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M. Tahenti, S. Gatfaoui, N. Issaoui, T. Roisnel, H. Marouani. A tetrachlorocobaltate(II) salt with 2-amino-5-picolinium Synthesis, theoretical and experimental characterization. *Journal of Molecular Structure*, 2020, 1207, pp.127781. 10.1016/j.molstruc.2020.127781 . hal-02472932

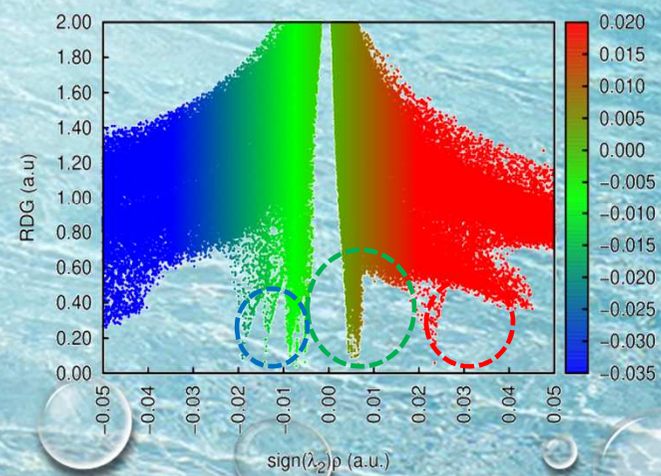
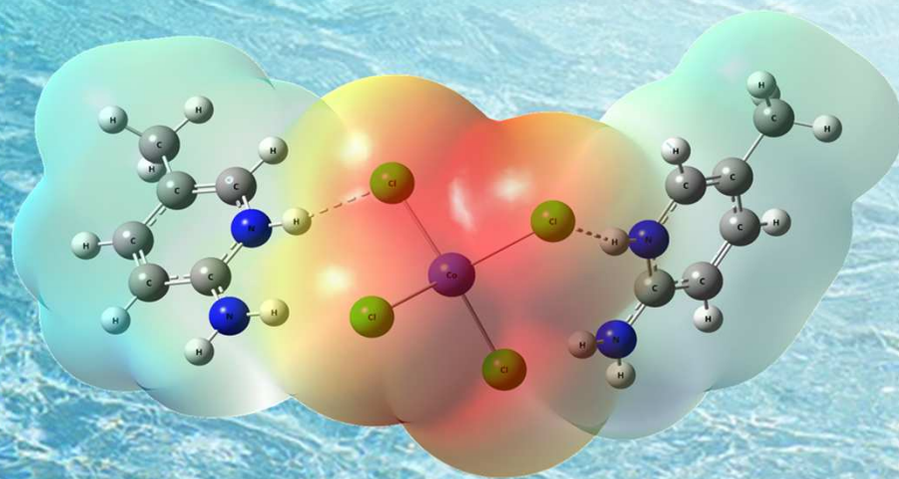
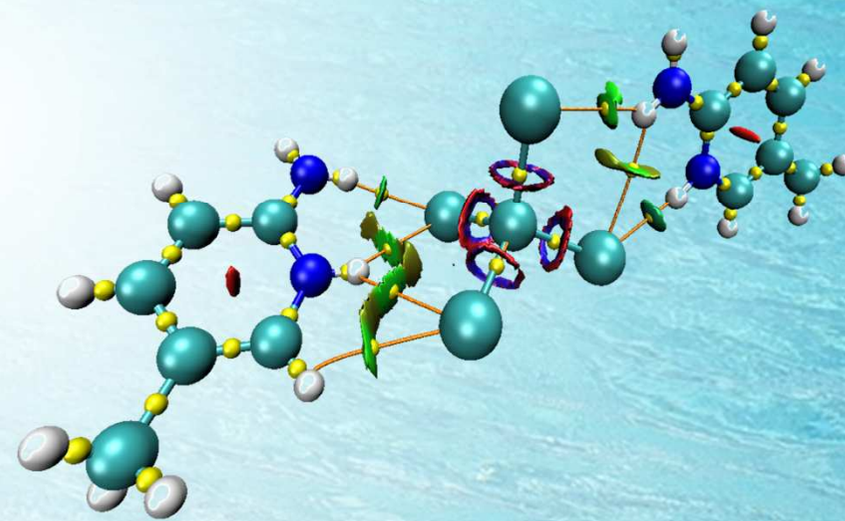
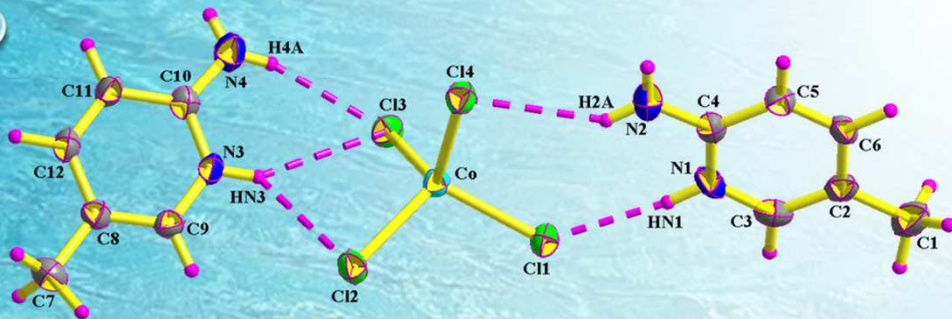
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Submitted on 28 May 2020

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A tetrachlorocobaltate(II) salt with 2-amino-5-picolinium: synthesis, theoretical and experimental characterization

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Abstract

The novel complex, (2-amino-5-picolinium) tetrachlorocobaltate(II), (2A5PCoCl₄) have been synthesized and crystallized to the monoclinic system with space group P2₁/c and the lattice parameters obtained are $a = 8.0734 (6) \text{ \AA}$, $b = 14.6297 (11) \text{ \AA}$, $c = 15.6913 (11) \text{ \AA}$, $\beta = 103.034 (4)^\circ$, $V = 1805.6 (2) \text{ \AA}^3$ and $Z = 4$. His atomic arrangement can be described as an alternation of inorganic and organic layers, [CoCl₄]²⁻ tetrahedral anions and 2-amino-5-picolinium cations are connected by bifurcated and no-bifurcated N-H...Cl and C-H...Cl hydrogen bonds. The intermolecular interactions in the crystal structure were quantified by using the three-dimensional Hirshfeld surfaces and two-dimensional fingerprint plots and interactions were analyzed by topological AIM, RDG and NBO approaches. Infrared spectrum confirms the existence of functional groups and used to gain more information of the elaborated material. The DTA/TG analyses were accomplished to inspect the thermal stability of 2A5PCoCl₄.

Keywords: X-ray diffraction; DFT calculations; Hirshfeld surface; AIM; IR spectroscopy

1. Introduction

An hybrid material is qualified when it comprises both organic and inorganic entities. Hybrids are very interest because of their wide range of technologically advantageous properties, their amazing composition and their exceptional structural diversity.

Thus, following the structural integration of organic cations and inorganic counterparts, electrical [1,2], magnetic [3,4], optical [5], and electroluminescence [6], as well as their ionic properties [7,8] appeared in this class of hybrid chemical systems, and penetrate several application fields such as optic, solid electrolytes, catalysis, biomaterials and biomedical applications [9,10].

The contribution of this work is on hybrid materials base on halogenometalate, have several advantages like include good electric mobility, tunable band gap and magnetic properties, and thermal stability [11] especially that based on cobalt(II).

The coordination numbers of inorganic compound can be three, four (tetrahedral) [12], five (trigonal bipyramidal), or six (octahedral) [13]. To date, salts of general formula $[\text{CoX}_4]^{2-}$ anion and organic cations (where $\text{X}=\text{Cl}, \text{Br}$), have been an interesting area of chemistry and have been extensively studied using spectroscopic in the literature [14], another purpose was to explore thermal [5] and magnetic [15] properties.

Particularly in recent years, much interest has been paid to the study of cobalt(II) halide salts containing tetrahedral CoX_4 geometries. However, the studies concerning this kind of hybrid material is still rare so far except of studies on structural phase transition and magnetic properties of these crystals [16].

In contrast, the hybrids based on cobalt halide possess good solubility and stability. The use of hydrogen bonding interactions with metal salts (such as $\text{N-H}\dots\text{Cl}$ hydrogen bonds) to control the crystal structure product has also recently received attention [14-17].

Considering the attractive and the attributes of halogenocobaltate(II), the present work reports on chemical preparation, crystal structure study of $(\text{C}_6\text{H}_9\text{N}_2)_2[\text{CoCl}_4]$ compound by Hirshfeld analysis, intermolecular interactions were studied by NBO and AIM analyses, infrared spectroscopy, thermal analysis of a novel organic tetrachlorocobaltate(II) have been investigated.

2. Experimental section

2.1 Materials

The cobalt chloride hexahydrate was a commercial product (Sigma–Aldrich), the amine 2-amino-5-picoline was a commercial product (Accros), as well as the Hydrochloric acid (HCl; 37%) was commercial by OXFORD.

