20.5
	O ₂₈ -H ₂₉ \cdots O ₄₇	1.93377	172.6	0.00702	-138.7
Epicat O ₂₈ -H ₂₉ \cdots Ala O _{sp} ³	O ₃₆ -H ₄₁ \cdots O ₂₈	2.85381	150.7	-0.00274	19.6
	C ₂₂ -H ₂₅ \cdots O ₄₇	3.04357	124.2	-0.00103	12.6
	O ₃₂ -H ₃₃ \cdots N ₃₉	1.81873	167.0	0.02739	-562.9
Epicat O ₃₂ -H ₃₃ \cdots Ala N _{sp} ³	C ₁ -H ₇ \cdots N ₃₉	2.93336	128.2	-0.00167	18.4
	O ₄₇ -H ₄₈ \cdots O ₃₂	1.86110	153.5	0.01348	-278.9
	O ₃₂ -H ₃₃ \cdots O ₄₀	1.91121	139.4	0.01459	-258.5
Epicat O ₃₂ -H ₃₃ \cdots Ala O _{sp} ²	O ₃₂ -H ₃₃ \cdots O ₄₇	1.96248	168.9	0.0058	-114.5
	C ₁ -H ₇ \cdots O ₄₇	2.98091	125.1	-0.00098	11.2
	O ₃₄ -H ₃₅ \cdots N ₃₉	1.84464	171.1	0.02576	-527.6
Epicat O ₃₄ -H ₃₅ \cdots Ala N _{sp} ³	C ₃₆ -H ₄₂ \cdots O ₃₄	2.80094	122.9	-0.00203	90.9
	C ₅ -H ₈ \cdots N ₃₉	2.98129	122.3	-0.00185	20.7
	O ₃₄ -H ₃₅ \cdots O ₄₀	1.90422	176.6	0.009	-164.8
Epicat O ₃₄ -H ₃₅ \cdots Ala O _{sp} ²	C ₅ -H ₈ \cdots O ₄₀	2.72268	126.6	-0.00162	20.7
	O ₃₄ -H ₃₅ \cdots O ₄₇	1.97709	171.3	0.00541	-107.3
	C ₃₆ -H ₄₁ \cdots O ₃₄	3.26338	133.9	-0.00274	110.5
Epicat O ₃₄ -H ₃₅ \cdots Ala O _{sp} ³	O ₅ -H ₈ \cdots O ₄₇	2.97008	125.0	-0.0009	12.4

Table 13. Hybridization of the *X* atom of the *X*-H bond in monomers and in the *X*-H...*Y* contact and the variations of the *S* character of the *X* atom ($\Delta S\%$) with complexation for the Cat... Ala complex.

Complexes	Hybridization sp^n (X-H)		$\Delta S\%$	$\Delta\sigma^*$ (10^{-3} e)	Complexes	Hybridization sp^n (X-H)		$\Delta S\%$	$\Delta\sigma^*$ (10^{-3} e)
	Monomers	Complexes				Monomers	Complexes		
Cat O ₂₈ – H ₂₉ ... Ala N _{sp} ³					Cat O ₂₈ – H ₂₉ ... Ala O _{sp} ²				
LP _{N39} → $\sigma_{O28-H29}^*$	sp ^{3.40}	sp ^{2.48}	6	63.39	LP _{O40} → $\sigma_{O28-H29}^*$	sp ^{3.40}	sp ^{2.67}	4.5	33.90
LP _{O40} → $\sigma_{C22-H25}^*$	sp ^{2.37}	sp ^{2.32}	0.49	1.98	LP _{O28} → $\sigma_{N39-H45}^*$	sp ^{2.93}	sp ^{2.78}	1.04	8.63
LP _{O28} → $\sigma_{C36-H42}^*$	sp ^{3.04}	sp ^{3.03}	0.05	-0.08	LP _{O40} → $\sigma_{C22-H25}^*$	sp ^{2.37}	sp ^{2.34}	0.27	0.27
Cat O ₂₈ – H ₂₉ ... Ala O _{sp} ³									
LP _{O47} → $\sigma_{O28-H29}^*$	sp ^{3.40}	sp ^{2.81}	3.48	22.69					
LP _{O28} → $\sigma_{N39-H46}^*$	sp ^{2.93}	sp ^{2.78}	1.04	5.46					
LP _{O47} → $\sigma_{C22-H25}^*$	sp ^{2.37}	sp ^{2.36}	0.14	0.45					
Cat O ₃₂ – H ₃₃ ... Ala N _{sp} ³					Cat O ₃₂ – H ₃₃ ... Ala O _{sp} ²				
LP _{N39} → $\sigma_{O32-H33}^*$	sp ^{3.53}	sp ^{3.58}	5.88	60.29	LP _{O40} → $\sigma_{O32-H33}^*$	sp ^{3.53}	sp ^{2.84}	3.97	32.53
LP _{N39} → σ_{C1-H7}^*	sp ^{2.24}	sp ^{2.24}	0.03	0.24	LP _{O32} → $\sigma_{O47-H48}^*$	sp ^{3.45}	sp ^{2.79}	3.9	28.10
Cat O ₃₂ – H ₃₃ ... Ala O _{sp} ³									
LP _{O47} → $\sigma_{O32-H33}^*$	sp ^{3.53}	sp ^{2.98}	3.09	19.67					
LP _{O47} → σ_{C1-H7}^*	sp ^{2.24}	sp ^{2.23}	0.14	0.42					
Cat O ₃₄ – H ₃₅ ... Ala N _{sp} ³					Cat O ₃₄ – H ₃₅ ... Ala O _{sp} ²				
LP _{N39} → $\sigma_{O34-H35}^*$	sp ^{3.56}	sp ^{2.57}	6.03	60.54	LP _{O40} → $\sigma_{O34-H35}^*$	sp ^{3.56}	sp ^{2.87}	3.86	31.49
LP _{O34} → $\sigma_{C36-H43}^*$	sp ^{3.14}	sp ^{3.01}	0.78	1.16	LP _{O34} → $\sigma_{O47-H48}^*$	sp ^{3.45}	sp ^{2.84}	3.57	28.27
LP _{N39} → σ_{C5-H8}^*	sp ^{2.28}	sp ^{2.28}	-	0.29					
Cat O ₃₄ – H ₃₅ ... Ala O _{sp} ³									
LP _{O47} → $\sigma_{O34-H35}^*$	sp ^{3.56}	sp ³	3.02	18.23					
LP _{O34} → $\sigma_{C37-H44}^*$	sp ^{3.13}	sp ^{3.11}	0.09	-0.29					
LP _{O47} ⁽¹⁾ → σ_{C5-H8}^*	sp ^{2.28}	sp ^{2.27}	0.11	0.6					

