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A proton transfer compound template phenylethylamine: synthesis, a collective experimental and theoretical investigations

Sofian Gatfaoui ^(a), Nouredine Issaoui ^(b), Thierry Roisnel ^(c) and Houda Marouani ^(a)

^(a)Laboratory of Material Chemistry, Faculty of Sciences of Bizerte, University of Carthage, 7021 Zarzouna, Tunisia

^(b)Laboratoire de physique quantique et statistique, Faculté des Sciences de Monastir, Université de Monastir, Monastir 5079, Tunisia

^(c)Centre de Diffraction X, UMR 6226 CNRS, Unité Sciences Chimiques de Rennes, Université de Rennes I, 263 Avenue du Général Leclerc, 35042 Rennes, France

ABSTRACT

Synthesis, crystalline architecture, vibrational spectroscopy, thermal analysis, electrical behavior and detailed theoretical DFT studies has been reported for the new inorganic-organic hybrid material phenylethylammonium trioxonitrate with general chemical formula $(C_6H_5(CH_2)_2NH_3)NO_3$ (PEAN). The determination of crystal structures plays an significant to understanding weak, intra and intermolecular, interactions that great influence crystal packing. PEAN has been crystallized to the monoclinic system with space group $P2_1/c$ and the network parameters obtained are $a = 5.9497(8) \text{ \AA}$, $b = 20.494(3) \text{ \AA}$, $c = 8.4528(11) \text{ \AA}$, $\beta = 113.812(8)^\circ$, $V = 942.9(2) \text{ \AA}^3$ and $Z = 4$. The atomic arrangement of the PEAN compound can be described by 3D framework. The organic cations are linked to the nitrate anions by means of an assortment of $N-H \cdots O$ hydrogen bonding trapped in layers at $y = \frac{1}{4}$ and $\frac{3}{4}$. These layers are interrelated via $C-H \cdots \pi$ interactions involving organic cations, an infinite chain are consequently created along the c -axis. Hirshfeld surfaces and 2D fingerprint plots estimate the weak intermolecular interactions accountable for the generation of crystal packing. The structural propriety and vibrational spectral analyze of PEAN have been investigated by using DFT calculations at B3LYP/6-311++G(d,p) level of theory. Intermolecular interactions were studied by NBO and AIM analyses. Thermal analysis (TG-DTA and DSC) and dielectric conductivity was also investigated. These measurements were realized to discuss the observed phase transition mechanism in the vicinity of 340 K.

Keywords: Crystal structure, Hirshfeld surface analysis, AIM, NBO, dielectric conductivity.

1. Introduction

Organic-inorganic hybrid materials have been increasingly attracted to recent research, hence the synergy between the thermal, electrical and/or optical properties of inorganic entities and the physico-chemical properties of organic materials through different interactions, electrostatic, non-bonding and van der Waals interactions penetrate several application fields such as biology [1,2], medicine [3], photo catalysis [4], also these hybrid materials flood the fields of optics and optoelectronics [5]. Phenylethylamine (PEA) is a monoaminated alkaloid that occurs in the nervous system of humans and other mammals, functioning as a transmitting neurohormone or a neurohormone moderator that maintains energy, attention and humor. This monoamine classified as a natural and physiological treatment against depression, for this reason studies and observations clinical have been made and show that (PEA) (at the dose of 10 mg once or twice a day) is effective in 60% of the patients. Its encounter with conventional antidepressant drugs has no adverse consequences like weight gain and sexual inhibition. (PEA) is structurally similar to amphetamine, it produces pharmacological effects and widely used in optical resolutions. (PEA) levels are shown to be very low in the biological fluids of depressed patients. Biochemically, PEA plays the role of an excitatory neurotransmitter and organizes neuronal potentials to promote glutamate activity and trigger neurotransmitters [6-8]. As part of our current study, we report our investigations on a new organic-inorganic hybrid compound: Phenylethylammonium nitrate with general chemical formula $(C_6H_5(CH_2)_2NH_3)NO_3$ (PEAN). Within this context chemical preparation, structural study and geometric optimization associate with an insight into Hirshfeld surface analysis are discussed. In addition, theoretical analyzes have been carried out along this work, which considerably enrich the available knowledge about the material, namely the vibrational properties, as well as the topological properties AIM [9] and orbital NBO [10]. The dielectric and thermal analysis study are also discussed.