2.2 Synthesis of $(C_6H_9N_2)_2[CoCl_4]$

The new compound $(C_6H_9N_2)_2[CoCl_4]$ was obtained by slow evaporation, at room temperature. $(C_6H_8N_2)$ was dissolved with either $CoCl_2 \cdot 6H_2O$ in 10 mL of distilled water and hydrochloric acid HCl (pH < 2.5) with metal/amine molar ratio 1:2. The clear solutions were stirred for 10 min until the complete dissolution and allowed to stand at room temperature. The crystals with the specific color of the metal appeared after few days. Then, the product was filtered off and washed with a small amount of distilled water before being dried in ambient air.



Crystallographic data (CIF) for the structure reported in this paper have been deposited in the Cambridge Crystallographic Data center as supplementary materials N°1942470 CCDC. Copies of the data can be obtained free of charge, from the Cambridge Crystallographic Data centre via www.ccdc.cam.ac.uk/data_request/cif.

2.3. Investigation techniques

2.3.1. X-ray crystallography

Small crystal of $2A5PCoCl_4$, was chosen for measurements on a diffractometer APEXII, Bruker-AXS with $MoK\alpha$ radiation ($\lambda = 0.71073 \text{ \AA}$). Data collection, absorption collections frame scaling and unit cell parameters refinements were carried out with multi scan technique with the SADABS program [18]. The total number of measured reflections was 15135 reflections which 4139 was independent and 3126 reflections with $I > 2\sigma(I)$. The structure an analysis was carried with the monoclinic symmetry, space groups $P2_1/c$, according to the automated search for space group available in WinGX [19]. This structure was solved with direct methods using SHELXL-97 [20]. H atoms bonded to C atoms were positioned geometrically and allowed to ride on their parent atoms, with $C-H = 0.95$. The hydrogen atoms bonded to nitrogen atoms were located from a difference map. The drawing

was made with DIAMOND program [21]. A final refinement on F^2 converged at $R(F^2) = 0.033$ and $wR(F^2) = 0.076$. The main crystallographic data and refinement parameters are presented in **Table 1**. An ORTEP [19] drawing of the molecular structure is shown in **Fig. 1a**.

2.3.2. Physical measurements

IR measurement was performed using a spectrometer. Sample was dispersed with spectroscopic KBr and pressed into a pellet. Scans were run over the range $400\text{-}4000\text{ cm}^{-1}$. The thermal analyses TG-DTA measurements was performed using a multimodule 92 setaram analyzed operating from 313 K up to 880 K in a platinum crucible at an average heating rate of $5\text{K}\cdot\text{min}^{-1}$ and the mass of the sample was 12.4 mg.

2.3.3. Theoretical studies

The intermolecular interactions were also studied by Hirshfeld surface analysis which was carried out using the Crystal Explorer 3.1 software [22] imported on a CIF file. The theoretical calculations have been performed with the Gaussian 09 software package [23] by using the Density functional theory (DFT) with the Becke-three-parameter hybrid exchange functional combined with the Lee-Yang-Parr correlation functional (B3LYP) [24, 25] levels, the 6-311++G(d,p) basis set was used for all atoms except for the cobalt atom, the LANL2DZ basis set and effective core potentials (ECPs) have been used in order to represent the metal. The GaussView was used as molecular visualization program [26]. All the calculations have converged to an optimized geometry that corresponds to the lowest energy. Positive vibrational frequencies obtained confirming the stability of the optimized geometries.

The vibrational assignments were performed considering the potential energy distribution components (PED) $\geq 10\%$ using the VEDA4 package [27]. The GaussView package was used to verify the assignments of the bands. In this study, NBO calculations [28] were employed to determinate the energy of intermolecular interactions, furthermore topological proprieties were determined with the Mutiwfn program [29] in accordance with the Bader's theory [30]. The results were plotted by the use of VMD programs [31].

3. Results and discussion

3.1. X-ray diffraction

The asymmetric unit of the title material, $C_{12}H_{18}N_4^{2+} \cdot CoCl_4^{2-}$ contains one tetrachlorocobaltate anion and two 2-amino-5-picolinium cations. The optimized geometry model of this compound is illustrated in **Fig. 1b**.

The Co(II) ion is tetrahedrally bound by four chlorine atoms, with Co–Cl bond distances ranging from 2.2473 (8) to 2.2953 (7) Å and Cl–Co–Cl bond angles between 105.23 (3) ° and 113.76 (3)° which are slightly deviated from the ideal value of 109.44° (**Table S1**). In the other hand, we propose a very simple geometry index in four-coordinate complex, τ_4 (eqn (1)).

$$\tau_4 = \frac{360 - (\alpha + \beta)}{141^\circ} \quad (1)$$

The formula for our four-coordinate index τ_4 is simply the sum of angles α and β the two largest θ angles in the four-coordinate species subtracted from 360°, all divided by 141°. Indeed, the four-coordinate cobalt(II) geometry index [32] provides a slightly distorted tetrahedral geometry of $[CoCl_4]^{2-}$.

$$\tau_4 = \frac{360 - (113.76 + 111.63)}{141} = 0.95$$

Indeed, the calculated average values of the distortion indices corresponding to the different angles and distances in the $CoCl_4$ tetrahedra according to the method described by Baur [33] are: $[DI(CoCl) = 0.0372; DI(ClCl) = 0.00632$ and $DI(ClCoCl) = 0.01976]$. The comparison of the distortion indices shows a small distortion of the Cl–Co–Cl angles with respect to the Co–Cl distances from which the $[CoCl_4]^{2-}$ anion has a tetrahedral geometry. It can be admitted that the $CoCl_4$ tetrahedron is formed by a regular arrangement of chlorine atoms, in which the cobalt atom is slightly displaced from the center of gravity of the tetrahedron.

The projection in the direction of the \vec{a} axis (**Fig. 2a**), can be described by a succession of anion-cation layered arrangement which alternate along of the \vec{c} axis and unfold parallel to the \vec{b} axis. On the same projection we can also notice that the mineral anions develop in zigzag along the \vec{b} axis in $z = 1/4$ and $3/4$, and the organic cations propagate in inclined rows along the \vec{c} axis $z = 1/2$. The cohesion of these layers is ensured by medium and weak hydrogen bonds which connect the $[CoCl_4]^{2-}$ and the 2-amino-5-picolinium cations. To

better understand the mode of arrangement of organic and inorganic entities in such an atomic arrangement, the structure is still projected in the direction of the axis \vec{c} (**Fig. 2b**). The projection of the anionic part $[\text{CoCl}_4]^{2-}$ on the (ab) plane shows that the latter develops in a chain according to the two directions \vec{a} and \vec{b} , in the same figure we notice that each anion $[\text{CoCl}_4]^{2-}$ connects to four other anions through type hydrogen bonds N-H...Cl thus forming a type of graphs $R_3^7(22)$ (**Fig. 2c**). The interatomic distances and the bonding angles of (CoCl_4) collected in **Table S1** do not show any particularity compared to those generally encountered in organic tetrachlorocobaltate known until now [34]. Concerning the organic cation, the geometric characteristics classified in **Table S1** are found in agreement with those observed for the same cation crystallizing with other types of anions [35]. The Hydrogen bonds N-H...Cl involving the hydrogen atoms of the ammonium groups of the organic entities to the chlorine atoms of the anionic group (**Table 2**) their distances $d_{\text{N}\dots\text{Cl}}$ are varying between 3.214 (2) Å and 3.478 (2) Å. We notice the appearance of the graphs $R_2^2(8)$, $R_1^2(4)$ and $R_2^1(6)$. Three bonds of type C-H...Cl attach the organic groups and the anionic groups $[\text{CoCl}_4]^{2-}$ their average distance $d_{\text{C}\dots\text{Cl}}$ is 3.719 Å.

The geometric parameters studied experimentally by X-ray diffraction, and theoretically by DFT(B3LYP/LANL2DZ), such as the bond lengths and the angles of the 2A5PCoCl₄ are tabulated in **Table S1**. Optimized molecular structure gives a comparison between the optimized geometric parameters and experimental X-ray crystallographic parameters. From this table, the C-C linkages in the pyridine of the 2-amino-5-picolinium group were calculated at the range of 1.384 Å-1.512 Å which is compatible with other pyridine rings [36]. The average value of calculated bond length of C-H an N-H is equal to 1.1 Å relative to the mean value observed by experimentally of 1.01. As regards the anionic group, we have a slightly difference in values because the investigational results characterize molecules at solid state; however the theoretical calculations refer to isolated molecules in the gas phase. Lastly, it can be noted that in spite of the differences between the experimental and optimal results, the second represent an excellent estimation.

3.2 Hirshfeld surface analysis

The Hirshfeld d_{norm} surface of 2A5PCoCl₄ is illustrated in **Fig. 3a** also the shape index and the curvedness cartography are shown respectively in **Figs. 3b** and **3c**. This analysis can be done by quantitative calculation of 2D fingerprint plots through the CrystalExplorer 3.1 program [22]. The full fingerprint plot and the fingerprints plots of the main contacts involved

in this compound are illustrated in **Fig. 4**. At each point of the Hirshfeld surface is associated a d_e and d_i value. For a pair (d_i , d_e), the sum of these components is equal to the distance separating the atoms contributing to this point.