3.4. Effect of Hybridization and “Hyperconjugative” Interactions

Tables 13 and 14 show the hybridization of the O_i atom of the O_i-H_j hydroxyl groups of EpiCat and Cat. It also illustrates the variation of the *s* character of O_i atoms. In the complexes EpiCat ... Ala and Cat... Ala, there is the hybridization of the O_i in the O_i-H_j... O and O_i-H_j... N. In monomers (EpiCat and Cat), O_i atoms are hybridized sp^{*x*} (*x* ∈ 3.39 – 3.67) while in complexes, other oxygen are hybridized sp^{*m*} (*m* ∈ 2.48 – 3.58). There is a decrease in character *p* and an increase in *s* one ($\Delta S\% > 0$). The reduction in O_i-H_j bond length is a consequence of this second hybridization [46].

However, this work notes the presence of the “hyperconjugative” interactions LP_O → $\sigma_{O_i-H_j}^*$ and LP_N → $\sigma_{O_i-H_j}^*$ with high stabilization energies *E*⁽²⁾; these vary from 7.45 kcal/mol to 31.54 kcal/mol. This interaction

increases the electron density of the antibonding orbital $\sigma_{O_i-H_j}^*$ polarized along the HB axis. Thereby, it weakens the O_i – H_j bonds; it increases their length [46]. Its effects outweigh those of hybridization [39].

For HB C_i-H_j...O or C_i-H_j...N interactions, the carbon C is hybridized sp^{*x*} (*x* ∈ 2.23 – 3.15) whereas in Ala, EpiCat and Cat its homologs are hybridized sp^{*m*} (*m* ∈ 2.24 – 2.93), sp^{*n*} (*n* ∈ 3.04 – 3.14), sp^{*t*} (*t* ∈ 2.24 – 2.37) respectively. These values indicate that its hybridization increases its character *s* and lowers that of *p*. This hybridization contributes to contracting the C_i-H_j bond. In these complexes, “hyperconjugative” interactions result in LP_O → $\sigma_{C_i-H_j}^*$ and LP_N → $\sigma_{C_i-H_j}^*$; they correspond to low stabilization energies *E*⁽²⁾; these go from 0.1 kcal/mol to 0.83 kcal/mol. This C_i-H_j bond shortening is illustrated in the data in tables 11 and 12. Its contraction ($\Delta d(C_i-H_j) < 0$) means that the effects of its hybridization outweigh those of “hyperconjugative” interactions.

Table 14. *X* Hybridization in the *X*-H bond in monomers and in the *X*-H...*Y* contact and the variations of the *s* character of the *X* atom ($\Delta S\%$) with complexation for the Cat... Ala.