2. Experimental

2.1. *Materials and physical measurements*

Measurements of X-ray diffracted intensities were recorded at 150 K using a diffractometer Bruker-AXS APEXII, working at molibdene wavelength ($\lambda = 0.71073 \text{ \AA}$). Absorption corrections were carried out by the multi-scan technique by the means of SADABS program [11]. Among the 6632 reflections measured only 2159 are independent, the remainder (1539) had an intensity $I > 2\sigma(I)$. The direct methods using the *SIR97* program were used to solve the structure [12]. The structure was, and then, refined with full-matrix least-square methods based on F^2 (*SHELXL-97*) [13] through the assist of the *WINGX*

program [14]. All non-hydrogen atoms were refined by means of anisotropic atomic displacement parameters. All the hydrogen atoms linked to carbon atoms were located in calculated positions and refined by the way of fixed individual displacement parameters [$U_{iso}(H) = 1.2U_{eq}$] awarded to the riding model (C–H (aromatic, methylene) bond lengths of 0.93 Å and 0.97 Å, respectively). The remained hydrogen atoms were located in a difference map.

A final refinement on F^2 converged at $R(F^2) = 0.049$ and $wR(F^2) = 0.125$. In **Table 1** we have recorded: the parameters used for the X-ray data, the strategy for the crystal structure determination and its final results. Fig. 1a illustrate an ORTEP [14] drawing of the molecular structure. Infrared (IR) spectrum, in the 4000-400 cm^{-1} region, was collected at room temperature by the means on a Nicolet IR 200 FTIR spectrophotometer. Thermal analysis, from room temperature up to 850K, was realized using a multimodule 92 Setaram analyzer. This later, operate at an average heating rate of 5K/min. In this study, the mass of the sample used was 5.63 mg for DTA/TG and 6.8 mg for DSC. An open circuit using two electrode configurations in the band of frequency ranging from 5 Hz to 13MHz and with signal amplitude of 50 mV were taken in the measurements of the impedance. The pellet surfaces were covered with silver pastes electrodes whereas the platinum wires linked to the electrodes were used as current collectors.

2.2. Theoretical studies

To confirm the arrangement of the hydrogen bonds and the inter and intra-molecular interactions in our material, we studied it by Hirshfeld surfaces analysis, this is an excellent approach used to generate unit cell packing diagrams of the compounds. This analysis is based on a 3D graph that represents the space region where the molecules come in contact and another 2D that summarizes the complexes information contained in a structure and makes it possible to identify each type of interaction. The analysis and calculations of this empirical method were carried out using the Crystal Explorer version 3.1 [15] software imported on a CIF file.

The theoretical calculations have been performed by using the hybrid B3LYP/6-311++G(d,p) method implanted in the Gaussian 09 software package [16]. The GaussView 06 was used as molecular visualization program [17]. Due to the anharmonicity effect, the calculated vibrational frequencies were scaled by using a scaling factor. In our study a factor 0.958 in the range of wavenumbers above 1700 cm^{-1} were scaled by 0.958 whereas the range of wavenumbers below 1700 cm^{-1} were scaled by a factor equal to 0.983 [18, 19]. The assignments of the bands observed in both spectrum (IR and Raman) were performed with

the help of the corresponding TEDs using the VEDA4 package [20]. The GaussView package was used to verify the assignments of the bands. In this work, NBO calculations [21] were employed to calculate the donor-acceptor energy interactions, topological properties were determined with the AIM2000 program [22] in harmony with the Bader's theory [23].