The interatomic contacts H...Cl/Cl...H have the greatest contribution to the Hirshfeld surface (42%), attributed to the N-H...Cl and C-H...Cl hydrogen-bonding interactions and represented by two sharp symmetric spikes in the two-dimensional fingerprint maps, with a maximum $d_e + d_i \sim 2,30 \text{ \AA}$ (**Fig. 4a**). This value is less than the sum of van der Waals radii of chlorine ($1,75 \text{ \AA}$) and hydrogen ($1,09 \text{ \AA}$) atoms; it confirms that the inter-contact is considered as being close contact. The fingerprint plots corresponding to the contacts C...C (1.9%) (**Fig. 4f**) shows that the sum ($d_i + d_e \sim 2.5 \text{ \AA}$) smaller than the sum of the van der Waals radii of the carbon atoms (3.4 \AA) confirm the presence of π - π interactions in the crystalline stack. This is also confirmed by the presence of the red and blue triangles on the Shape index cartography (**Fig.3b**) and in the plane regions delimited by a blue outline on the Hirshfeld surfaces mapped with Curvedness (**Fig. 3c**).

The enrichment ratio (ER) of a pair of elements is defined as the ratio between the percentage of the actual contacts in the crystal and the theoretical percentage of the random contacts [37]. The set of enrichment ratios of the various intermolecular contacts occurring in the 2A5PCoCl₄ compound are recorded in **Table S2**. The list of enrichment ratios highlights the contacts H ... Cl / Cl ... H ($ER_{HCl} = 1.47$) clearly provides evidence of the formation of hydrogen bonds of type N-H ... Cl and C-H ... Cl. The H...H contacts are the second most frequent interactions due to the large quantity of hydrogen on the molecular surface (34.2 %), but they have too small degree figured with an enrichment ratio of 0.84. The ER relates to the contacts of Cl...Cl and Cl...C/C...Cl are of low meaning as they are derived from less important interactions with small contributions in the all parts Hirshfeld surface. The contacts C ... H / H ... C have an enrichment ratio equal to 0.98, which also proves the presence of the interactions C-H ... π [37, 38].

Such visual analysis for intermolecular interactions is coherent with those noticed by the X-ray diffraction analysis.

3.3. AIM topological, reduced density gradient (RDG) and NBO analyses

The AIM approach consists of defining the different properties of chemical bonds, including hydrogen bonds. For this purpose, the critical point BCP is used for the

identification of the chemical bonds between the atoms and the interatomic interaction. Geometric and topological parameters are an effective way to characterize the strength and nature of the H bond, but these criteria for the existence of hydrogen bonds are often considered insufficient, so we must rely on Koch and Popelier [39] to confirm the existence of a hydrogen bond, therefore these criteria are the existence of a critical binding point for the contact of the proton acceptor H (A), thus confirming the existence of a hydrogen bonding interaction, these criteria are the existence of a critical binding point for the proton (A) acceptor contact (H) confirming the existence of a hydrogen bonding interaction, the value of the electron density $\rho(\text{H}\dots\text{A})$ must be in the range of 0.002 to 0.004 a.u., the corresponding Laplacian is positive in the range of 0.024 to 0.0139 a.u according to Rozas et al. [40], the interaction can be classified as follows:

- Strong H bonds are characterized by $\nabla^2\rho(\text{BCP}) < 0$, $H(\text{BCP}) < 0$, $E_{\text{HB}} > 24.0$ Kcal/mol and their covalent character is established.
- Moderate H bonds are characterized by $\nabla^2\rho(\text{BCP}) > 0$, $H(\text{BCP}) < 0$, $12.0 < E_{\text{HB}} < 24.0$ Kcal/mol and their partially covalent character is established.
- The weak H bonds are characterized by $\nabla^2\rho(\text{BCP}) > 0$, $H(\text{BCP}) > 0$, $E_{\text{HB}} < 12.0$ Kcal/mol and they are mainly electrostatic and the distance between the interacting atoms is greater than the sum of the van der Waals rays of these atoms.

The graphical representation of AIM analysis of 2A5PCoCl₄ is illustrated in **Fig. 5**, whereas the calculated topological parameters are listed in **Table S3**. According to this table, Cl36 ... H9, Cl37 ... H24, Cl37 ... H26, Cl37 ... H17, Cl38 ... H24, Cl38 ... H33 and Cl39 ... H17 are weak hydrogen bonds. AIM analysis shows that our material is stable by two hydrogen bonds Cl ... H. In this study, the BCPs along Cl...H and the Laplacian of the total electron density $\nabla\rho(r_{\text{BCP}})$ are between 0.0238 and 0.0576 a.u with electron energy densities $H(r_{\text{BCP}})$ between 0.0009 and 0.0284 a.u respectively for the two hydrogen bonds N32...H33 and N25...H26. The energy of the hydrogen bond is calculated using the Espinosa relation [41]: $E_{\text{Cl}\dots\text{H}} = 1/2 V_{\text{BCP}}$. Therefore the hydrogen bond energies of our compound vary between -14.10 and -3.24 kJ/mol. In our case, based on the energy density values of the topological parameters, it can be concluded that the bonds are weak bonds since the values of the Laplacian and the density of the total energy are positive. The AIM confirms the presence of the H...H contacts already observed by Hirshfeld surface analysis, so the existence of its intermolecular interactions is also a proof of the critical points BCP. In addition, the presence

of a new ring critical point 'RCP' confirms the cyclic character of an atomic chain. As shown in **Fig. 5**, the interactions between the organic group and the $[\text{CoCl}_4]^{2-}$ anion via hydrogen bonds give rise to the formation of the rings NRCP1 and NRCP2.

The existing intermolecular interactions in 2A5PCoCl_4 compound experimentally processed by X-ray diffraction and confirmed theoretically by Hirshfeld surface analysis, geometric optimization and by AIM topological analysis, have been also validated and evaluated using the multi-wavefunction analysis, this latter was carried out by a RDG plot analysis and isosurface of interactions. The RDG approach is a dimensionless elementary amount obtained from the density and its first derivative developed by Johnson et al. [42]:

$$\text{RDG}(r) = \frac{1}{2(3\pi^2)^{1/3}} \frac{|\nabla\rho(r)|}{\rho(r)^{4/3}}$$

Based on the density $\rho(r)$ against sign λ_2 we have tried to understand and indicate the nature of strength of the different interactions existing in our crystalline structure:

- If $\lambda_2 < 0$: Attractive and binding interactions (hydrogen bonds).
- If $\lambda_2 > 0$: repulsive and non-binding interactions (steric effect in ring and cage).
- Values close to zero indicating van der Waals interactions.

The results were collected in **Figs. 6a** and **6b** and illustrate the attractive, van der Waals and repulsive interactions in our compound structure. The interactions appear in the form of green plates are attributed to van der Waals interactions; its plates are located exactly between the hydrogen and the chlorine atoms. The elliptic red plate located at the center of the aromatic nucleus [43] is related to $\pi - \pi$ stacking interactions between the pyridine rings and show a strong steric effect. The green plates signify the strong attractive interaction N-H ... Cl and C-H ... Cl. The RDG peak at -0.22 corresponds to a strong interaction between chlorine and hydrogen, which was also confirmed by the blue color in the color scaling. These results are comparable to those found in the AIM calculation and in other previous researches [44-45].

Natural bond analysis (NBO) is a powerful method that allows us to study and identify the specific bonds and energies associated with one-pair electrons, which play a very important and essential role in physico-chemical processes [46]. This method has been used to establish electronic exchanges, transfer reactions, hydrogen bonding between donor-acceptor

compounds and hyperconjugative interactions. The difference of energy between interacting orbitals give an idea for the interaction force between acceptor electrons and donor electrons or otherwise the stability of the orbital interactions. The most stable interaction can be estimated by calculating the second-order stabilization energy $E^{(2)}$ [47]:

$$E^{(2)} = q_i \frac{F^2(i, j)}{\varepsilon_i - \varepsilon_j}$$

Where q_i is the occupation of i orbital, ε_i and ε_j are the diagonal elements of orbital energies i and j and $F(i, j)$ is the Fock matrix element outside diagonal. The value of $E^{(2)}$ energy is directly related to the intensity of interaction between donor and electron acceptor, more $E^{(2)}$ is large more the interaction between donor and electrons acceptor is intense.

Table S4 presents the stabilization energies $E^{(2)}$, which indicate the electron delocalization from the donor NBO to acceptor. It is noticed that hydrogen bond interactions between the lone pair LP (C136) orbital and the σ^* (N8-H9) antibonding orbital with stabilization energy equal to $9.71 \text{ kcal mol}^{-1}$, this is reflected by the strong interaction between the anionic and cationic groups which is manifested by the formation of a hydrogen bond N8-H9...C136. The stabilization energy $E^{(2)}$ of the hyperconjugant interactions of pyridine rings associated to π (N8-C10) $\rightarrow \pi^*$ (C5-C6) and π (N25-C27) $\rightarrow \pi^*$ (C22-C23) interactions is in the order of 11 kcal mol^{-1} . Finally, we can conclude that the NBO analysis confirms that the hydrogen bonding in the 2A5PCoCl₄ compound leads to his crystal structure.