Complexes	Hybridization sp^n (X-H)		$\Delta S\%$	$\Delta\sigma^*$ (10^{-3} e)	Complexes	Hybridization sp^n (X-H)		$\Delta S\%$	$\Delta\sigma^*$ (10^{-3} e)
	Monomers	Complexes				Monomers	Complexes		
Epicat O ₂₈ – H ₂₉ ... Ala N _{sp} ³					Epicat O ₂₈ – H ₂₉ ... Ala O _{sp} ²				
LP _{N39} → $\sigma_{O28-H29}^*$	sp ^{3.39}	sp ^{2.48}	6	66.62	LP _{O40} → $\sigma_{O28-H29}^*$	sp ^{3.39}	sp ^{2.66}	4.54	34.56
LP _{O28} → $\sigma_{C36-H42}^*$	sp ^{3.04}	sp ^{3.05}	-	-0.22	LP _{O28} → $\sigma_{N46-H48}^*$	sp ^{2.93}	sp ^{2.78}	1.01	8.40
			0.06		LP _{O40} → $\sigma_{C22-H25}^*$	sp ^{2.32}	sp ^{2.35}	-	0.97
								0.21	
Epicat O ₂₈ – H ₂₉ ... Ala O _{sp} ³									
LP _{O47} → $\sigma_{O28-H29}^*$	sp ^{3.39}	sp ^{2.84}	3.26	20.76					
LP _{O28} → $\sigma_{C36-H41}^*$	sp ^{3.04}	sp ^{3.01}	-	-0.52					

	Hybridization sp ⁿ (X-H)		Δ%S	Δσ* (10 ⁻³ e)	Complexes	Hybridization sp ⁿ (X-H)		Δ%S	Δσ* (10 ⁻³ e)
	Monomers	Complexes				Monomers	Complexes		
LP _{O47} → σ _{C22-H25} [*]	sp ^{2,32}	sp ^{2,37}	0.12	1.35					
Epicat O ₃₂ - H ₃₃ ... Ala N _{sp³}			-		Epicat O ₃₂ - H ₃₃ ... Ala O _{sp²}				
LP _{N39} → σ _{O32-H33} [*]	sp ^{3,56}	sp ^{2,58}	5.95	59.72	LP _{O32} → σ _{O47-H48} [*]	sp ^{3,45}	sp ^{2,84}	3.55	28.32
LP _{N39} → σ _{C1-H7} [*]	sp ^{2,28}	sp ^{2,24}	0.38	-0.63	LP _{O40} → σ _{O32-H33} [*]	sp ^{3,56}	sp ^{2,86}	3.98	31.15
Epicat O ₃₂ - H ₃₃ ... Ala O _{sp³}									
LP _{O47} → σ _{O32-H33} [*]	sp ^{3,56}	sp ^{2,97}	3.21	19.20					
LP _{O32} → σ _{C37-H44} [*]	sp ^{3,13}	sp ^{3,15}	-	-2.45					
Epicat O ₃₄ - H ₃₅ ... Ala N _{sp³}					Epicat O ₃₄ - H ₃₅ ... Ala O _{sp²}				
LP _{N43} → σ _{O34-H35} [*]	sp ^{3,67}	sp ^{2,61}	6.24	59.60	LP _{O40} → σ _{O34-H35} [*]	sp ^{3,67}	sp ^{2,93}	3.99	32.79
LP _{O34} → σ _{C36-H41} [*]	sp ^{3,04}	sp ^{3,02}	0.12	-0.13	LP _{O40} → σ _{C5-H8} [*]	sp ^{2,24}	sp ^{2,23}	0.07	1.84
LP _{N43} → σ _{C5-H8} [*]	sp ^{2,24}	sp ^{2,28}	-	1.02					
Epicat O ₃₄ - H ₃₅ ... Ala O _{sp³}									
LP _{O47} → σ _{O34-H35} [*]	sp ^{3,67}	sp ^{3,02}	3.45	19.28					
LP _{O34} → σ _{C36-H41} [*]	sp ^{3,04}	sp ^{3,02}	0.09	-1.19					
LP _{O47} → σ _{C5-H8} [*]	sp ^{2,24}	sp ^{2,26}	-	1.29					

3.5. AIM Analysis

Figures 6 and 7 illustrate, as an example, the AIM-molecular graphs of Epicat O₂₈ - H₂₉ ... Ala N_{sp³}, Epicat O₂₈ - H₂₉ ... Ala O_{sp²} and Epicat O₂₈ - H₂₉ ... Ala O_{sp³} and then Cat O₂₈ - H₂₉ ... Ala N_{sp³}, Cat O₂₈ - H₂₉ ... Ala O_{sp²} and Cat O₂₈ - H₂₉ ... Ala O_{sp³} complexes, respectively; the small green spheres are BCP.