2.3. Chemical preparation of $(C_8H_{12}N)NO_3$

$(C_8H_{12}N)NO_3$ was synthesized according to the following procedure. A solution containing 1 mmol of nitric acid (HNO_3) in 10 mL of distilled water was added to 1 mmol of phenylethylamine in 10 mL of ethanol. The mixture was stirred at room temperature for one hour and then left to evaporate. After a few days of evaporation, the solution reaches saturation and gives rise to colorless crystals having a prismatic shape, stable and of suitable size for a structural study. Schematically the equation of the reaction can be written as follow:



3. Results and discussion

3.1. X-ray diffraction and geometric optimization

The unit formula of the title material (PEAN) (Fig. 1a) demonstrates a monoprotinated organic cation which is an inorganic anion (NO_3^-) and 2-phenethylammonium. The optimized geometry model of this compound is illustrated in Fig. 1b. To describe the atomic arrangement of our crystal structure, we have projected it along the \vec{a} axis (Fig. 2a). This projection shows the arrangement of NO_3^- anions in a corrugated manner in the $y = 1/4$ and $3/4$ positions between which the 2-phenethylammonium groups are interleaved, which are interconnected by C–H ... π interactions [24, 25], thus forming a cationic chains along the axis \vec{c} to $y = 0$ and $1/2$. These anionic and cationic groups are interconnected via weak hydrogen bonds N–H ... O. The hydrogen bonding strength can be interpreted according to Brown and Blessing criteria [26, 27] based on the donor-acceptor distance ($d > 2.73 \text{ \AA}$: weak; $d < 2.73 \text{ \AA}$: strong). Thus, in PEAN, there are three N–H ... O hydrogen bonds, two to three centers (bifurcated) generating cycles $R_1^2(4)$ [28] and one at two centers (Table. 2). These bonds are considered to be weak according to the donor-acceptor distances which vary between 2.815 (2) and 3.254 (2) \AA . Each organic moiety is bonded to three different nitrate anions through five hydrogen bonds, thus forming $R_6^4(14)$ and $R_1^2(4)$ units [28] (Fig. 2b). The interatomic distances and the bonding angles of (NO_3^-) collected in Table 3 do not show any particularity compared to those generally encountered in nitrates known until now [29, 30]. Concerning the organic cation, the geometric characteristics classified in Table 3 are comparable to those observed for the same cation crystallizing with other types of anions [31]. The C– NH_3^+ bond

length (1.48 (2) Å) of the ammonium group is considerably higher than the C–NH₂ bond of the amine, this being explained by the protonation of 2-phenylethylamine. The benzene ring of the 2-phenethylammonium cation is perfectly plane, the mean deviation from its mean plane is ± 0.0019 Å. Packing of organic cations in the crystalline stack is ensured *via* C–H... π interactions (Fig. 2c) with donor-acceptor distances (C...Cg) ranging from 3.554 (2) to 3.704 (2) Å (Table 2).

The geometric parameters studied experimentally by X-ray diffraction, and theoretically by BLYP methods with 311 ++ G (d, p) and 311 ++ G (3df, 3pd) basis, such as the bond lengths and the angles of the PEAN are tabulated in **Table 3**. Optimized molecular structure has a dipolar moment equal to 6.0203 Debye and energy worth -647.33030001 a.u. Protonation on the nitrogen atoms N2 of the cation and the loss of a proton of nitric acid are confirmed by the extension of C1-N2, O2-N1 and O1-N1 bonds respectively. These differences lead to the conversion of neutral state to ionic state. The C-C linkages in the phenyl of the 2-phenylethylammonium group were calculated at the range of 1.3899 Å-1.3993 Å, while the C-C-C binding angles were calculated between 118.3939 ° and 120.94 °. These values do not differ greatly from those observed by X-ray diffraction, which allows us to conclude that the aromatic nucleus is somewhat irregular. The average value of the calculated bond length of C–H and N–H is equal to 1.11 and elongated by about 0.1 Å relative to the mean value observed by experimentally of 1.01. This later is due to the difference between the theoretical calculations which done to an isolated molecule in gaseous phase and experimental results which molecules is in solid state, where the crystal structure is related to intra and intermolecular interactions, such as van der Waals and hydrogen bonding interactions. The synergy between the organic group and the nitrate anion is carried out through hydrogen atom H2N which provide an intermolecular interaction between the two entities playing a very important role in the maintenance and the cohesion of the crystalline edifice. The angle of the intermolecular is correlated to be linear in both experimental and computational methods. Also the remarkable variation of O3 ... H2N distance experimentally compared with the results observed theoretically. This one suggests the importance and the strong character of the hydrogen bond N2-H2N ... O3.