3.4 Frontier molecular orbital analysis

In our study, we generally not focus on a comparison between the MOs but instead just take the MOs as a set of orbitals for the whole molecule. So we will always focus simply on the electrons which are the most important. Indeed, there are the Highest Occupied Molecular Orbital (HOMO) which is the highest energy MO that has any electrons in it and the lowest un-Occupied Molecular Orbital (LUMO) is the next highest energy orbital, it will be empty and is the lowest energy place to put or excite an electron. These orbitals play an important role in chemical stability of the molecular edifice. The difference in energy between the HOMO and LUMO is called the HOMO–LUMO gap. This latter between these two frontier orbitals can be used to predict the strength and stability of transition metal complexes and can tell us about what wavelengths the compound can absorb. A high gap indicates a high stability of the molecule. Low gap energy is criterion of high reactivity. The picture

representations of the (HOMO, HOMO-1 and LUMO, LUMO+1) and the contribution of the organic and inorganic units to these orbitals of 2A5PCoCl₄ are mapped in **Fig. 7** and the results of this survey are gathered in **Table S4**. The red and the green colors of the MO plot show the positive and negative phase, respectively. It is observed that the highest occupied orbitals are localized on the organic cation but the lowest unoccupied orbital's is located on the anionic part. The calculated energy values of HOMO and LUMO are around -5.37 eV and -2.49 eV respectively. The energy gap between the HOMO and LUMO is +2.88 eV. This value agree very well with the corresponding energy gap reported earlier of others organic tetrachlorocobaltate(II) compounds [48-49].

3.5. Vibrational IR spectra and assignments

For the determination of vibrational characteristics of 2A5PCoCl₄ we add a vibrational study using infrared spectroscopy at room temperature. The experimental infrared wavenumber compared to the calculated from DFT method assigned by VEDA4 program [27] and GaussView software [26] were listed in **Table S5** along with a tentative assignment of the observed bands. In this table, we have presented the scaled and the unscaled calculated harmonic vibrational frequencies, infrared intensity and detailed PED assignment. In our study, wavenumbers are in the ranges from 4000 to 400 cm⁻¹. The experimental IR spectrum according the calculated is illustrated in **Fig. 8**. By reference to the results of the assignment and to the bibliographic study [50-54] we have ascribed the organic cation to the set of bands that appeared in IR spectrum.

3.5.1 Vibrational modes of the anion

The compound crystallizes in the orthorhombic system P2₁/c, the elemental cell contains four form units occupying four crystallographically equivalent sites. The [CoCl₄]²⁻ anion has C₁ site symmetry. Work perform on infrared absorption spectroscopy allowed us on one hand to determine the vibrational characteristics of the atomic groups of compound, and the enumeration of the modes of vibration of the [CoCl₄]²⁻ groups on the other hand. The tetrahedral anion [CoCl₄]²⁻ contains 5 atoms so it has (5 x 3) - 6 = 9 modes of vibration. The six subtracted modes correspond to the three rotations (Rx, Ry, Rz) and to the three translations (Tx, Ty, Tz) of the whole ion. Based on the correlation method we find that the CoCl₄²⁻ anion has 9(A_u+B_u+A_g+B_g) active modes in IR and 9(A_u+B_u+A_g+B_g) modes active in Raman, they are shared in four fundamental modes ν_1 , ν_2 , ν_3 et ν_4 . We have distinguished the four fundamental modes (ν_1 , ν_2 , ν_3 and ν_4) of the [CoCl₄]²⁻ anion in the wave number region

from 50 to 300 cm^{-1} . In fact, symmetric (ν_2) and asymmetric (ν_4) stretching vibrations are attributed to the Raman peaks observed at 107 and 132 cm^{-1} , respectively. The same stretching vibration is predicted at 1111 and 137 cm^{-1} by DFT method. Their corresponding bending vibration modes are respectively 220 and 276 cm^{-1} . This mode is theoretically calculated between 237 and 268 cm^{-1} by DFT method.

3.5.2 Vibrational modes of the cation

Based on the work reported above we have estimated the vibration domain of the organic group which is located between 541 and 3324 cm^{-1} (**Fig. 8**) and shows the bands corresponding to the characteristic vibrational modes of 2A5PCoCl₄ which compared with those of similar salts.

In fact, the intense bands in 3324 and 3211 cm^{-1} correspond to the valence vibrations of the groups ($-\text{NH}^+$), ($-\text{NH}_2$) and (C-H), so the broadening of these bands is due to the establishment of the hydrogen bonds, which effect the lowering of the frequencies of these groups. The same vibrations are calculated between 3231 and 3697 cm^{-1} by DFT level. The bands between 1674 and 1588 cm^{-1} are attributed to the valence vibrations ν (C = C) of the aromatic ring and the symmetric and asymmetric vibrations of the ($-\text{NH}^+$), ($-\text{NH}_2$) and ($-\text{CH}_3$) groups. The corresponding bands are calculated at range 1593 and 1708 cm^{-1} . In the range 1355-1155 cm^{-1} , the bands attributed to the valence vibrations of the group ν ($\text{C}_{\text{aryl}}-\text{C}$) and $\nu(\text{C}-\text{N})$ appear, nearly at the calculated one among 1155 and 1355 cm^{-1} .

Finally, the observed weak band which is located towards 996 cm^{-1} and 514 cm^{-1} , and between 499 and 1017 cm^{-1} in calculated IR spectrum, are attributed to the (C-H) group of the pyridine ring its off-plane deformation.

3.6 Molecular Electrostatic Potentials (MEP)

The Molecular Electrostatic potentials (MEP) play a key role in the characterization of interaction energy between the electric charge produced by the electrons and the nuclei of the molecule. MEP is one of the best suitable tools for prediction of the presence of inter- and intramolecular interactions of the studied compounds. It is a useful descriptor to determine the active sites of electrophilic and nucleophilic attacks over the molecule for the study of biological recognition process as well as hydrogen-bonding interactions [55]. The MEP surface of 2A5PCoCl₄ was made by using the software Gauss view (**Fig. 9**). The mapping represent the donor atoms in blue regions with positive potential, is the most positive

electrostatic potential (nucleophilic site) and it is localized hydrogen atoms and the acceptor atoms with negative potential in red regions represents the most electronegative potential (electrophilic sites) and it is mainly localized over anionic group (red, chlorine atoms), whereas the green or yellow colors indicates the neutral region [56]. This graph shows that the electrophilic sites are located around the hydrogen atoms, whereas the nucleophilic sites are located around the chlorine atoms. MEP surface represents a proof of the formation of the hydrogen bond [57]; it is defined as being a donor acceptor interaction. It proves that the electrostatic potential of the donor becomes less negative as the acceptor becomes more negative.

3.7 Thermal behavior

To understand the behavior of the title compound, thermal analysis DTA/TG (**Fig. 10**) of $2A5PCoCl_4$ carried out shows that our compound undergoes a phase transition at 327 K. Then a series of endothermic peaks are observed on the DTA curve accompanied by a weight loss on the TG curve which can be attributed to the pyrolysis of the organic molecule. Finally, an endothermic peak without loss of mass appears at 521 K which can correspond to melting. This phenomenon is further confirmed by the heating of our compound on a Kofler bench which makes it possible to estimate the melting temperature of a material, this temperature 517 K is close to that found on the DTA curve. After that the remaining product undergoes a total combustion which leads to a volatile gas and a black carbonaceous residue.

4. Conclusion

The present study reports on the synthesis and subsequent characterization of a new organic tetrachlorocobaltate(II), $2A5PCoCl_4$. This novel Co(II) complex has been successfully synthesized at room temperature by slow evaporation solution growth technique in the monoclinic system with the $P2_1/c$ space group. The structural level the atomic arrangement can be described by the alternation of organic and inorganic layers. The anionic arrangement is built up by tetrahedral of $[CoCl_4]^{2-}$ arranged in sandwich between the organic layers, interconnected by means of the ionic N–H...Cl hydrogen bonds. The crystal packing is stabilized by intermolecular H-bonding and π - π stacking interactions which were confirmed by topological AIM approaches. Moreover, the Hirshfeld surface analysis reveals the percentage of intermolecular contacts and highlights the non-covalent interactions in $2A5PCoCl_4$. The TG/DTA thermal analysis was performed to establish the thermal stability of the crystal. The vibrational wavenumbers of the fundamental modes of $2A5PCoCl_4$ have been

precisely assigned, analyzed, and the theoretical results were compared with the experimental values. NBO analysis explained the stability of the synthesized compound. HOMO-LUMO, energy gap, molecular hardness, ionization energy, electron affinity were predicted by using DFT level.

Acknowledgements

This work was supported by the Ministry of Higher Education and Scientific Research of Tunisia.

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- A new Co(II) complex compound, $2A5PCoCl_4$, was prepared at room temperature.
- The atomic arrangement of the $2A5PCoCl_4$ shows three-dimensional network.
- The optimized geometry and vibrational wavenumbers are computed using DFT(B3LYP/LANL2DZ) method.
- $2A5PCoCl_4$ was characterized by X-ray, FT-IR and DTA/TG analyses.
- The intermolecular interactions were analyzed by Hirshfeld surfaces, AIM, NBO, RDG and MEP.

Figure captions

Fig. 1. *ORTEP* of 2A5PCoCl₄ with atoms-labeling scheme. Displacement ellipsoids are drawn at the 30% probability level. Hydrogen atoms are represented as small spheres (a) and the optimized molecular structure (b).