Tables 15 and 16 present results of the X-H...Y BCP (X=O, N, C) and (Y=O, N) interactions topological analyzes for EpiCat ... Ala and Cat ... Ala complexes. In these, the electronic density ρ(r) between the hydroxyl groups O_i - H_j with the Nsp³, Osp² and Osp³ heteroatoms of Ala, vary between 0.004408 ea₀⁻³ and 0.043552 ea₀⁻³. Those of ∇² ρ(r) range from 0.017568 ea₀⁻⁵ to 0.101413 ea₀⁻⁵. Values of ∇² ρ(r) are positive. They indicate that these interactions are HB. Their total negative electron energy density H(r) O_i-H_j...N_{sp³}, O_i-H_j...O_{sp²} and O_i-H_j...O_{sp³} reflects moderate HB [40]. Moreover, this work adopts ratios - $\frac{G(r)}{V(r)}$ between 0.5 and 1. These moderate HB are partially covalent.

The total electron energy density H(r) is slightly positive for the interactions O₃₄-H₃₅...O₄₀, O₂₈-H₂₉...O₄₀ of EpiCat ... Ala complexes and O₂₈-H₂₉...O₄₀, O₂₈-H₂₉...O₄₇ for those formed from Cat ...Ala Their - $\frac{G(r)}{V(r)}$ ratios are greater than 1. These values

correspond to interactions which are within the moderate HB limit with a partially covalent character. Those of the HB E_{HB} energies calculated using Espinoza's method are reported in the last columns of tables 15 and 16. For the interactions linked to hydroxyl groups O_i - H_j, they vary from 5.0929938 kcal/mol to 10.0687117 kcal/mol. These interactions are moderate HB; they meet Espinoza's criteria (4 kcal/mole < E_{HB} < 15 kcal/mole) [51]. They include the interactions O₃₂-H₃₃...N₃₉ (E_{HB}= 8.80 kcal/mol), O₃₂-H₃₃...O₄₀ (E_{HB}= 7.22 kcal/mol), O₃₂-H₃₃...O₄₇ (E_{HB}= 4.60 kcal/mol), O₃₄-H₃₅...N₃₉ (E_{HB}= 8.35 kcal/mol), O₃₄-H₃₅...O₄₀ (E_{HB}= 7.14 kcal/mol), O₃₄-H₃₅...O₄₇ (E_{HB}= 5.27 kcal/mol), O₂₈-H₂₉...N₃₉ (E_{HB}= 8.71 kcal/mol), O₂₈-H₂₉...O₄₀ (E_{HB}= 7.47 kcal/mol) et O₂₈-H₂₉...O₄₇ (E_{HB}= 6.08 kcal/mol) of Cat ... Ala complexes (Table 15). For the EpiCat... Ala ones (Table 16), they are the O₃₂-H₃₃...N₃₉ (E_{HB} = 8.69 kcal/mol), O₃₂-H₃₃...O₄₀ (E_{HB} = 7.05 kcal/mol), O₃₂-H₃₃...O₄₇ (E_{HB}= 5.44 kcal/mol), O₃₄-H₃₅...N₃₉ (E_{HB}= 8.05 kcal/mol), O₃₄-H₃₅...O₄₀ (E_{HB}= 5.71 kcal/mol), O₃₄-H₃₅...O₄₇ (E_{HB}= 5.29 kcal/mol), O₂₈-H₂₉...N₃₉ (E_{HB}= 9.16 kcal/mol), O₂₈-H₂₉...O₄₀ (E_{HB}= 7.55 kcal/mol), O₂₈-H₂₉...O₄₇ (E_{HB}= 5.62 kcal/mol). However, for C-H...O and N-H...O interactions, the values of ∇² ρ(r) and H(r) are positive. Those of C-H...O and N-H...O are weak. Moreover, the - $\frac{G(r)}{V(r)}$ ratios are greater than 1. They suggest that their characters be non-covalent or electrostatic. E_{HB} values are less than 4 kcal/mol.

Table 15. AIM parameters of BCP of Cat ...Ala interactions.

Contact	ρ(r) _{X-H...Y} (ea ₀ ⁻³)	∇ ² ρ(r) _{X-H...Y} (ea ₀ ⁻⁵)	V(r) (ua)	G(r) (ua)	H(r) (ua)	- $\frac{G(r)}{V(r)}$	E _{HB} (kcal/ mole)
Cat O ₂₈ -H ₂₉ ... Ala N _{sp³}							
O ₂₈ -H ₂₉ ...N ₃₉	0.04182	0.090873	-0.027754	0.025236	-0.002518	0.91	8.71
C ₂₂ -H ₂₅ ...O ₄₀	0.006168	0.021917	-0.003838	0.004658	0.00082	1.21	1.20
Cat O ₂₈ -H ₂₉ ... Ala O _{sp²}							
O ₂₈ -H ₂₉ ...O ₄₀	0.032491	0.100229	-0.023805	0.024431	0.000626	1.03	7.47