3.2. 3D cartographies and 2D fingerprint plots

The analysis of the three-dimensional graphic representation in d_{norm} illustrated in (**Fig. 3a**) makes it possible to identify the contacts which take place in our structure, namely the close contacts indicated by the red spots taking place between the hydrogen and oxygen atoms (H ... O/O ... H) which are attributed to hydrogen bonds N–H ... O. On the other hand, the large red

spot on the aromatic nucleus and the blue spot surrounding the H1A and H2B hydrogen observed on the Shape index cartography (**Fig. 3b**), the large flat region delimited by a blue outline observed on the Curvedness graph (**Fig. 3c**), and the important contribution of the contacts C ... H/H ... C (17.3%) on the Hirshfeld surface (**Fig. 4d**) are compatible with the presence of C–H ... π interactions in our structure. Noting the absence of the red and blue triangles on the Shape index cartography, the low contribution of the contacts C ... C (0.8%) (**Fig. 4e**) thus the sum ($d_i + d_e \sim 3.6\text{\AA}$) greater than the sum of the van der Waals radii of the carbon atoms (3.4\AA) confirm the absence of π - π interactions in the crystalline stack. The full fingerprint plot and the fingerprints plots of the main contacts involved in the PEAN 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. There are close contacts when the sum of these components is smaller or equal to the sum of the van der Waals rays of the involved atoms. **Fig. 4b** is a simplified illustration 2D that takes into account only the contacts between hydrogen and oxygen atoms, this graph reveal that these contacts occupy the largest space on the Hirshfeld surface (38.8%) and appear as two narrow symmetric points located around a sum ($d_e + d_i = 1.9\text{\AA}$) less than the sum of the van der Waals radii of the oxygen (1.52\AA) and hydrogen (1.09\AA) atoms. These contacts clearly demonstrate the formation of hydrogen bonds N-H ... O. The graph depicted in **Fig. 4c** represents the relationship between the hydrogen atoms (H...H) which comprise 36.9% of the total surface area of Hirshfeld. It is observed for its contacts that a single quantity of pairs (d_e, d_i) of high density located in the zone centered on the point ($d_i = 1.1\text{\AA}$, $d_e = 1.1\text{\AA}$) use of color green. This value is almost equal to the van der Waals radii value of the hydrogen atom (1.09\AA).

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

R. Bader proposed an AIM topological approach [23, 32] which considers the electronic density $\rho(\mathbf{r})$ as a local function of the molecular system. This approach is a model which characterizes the chemical bonding of a system based on the topology of the charge density in the space $\rho(\mathbf{r})$. The AIM quantum theory allows characterizing the nature of interatomic interaction between two atoms involved, the non-binding interactions, the electronic structure and the chemical reactivity of molecule, from different topological properties (ρ , λ_i (1, 2, 3), $\nabla^2 \rho(\mathbf{r})$), energy (the kinetic energy densities G, potential V and total $H = G + V$) and the electronic distribution. Recall that the Laplacian of electronic density, defined as the sum of the three eigenvalues of Hessian matrix, given a valuable information on the charge

concentration ($\nabla^2\rho > 0$: depletion of load, $\nabla^2\rho < 0$: load concentration areas). The ellipticity $\varepsilon = \left| \frac{\lambda_1}{\lambda_2} \right| - 1$, given also an idea of the accumulation (depletion) of charges in a given plane.