Fig. 2. Projection along the \vec{a} axis of atomic arrangement of 2A5PCoCl₄ (a). Projection along the \vec{c} axis showing the interactions between inorganic and ammonium groups in 2A5PCoCl₄ (b). Hydrogen bond motifs observed in 2A5PCoCl₄ (c).

Fig. 3. Hirshfeld surfaces mapped with d_{norm} (-0.395 – 1.151) (a) (dotted lines “red” represent hydrogen bonds), shape index (-1 – 1) (b) and curvedness (-4 – 4) (c) for 2A5PCoCl₄.

Fig. 4. Fingerprint plots of the full and the significant intercontacts and percentage of various intermolecular contacts contributed to the Hirshfeld surface in 2A5PCoCl₄ compound.

Fig. 5. AIM molecular graphic showing the different critical points of 2A5PCoCl₄ (BCP: small red sphere, RCP: small yellow sphere, bond paths: yellow lines).

Fig. 6. Representation of different types of interactions in 2A5PCoCl₄ compound (a). The map of Reduced Density Gradient (RDG) defines the interaction limits for 2A5PCoCl₄ (b).

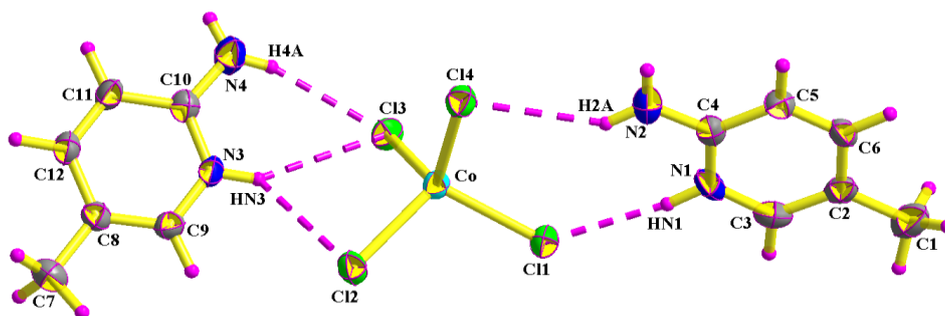
Fig. 7. The frontier molecular orbitals of 2A5PCoCl₄ computed with TD-DFT (B3LYP)/6-311++G(d,p) method.

Fig. 8. Theoretical (blue) and experimental (red) IR spectra of 2A5PCoCl₄.

Fig. 9. Molecular Electrostatic Potentials (MEP) surface mapped of 2A5PCoCl₄.

Fig. 10. DTA and TG curves of 2A5PCoCl₄.

(a)



(b)

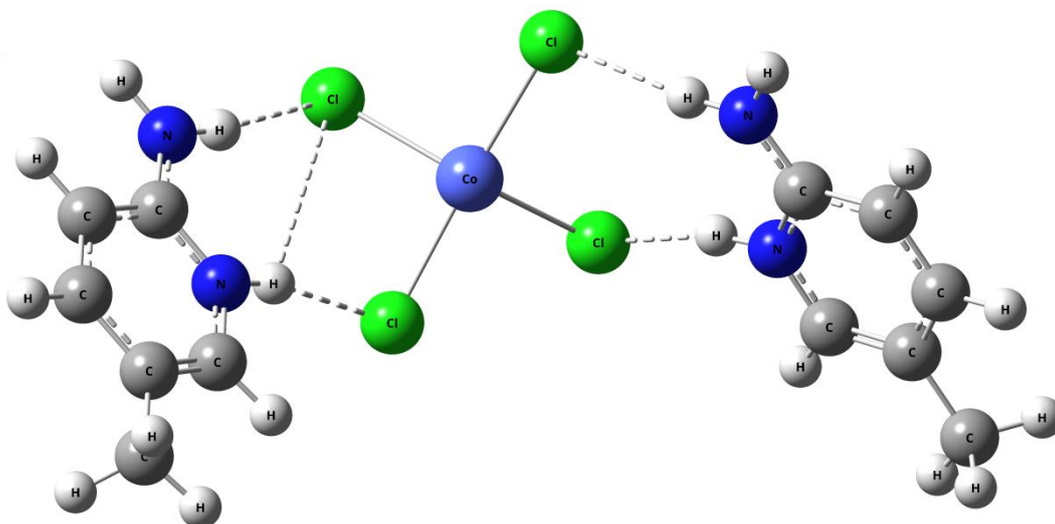


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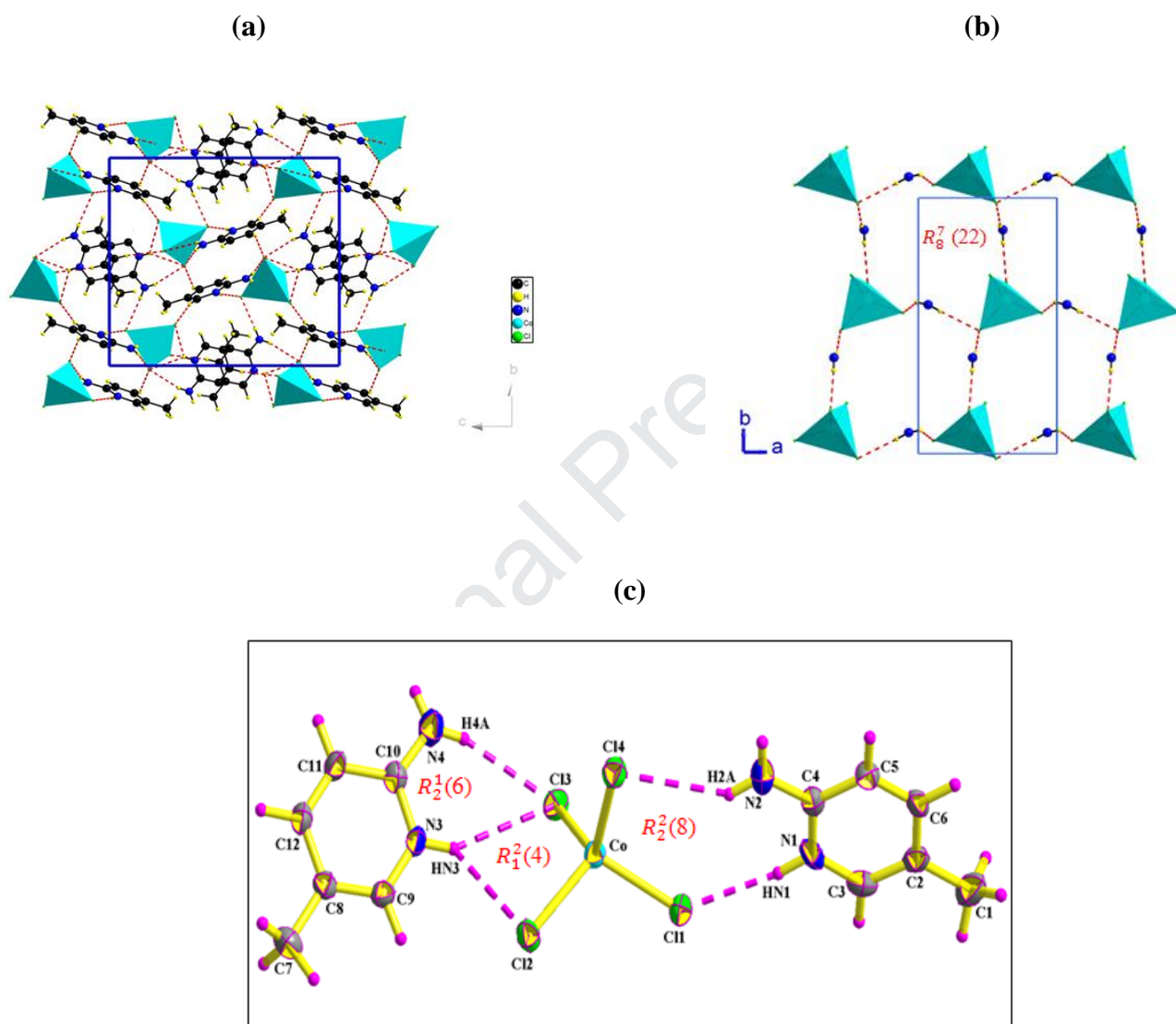