Contact	$\rho(r)_{X-H...Y} (ea_0^{-3})$	$\nabla^2\rho(r)_{X-H...Y} (ea_0^{-5})$	$V(r) (ua)$	$G(r) (ua)$	$H(r) (ua)$	$-\frac{G(r)}{V(r)}$	$E_{HB} (kcal/mole)$
N ₃₉ -H ₄₅ ...O ₂₈	0.011022	0.036611	-0.008111	0.008632	0.000521	1.06	2.54
Cat O ₂₈ -H ₂₉ ... Ala O _{sp} ³							
O ₂₈ -H ₂₉ ...O ₄₇	0.025778	0.078407	-0.019391	0.019496	0.000105	1.01	6.08
N ₃₉ -H ₄₆ ...O ₂₈	0.011117	0.035634	-0.008101	0.008505	0.000404	1.05	2.54
Cat O ₃₂ -H ₃₃ ... Ala N _{sp} ³							
O ₃₂ -H ₃₃ ...N ₃₉	0.041844	0.092378	-0.028034	0.025564	-0.00247	0.91	8.80
Cat O ₃₂ -H ₃₃ ... Ala O _{sp} ²							
O ₃₂ -H ₃₃ ...O ₄₀	0.02975	0.085458	-0.023009	0.022187	-0.000822	0.96	7.22
O ₄₇ -H ₄₈ ...O ₃₂	0.030158	0.088992	-0.022603	0.022425	-0.000178	0.99	7.09
Cat O ₃₂ -H ₃₃ ... Ala O _{sp} ³							
O ₃₂ -H ₃₃ ...O ₄₇	0.018988	0.057703	-0.014663	0.014544	-0.000119	0.99	4.60
C ₁ -H ₇ ...O ₄₇	0.006556	0.022644	-0.004028	0.004844	0.000816	1.20	1.26
Cat O ₃₄ -H ₃₅ ... Ala N _{sp} ³							
O ₃₄ -H ₃₅ ...N ₃₉	0.040465	0.089222	-0.026609	0.024457	-0.002152	0.92	8.35
Cat O ₃₄ -H ₃₅ ... Ala O _{sp} ²							
O ₃₄ -H ₃₅ ...O ₄₀	0.029382	0.084394	-0.02275	0.021924	-0.000826	0.96	7.14
O ₄₇ -H ₄₈ ...O ₃₄	0.029843	0.087166	-0.022276	0.022034	-0.000242	0.99	6.99
Cat O ₃₄ -H ₃₅ ... Ala O _{sp} ³							
O ₃₄ -H ₃₅ ...O ₄₇	0.022293	0.066072	-0.016785	0.016651	-0.000134	0.99	5.27
C ₃₆ -H ₄₁ ...H ₈ -C ₅	0.001478	0.004523	-0.000526	0.000829	0.000303	1.58	0.17

Table 16. AIM parameters of the bond critical point (BCP) of EpiCat ... Ala interactions.

Contact	$\rho(r)_{X-H...Y} (ea_0^{-3})$	$\nabla^2\rho(r)_{X-H...Y} (ea_0^{-5})$	$V(r) (ua)$	$G(r) (ua)$	$H(r) (ua)$	$-\frac{G(r)}{V(r)}$	$E_{HB} (kcal/mole)$
Epicate O ₂₈ -H ₂₉ ... Ala N _{sp} ³							
O ₂₈ -H ₂₉ ...N ₃₉	0.043552	0.093384	-0.029193	0.02627	-0.002923	0.900	9.16
Epicate O ₂₈ -H ₂₉ ... Ala O _{sp} ²							
O ₂₈ -H ₂₉ ...O ₄₀	0.032866	0.101413	-0.024061	0.024707	0.000646	1.027	7.55
N ₃₉ -H ₄₆ ...O ₂₈	0.010796	0.036099	-0.007924	0.008474	0.000550	1.069	2.49
Epicate O ₂₈ -H ₂₉ ... Ala O _{sp} ³							
O ₂₈ -H ₂₉ ...O ₄₇	0.024023	0.072554	-0.017924	0.018031	0.000107	1.006	5.62
C ₃₆ -H ₄₁ ...O ₂₈	0.004408	0.017568	-0.002366	0.003379	0.001013	1.428	0.74
Epicate O ₃₂ -H ₃₃ ... Ala N _{sp} ³							
O ₃₂ -H ₃₃ ...N ₃₉	0.041471	0.091515	-0.027694	0.025286	-0.002408	0.913	8.69
Epicate O ₃₂ -H ₃₃ ... Ala O _{sp} ²							
O ₄₇ -H ₄₈ ...O ₃₂	0.030009	0.087435	-0.022426	0.022142	-0.000284	0.987	7.04
O ₃₂ -H ₃₃ ...O ₄₀	0.029005	0.083394	-0.02248	0.021664	-0.000816	0.964	7.05
Epicate O ₃₂ -H ₃₃ ... Ala O _{sp} ³							
O ₃₂ -H ₃₃ ...O ₄₇	0.022996	0.068124	-0.017323	0.017177	-0.000146	0.992	5.44
C ₃₆ -H ₄₁ ...H ₇ -C ₁	0.001642	0.005001	-0.000593	0.000922	0.000329	1.555	0.19
Epicate O ₃₄ -H ₃₅ ... Ala N _{sp} ³							
O ₃₄ -H ₃₅ ...N ₃₉	0.039207	0.086047	-0.025642	0.023577	-0.002065	0.919	8.05
Epicate O ₃₄ -H ₃₅ ... Ala O _{sp} ²							
O ₃₄ -H ₃₅ ...O ₄₀	0.024635	0.078477	-0.018194	0.018906	0.000712	1.039	5.71
C ₅ -H ₈ ...O ₄₀	0.006104	0.022354	-0.003629	0.004609	0.000980	1.270	1.14
N ₃₉ -H ₄₅ ...C ₅ = C ₆	0.005379	0.017954	-0.002411	0.00345	0.001039	1.431	0.76
Epicate O ₃₄ -H ₃₅ ... Ala O _{sp} ³							
O ₃₄ -H ₃₅ ...O ₄₇	0.02241	0.065291	-0.016853	0.016588	-0.000265	0.984	5.29
C ₃₆ -H ₄₁ ...H ₈ -C ₅	0.001906	0.006106	-0.000723	0.001125	0.000402	1.556	0.23