According to AIM theory, a bond is characterized by the existence of a BCP (critical point of binding). In this context, the properties of BCP are analyzed according to the parameters described above. To calculate the properties of the bonds in our compound, in particular the hydrogen bonds Koch and Popelier [33] proposed three criteria to show the existence of a hydrogen bond:

- The presence of a BCP between the acceptor group and the hydrogen atom of the donor group confirms the existence of a hydrogen bond.
- The value of $\rho(\text{BCP})$ at the BCP point should be low and located in the range 0.0020–0.0400 a.u.
- The value of $\nabla^2\rho(\text{BCP})$ should be positive and located in the domain 0.0240–0.1390 a.u.

Based on the Rozas and al [34] criterion we have classified into three categories the hydrogen bonds:

- Weak hydrogen Bonds: $\nabla^2\rho(r_{\text{BCP}}) > 0$ and $H(r_{\text{BCP}}) > 0$
- Moderate hydrogen bonds: $\nabla^2\rho(r_{\text{BCP}}) > 0$ and $H(r_{\text{BCP}}) < 0$
- Strong or very strong hydrogen bonds: $\nabla^2\rho(r_{\text{BCP}}) < 0$ and $H(r_{\text{BCP}}) < 0$

The graphical representation of AIM analysis of the PEAN compound is illustrate in **Fig. 5**. whereas the calculated topological parameters are listed in **Table 4**.

The AIM topological analysis reveals that our crystal is stabilized by three hydrogen bonds O1 ... H1, O1 ... H2 and O2 ... H. According to the BCP analysis of Table 4, O1 ... H1 and O1 ... H2 are considered weak with hydrogen bond energies of -1.18171 and -1.03322 kcal / mol, the strength of these two hydrogen bonds is also confirmed by the Laplacian of electron density (0.02270, 0.01860) and the energy density (0.60600, 0.43177) with values greater than zero follow the Rozas criterion. Concerning the bond O2 ... H, this bond has an energy $E_{\text{int}} = -8.97874$ kcal / mol, a Laplacian of the electronic density $\nabla^2\rho(r) = 0.08946 > 0$ and an electronic density $G(r) + V(r) = -1.93379 < 0$, indicating that O2 ... H is a mean hydrogen bond. Positive Laplacian values indicate electronic charge depletion along the path of its links. 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 nitrate anion via hydrogen bonds give rise to the formation of the rings NRCP1

and NRCP2. The very low value of ellipticity ($\epsilon = -1.2073$) at the RCP point confirms that there is delocalization of electron in aromatic nucleus. Indeed the overall ellipticity suggests that there is a strong delocalization in PEAN crystalline structure [$\epsilon(\text{NRCP1}) = -1.8530$, $\epsilon(\text{NRCP2}) = -1.7866$]. Electron localization function (ELF), local information entropy, reduced density gradient (RDG), localized orbital locator (LOL), average local ionization energy, ellipticity (ϵ) and the eta index (ζ) of the hydrogen bonds in the PEAN material at the critical points BCP are detailed in **Table 4**.

The existing intermolecular interactions in PEAN 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. [35]

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

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 performed and plotted respectively by the use of Multiwfn and VMD programs [36, 37]. These results were collected in **Figs. 6a** and **6b**. **Fig. 6a** illustrates the attractive, van der Waals and repulsive interactions in the PEAN structure. These interactions appear as zones marked respectively by blue, green and red circles. Turning to **Fig. 6b**, a clear blue spot is checked between the hydrogen and oxygen atoms, which signify the strong attractive interaction N-H ... O. The interactions appear in the form of green plates are attributed to van der Waals interactions; its plates are located exactly between the hydrogen atoms. The elliptic red plate located at the center of the aromatic nucleus is related to repulsive interactions which show a strong steric effect.

3.4 Natural Bond order analysis

Natural bond analysis (NBO) is an 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 [38, 39]. 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 $E^{(2)}$ [40]:

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

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.