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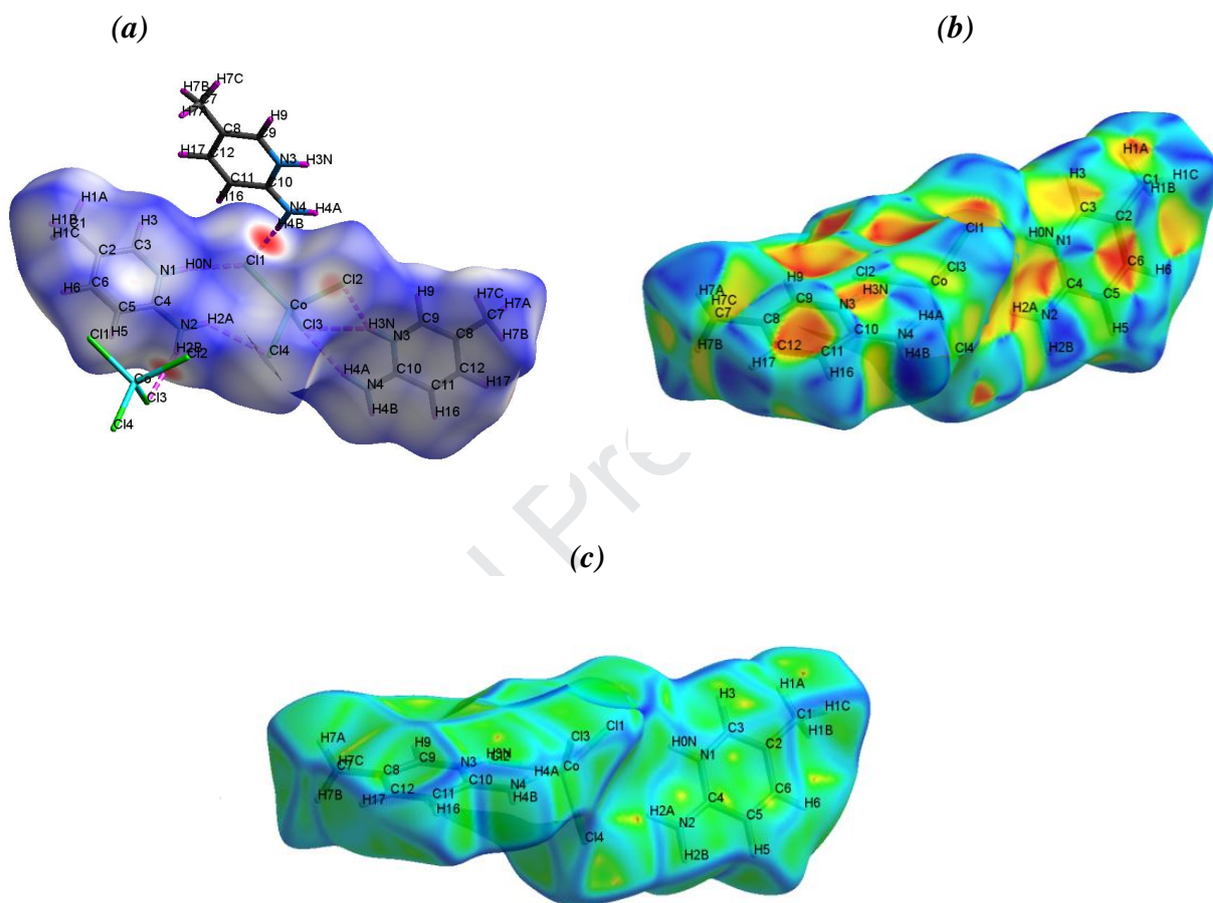


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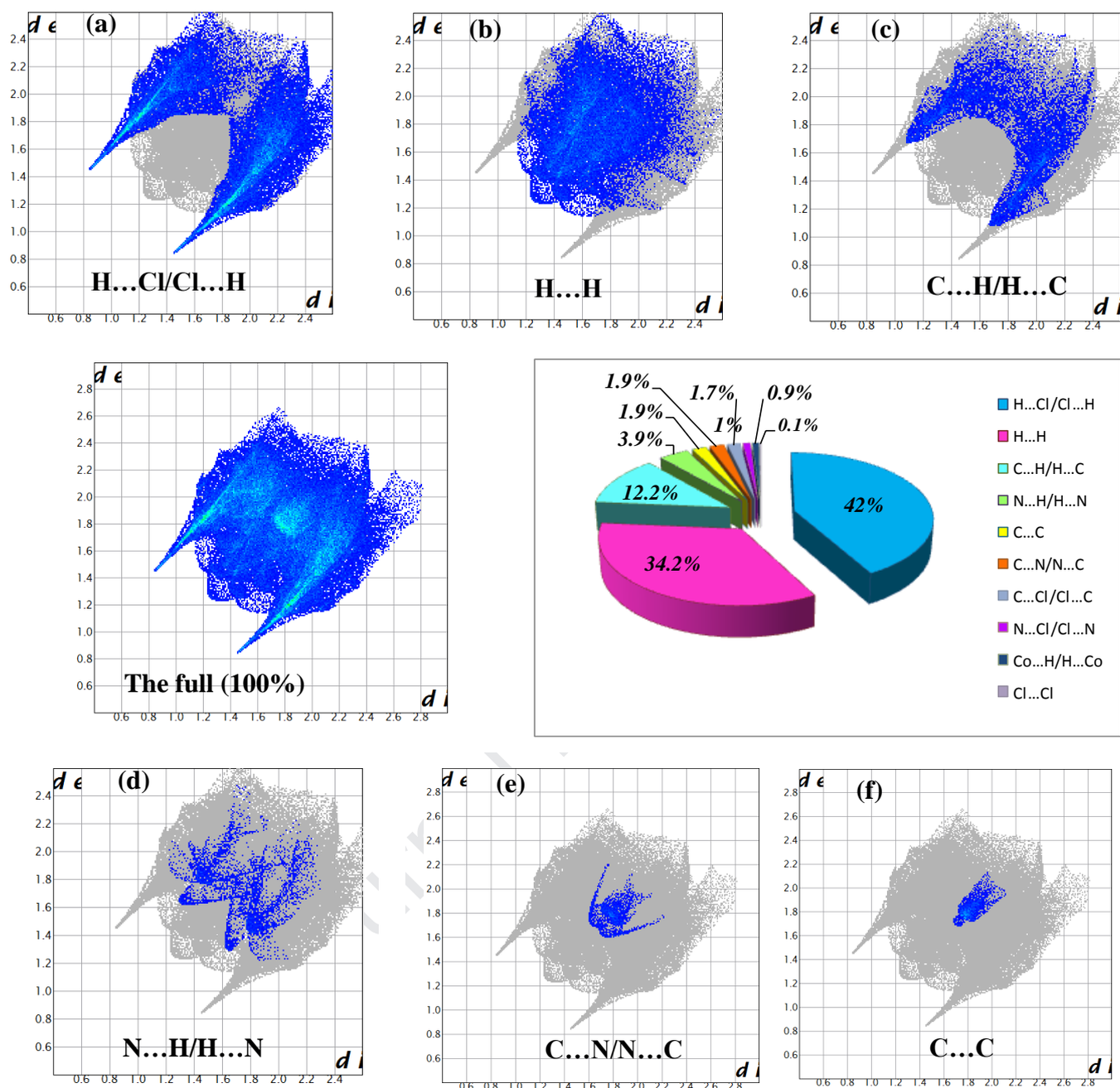


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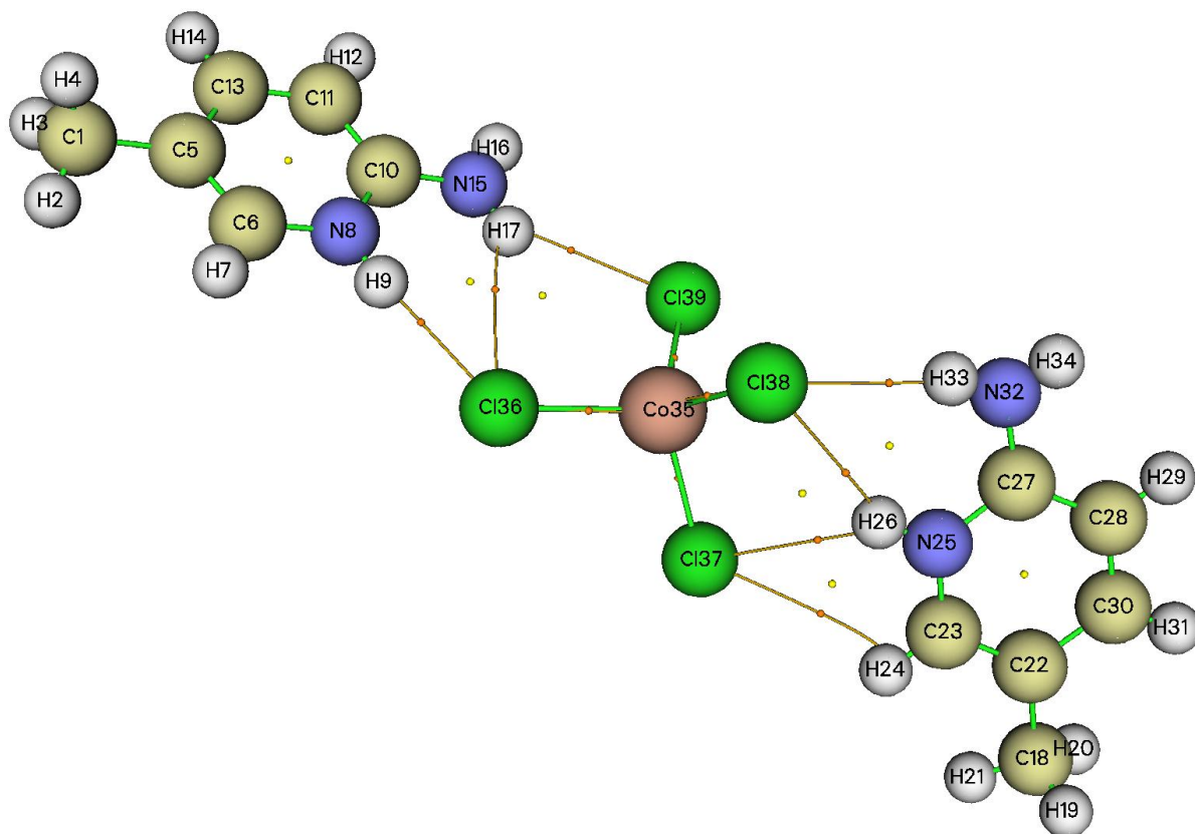