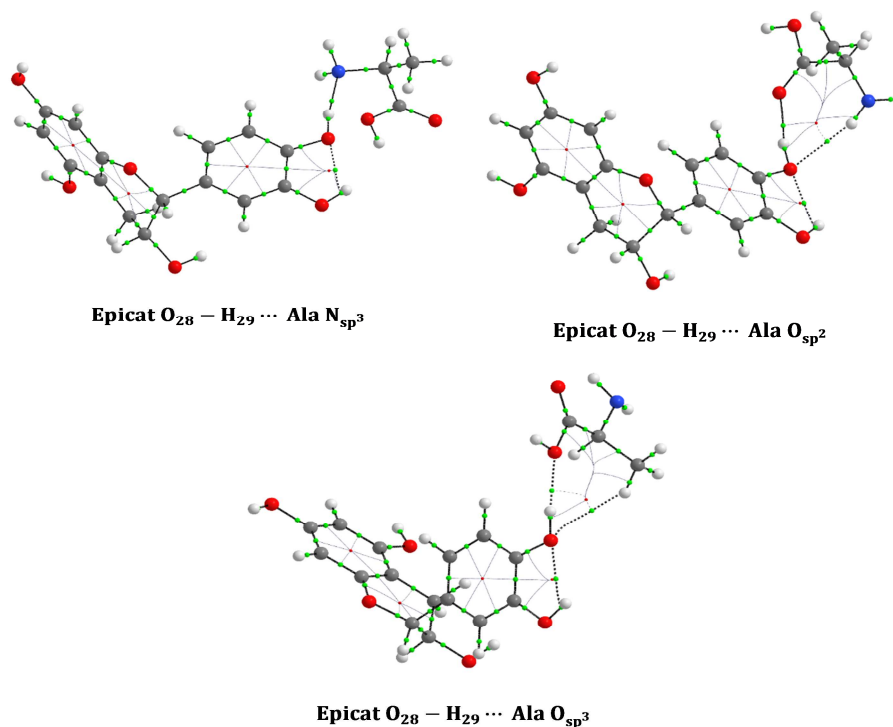


Figure 6. AIM molecular plots of the Cat O₂₈-H₂₉...Ala N_{sp³}, Cat O₂₈-H₂₉...Ala O_{sp²} and Cat O₂₈-H₂₉...Ala O_{sp³}.