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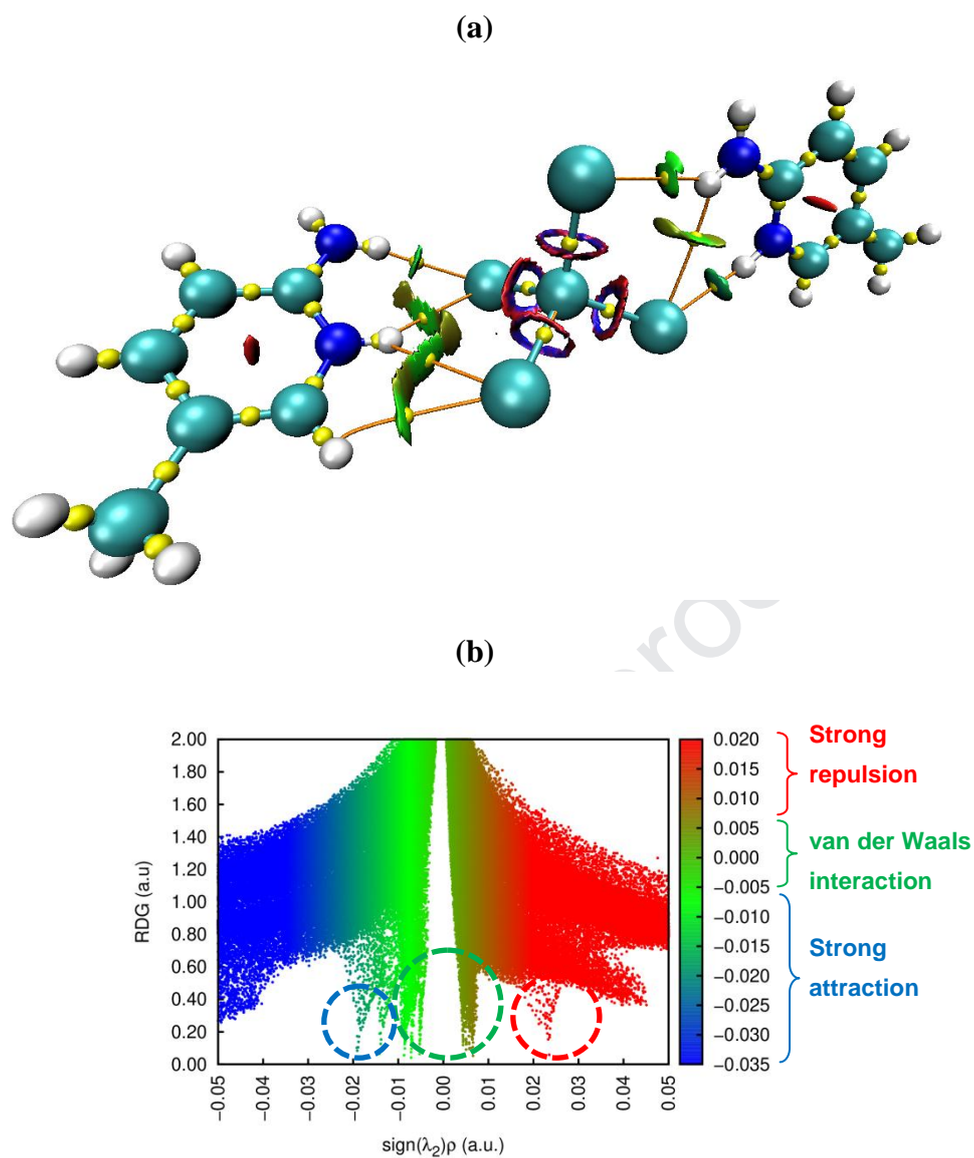


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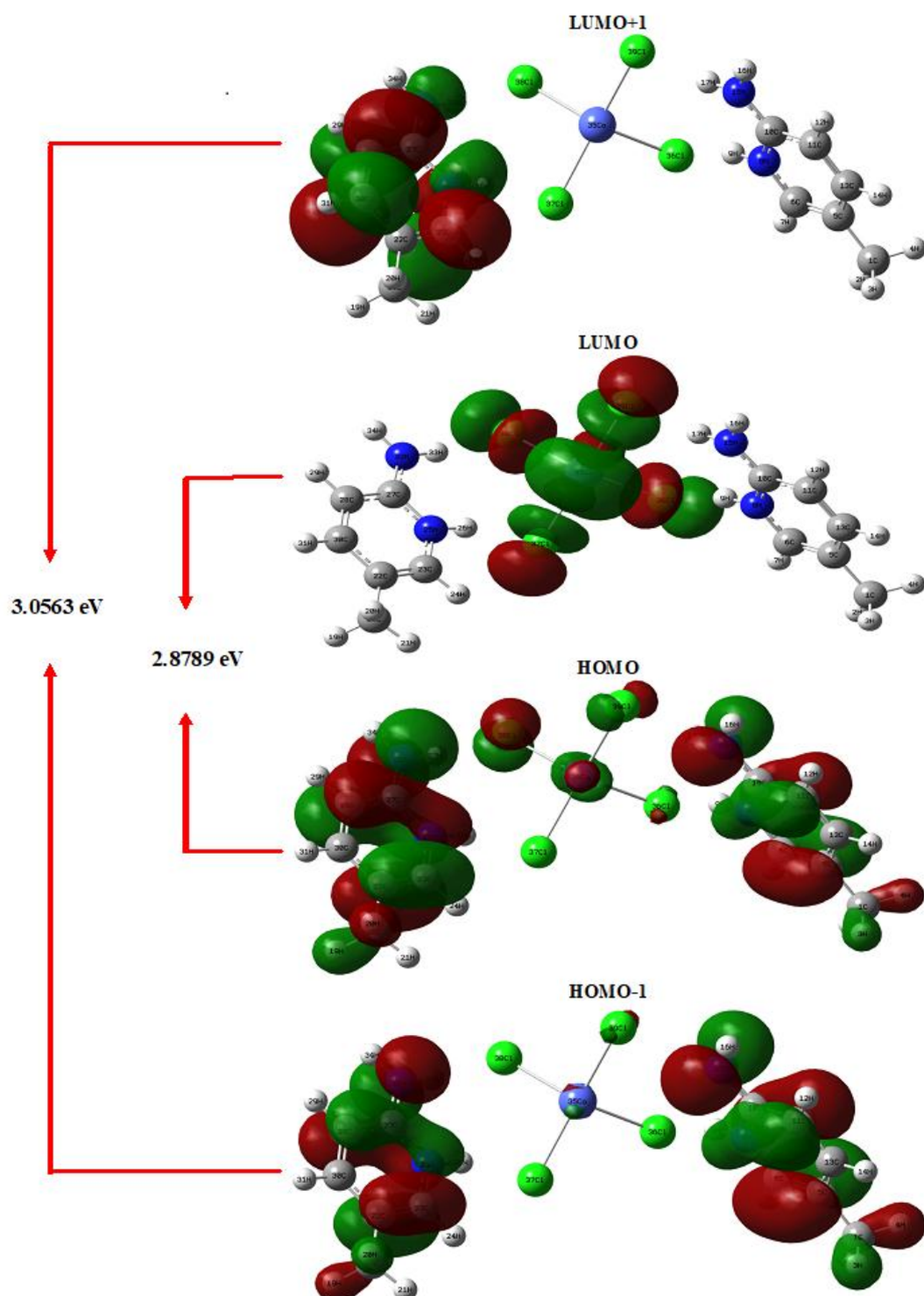


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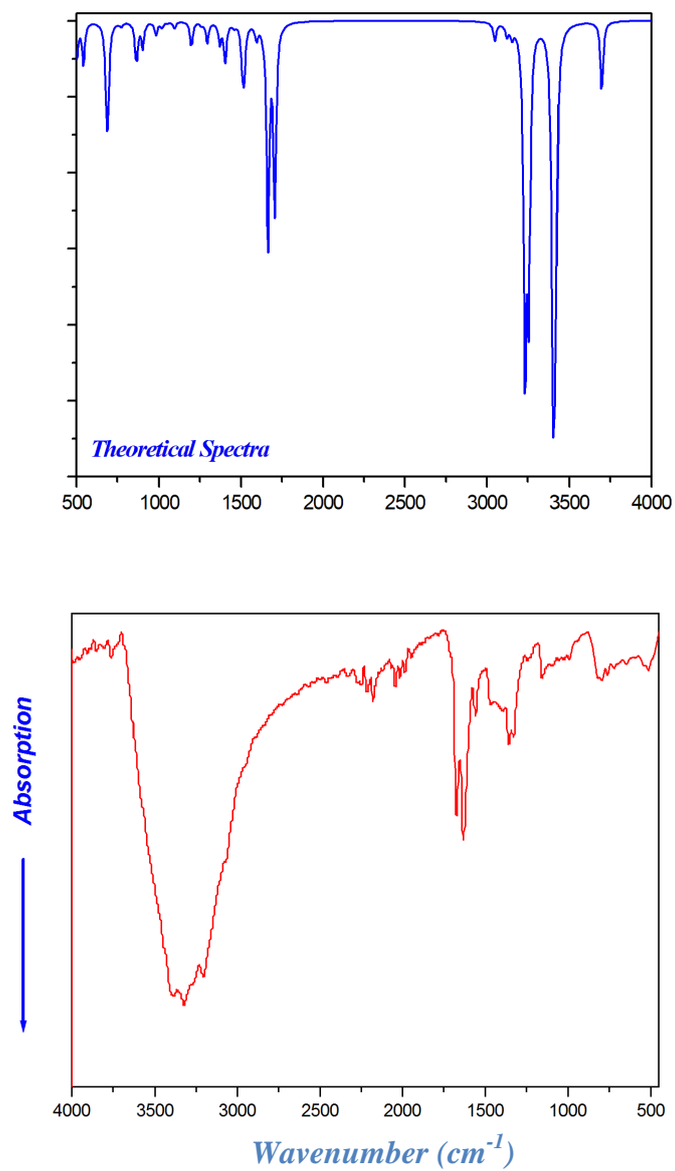


Fig. 8. Theoretical (blue) and experimental (red) IR spectra of 2A5PCoCl₄.