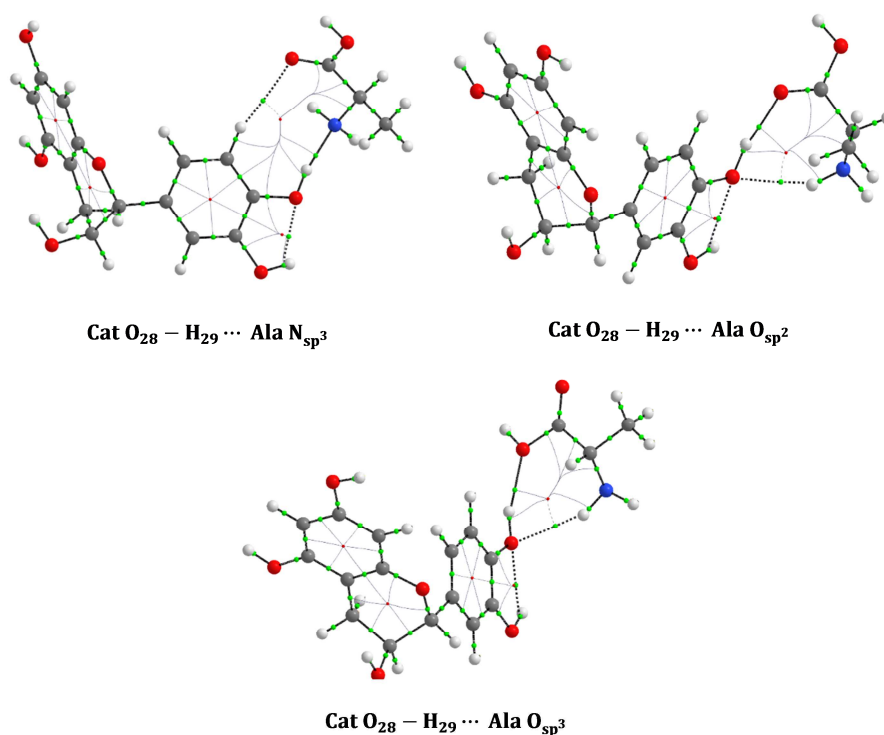


Figure 7. AIM molecular plots of the Epicat O₂₈-H₂₉...Ala N_{sp³}, Epicat O₂₈-H₂₉...Ala O_{sp²} and Epicat O₂₈-H₂₉...Ala O_{sp³} complexes.

3.6. NCI Analysis

The NCI analysis is restricted to the interactions of the O₂₈ – H₂₉ hydroxyl group of EpiCat and Cat with the N_{sp³}, O_{sp²} and O_{sp³} heteroatoms of Ala. These are the Epicat O₂₈-H₂₉...Ala N_{sp³}, Epicat O₂₈-H₂₉...Ala O_{sp²}, Epicat O₂₈-H₂₉...Ala O_{sp³}, Cat O₂₈-H₂₉...Ala N_{sp³}, Cat O₂₈-H₂₉...Ala O_{sp²} and Cat O₂₈-H₂₉...Ala O_{sp³} complexes.

Figure 8 shows the NCI iso-surfaces and RDG (Reduced Density Gradient) plots as a function of $(\text{sign } \lambda_2) \times \rho$ of the Epicat...Ala and Cat...Ala complexes.

In Epicat O₂₈-H₂₉...Ala N_{sp³} and Cat O₂₈-H₂₉...Ala N_{sp³} complexes, NCI analysis suggests the presence of HB and repulsive interaction. The bonds O-H ... N, O-H ... O and C-

H \cdots O are revealed by peak appearance at $-(\text{sign } \lambda_2) \times \rho < -0.010$ a.u. while the repulsive interactions O \cdots O, H \cdots H, C \cdots C have their peaks appearing at 0.010 a.u. $< (\text{sign } \lambda_2) \times \rho < 0.020$ a.u. The blue “NCI” between the H and N; H and O atoms characterize the presence of the previously mentioned O-H \cdots N, O-H \cdots O and C-H \cdots O HB. Similarly, the red-coloured “isosurfaces” between O and O; C and C and within the rings are associated with the

existence of repulsive interactions.

For complexes Epicat O₂₈-H₂₉ \cdots Ala O_{sp}², Epicat O₂₈-H₂₉ \cdots Ala O_{sp}³, Cat O₂₈-H₂₉ \cdots Ala O_{sp}² and Cat O₂₈-H₂₉ \cdots Ala O_{sp}³, research notes that VDW interactions overlap with HB and repulsive ones; the presence of peaks at $(\text{sign } \lambda_2) \times \rho = 0$ attests to this. Moreover, the green-coloured “isosurface” between the hydrogen H atoms reinforces this hypothesis.

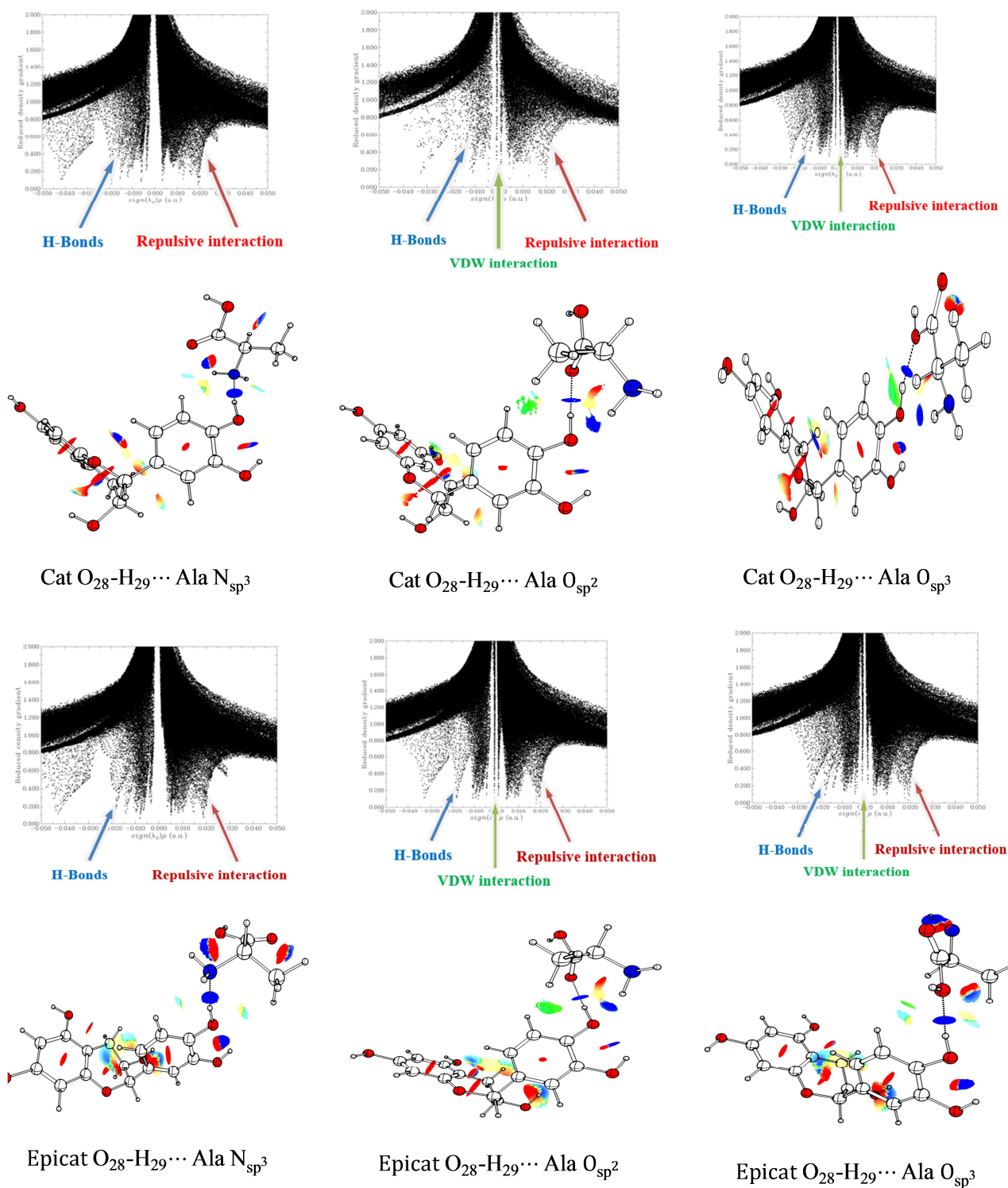


Figure 8. NCI “isosurface” and gradients plot of the RDG as a function of $(\text{sign } \lambda_2) \times \rho$ of some Cat... Ala and EpiCat... Ala complexes.

4. Conclusion

This study focused on the interactions between EpiCat and Ala and between Cat and Ala. According to the optimized geometries of the EpiCat... Ala and Cat complexes... Ala formed, the energies, vibrational frequencies, CT, and topological parameters were analyzed. It aims to identify the nature and category of interactions in the EpiCat... Ala and Cat... Ala complexes. It intends to specify whether they are covalently based on quantitative data. Following the optimization of their geometries, it focuses on the energetic and vibrational aspects associated with these two complexes. It evaluates CT within them and their topological parameters. Right off the bat, the assessment of this data makes it possible to discuss their stability. The enthalpies of formation and free Gibbs enthalpies show that the synthesis of EpiCat... Ala and Cat... Ala is exothermic and spontaneous in the gas phase. Afterwards, NBO analysis reveals the presence of several non-covalent interactions including O – H...O, O – H...N, C – H...O, C – H...N, N – H...O, O – H... π , C – H ... π , N – H ... π . Furthermore, this work establishes that O – H ... O and O – H ... N are the main ones; these possess the highest stabilization energies. Moreover, examination of the complexes vibrational frequencies suggests that O – H...O, O – H...N and N – H...O are appropriate or conventional HB. It proves that C – H ... O and C – H ... N are inappropriate or unconventional HB. In addition, this work is based on the parameters obtained at the various BCP and on the AIM analysis results to categorize these latter of the EpiCat... Ala and Cat... Ala complexes to classify the HB. It demonstrates that O – H...O and O – H...N are weak and partially covalent whereas C – H...O, C – H...N and N – H...O are also feeble and non-covalent. Finally, the study of the stabilization energies $E^{(2)}$, of the CT and of the E_{HB} establishes that the most active sites are the hydroxyl groups O₂₈ – H₂₉, O₃₂ – H₃₃ and O₃₄ – H₃₅ for EpiCat and Cat and the N_{sp³} and O_{sp²} heteroatoms of Ala. These sites lead to the most stable complexes.

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