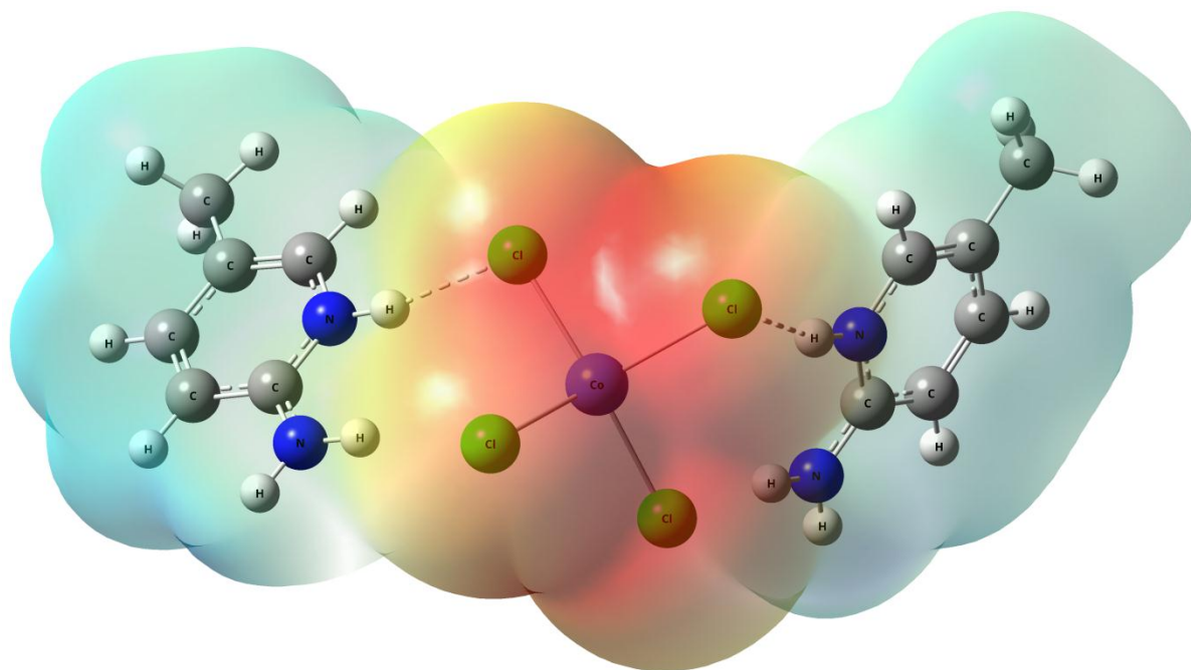


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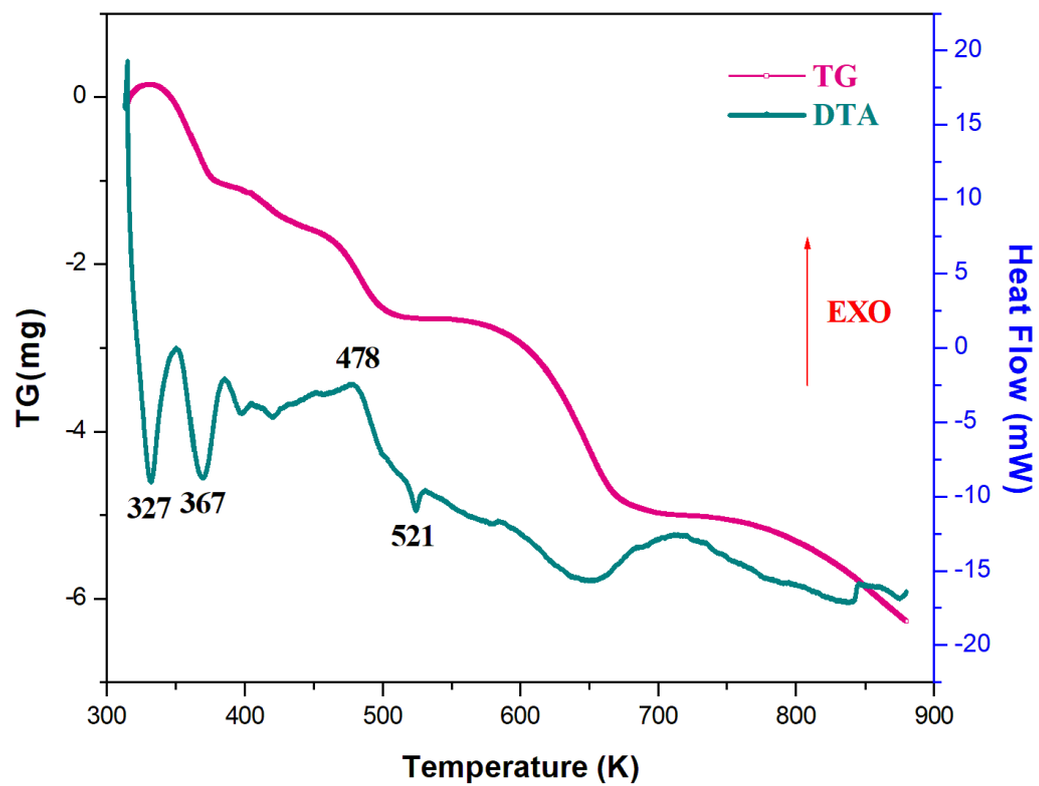


Fig. 10. DTA and TG curves of 2A5PCoCl₄.

Table 1

Crystal data and experimental parameters used for the intensity data collection strategy and final results of the structure determination.

Temperature	150K
Empirical formula	(C ₆ H ₉ N ₂) ₂ [CoCl ₄]
Formula weight (g mol ⁻¹)	184.2
Crystal system	Monoclinic
Space group	P2 ₁ /c
A	8.0734 (6) Å
b	14.6297 (11) Å
c	15.6913 (11) Å
β	103.034 (4) °
Z	4
V	1805.6 (2) Å ³
F(000)	852
μ(Mo Kα)	0.71073 Å
Index ranges	-9 < h _{max} < 10, -14 < k _{max} < 18, -20 < l _{max} < 16
Reflections collected	15135
Independent reflections	4139
Reflections with I > 2σ(I)	3126
R _{int}	0.033
Absorption correction: <i>multi-scan</i>	T _{min} = 0.697, T _{max} = 0.806
Refined parameters	210
R[F ² > 2σ(F ²)]	0.033
wR(F ²)	0.076
Goodness of fit	1.06
Δρ _{max} = 0.05e Å ⁻³	Δρ _{min} = -0.41e Å ⁻³

Table 2

Hydrogen-bond geometry (Å, °)

D—H...A	D—H	H...A	D...A	D—H...A
N1—HN1...C11	0.89 (3)	2.34 (3)	3.214 (2)	170 (3)
N2—H2A...C14	0.84 (3)	2.68 (3)	3.369 (3)	140 (3)
N2—H2B...C13 ⁱ	0.84 (3)	2.50 (3)	3.308 (3)	163 (3)
N3—HN3...C13	0.86 (2)	2.73 (2)	3.478 (2)	146 (2)
N3—HN3...C12	0.86 (2)	2.75 (2)	3.374 (2)	130 (2)
N4—H4A...C13	0.79 (3)	2.46 (3)	3.230 (3)	164 (3)
N4—H4B...C11 ⁱⁱ	0.79 (3)	2.52 (3)	3.303 (3)	176 (3)
C3—H3...C12 ⁱⁱⁱ	0.95	2.75	3.688 (3)	170
C6—H6...C13 ^{iv}	0.95	2.86	3.716 (3)	151
C12—H17...C14 ^v	0.95	2.81	3.754 (3)	173

Symmetry codes: (i) $x-1, y, z$; (ii) $-x+1, y+1/2, -z+1/2$; (iii) $x, -y+1/2, z+1/2$; (iv) $-x, -y+1, -z+1$; (v) $-x+1, -y+1, -z$.