The computed results are collated in **Table 5**. The examination of this table shows clearly the existence of a hydrogen bond N-H ... O. This is explained by the important interactions between the free pair of LP(O3) and LP(O2) orbitals and the antibonding orbital $\pi^*(O1-N1)$ with stabilizing energies respectively equal to 47.66 and 55.1 kcal.mol⁻¹. The total stabilization energy of LP(O3) \rightarrow $\sigma^*(N2-H2N)$ interaction is 14.04 kcal.mol⁻¹ order. Interactions between the $\pi(C3-C4)$, $\pi(C3-C4)$, $\pi(C7-C8)$, $\pi(C7-C8)$, $\pi(C5-C6)$ and $\pi(C5-C6)$ bonding orbitals and antibonding orbitals $\pi^*(C7-C8)$, $\pi^*(C5-C6)$, $\pi^*(C3-C4)$, $\pi^*(C5-C6)$, $\pi^*(C3-C4)$ and $\pi^*(C7-C8)$ lead to an average stabilization energy around 20.5 kcal.mol⁻¹. It is obvious that its interactions significantly affect the stabilization and consolidation of the crystalline architecture of PEAN compound.

3.5. Molecular electrical potential surface and electron localization function

The electrostatic potential surface is used to expect the behavior and the chemical reactivity of molecules. This property is a valuable tool for understanding molecular interactions. Its importance lies in the fact that it displays several molecular properties namely the regions of positive electrostatic potentials, negative and neutral in terms of a color code. The blue color indicates the positive values (region of strong attraction), the red indicates the negative values (region of strong repulsion) and the green indicates the neutral zone. The map of electrostatic potential surfaces of the PEAN compound is illustrate in **Fig. 7a**. These areas are plotted in the region between $-3.8e-2$ (red) and $3.8e-2$ au (blue). The blue color reveals the

strongest attraction whereas the red color indicates the strongest repulsion. The area of negative potentials (excess in electrons) are associated with the free doublets of oxygen atoms acceptor of H^+ proton of NO_3^- group while the regions of the positive potentials (deficient in electrons) are associated with the $-NH_3^+$ group of the phenylethylammonium cation. This confirms the existence of intermolecular $N-H \cdots O$ interactions. The electrophilic and nucleophilic sites explain the formation of the H-bonds between the nitrate and phenylethylammonium groups in the compound under study and its important role in its stabilization. The surface of the electrostatic potential is very useful in the search for a molecular structure with its physicochemical properties. A new map based on the critical points located inside this surface has been used in the **Fig. 7b**. Using the Monte Carlo method to calculate the values of the red and green spheres correspond respectively to the maximums and the minimums inside the surface, these data are collated in **Table 6**. The maximum energies located on this surface at the 6th and 7th green spheres as well as the minimum energies for the 3th and 4th blue spheres are derived from the interactions between the phenylethylammonium group and nitrate anion via $N-H \cdots O$ hydrogen bonds.

In quantum chemistry, the ELF is a measure of the possibility of discovering an electron in the locality of a reference electron density at a given point and with the same spin. The bifurcation diagram of the ELF localization domains generally gives information on the nature of the interactions by observing the order of domain separation (reduction process) when the value of ELF isosurface varies between 0 and 1. The ELF value close to 1 is relative to the maximum Pauli repulsion regions, while the minimum Pauli repulsion regions have the ELF value near 0. **Fig. 8** clearly shows the topology of ELF function for the PEAN crystal. The red colored field located in the range (0.8 - 1.0) is assigned to the high ELF values. The zone between 0.8 and 0.5 (yellow to green) corresponds to the average ELF values. Finally the series goes down from 0.5 to 0 represented by the blue color shows low ELF values. In conclusion we find that the red regions surrounded by hydrogen atoms with a single electron are the strongest with a maximum Pauli repulsion, while the other blue areas occupied by the nitrogen atom of nitrate group and the C3 and C6 atoms of organic cation having similar spin electrons close with a minimum Pauli repulsion.

3.6. Vibrational IR spectra and assignments

Work perform on nitrates [41, 42] allowed us on the one hand to distinguish the four domains of modes relating to valence vibrations and their separation of deformation modes on the other hand. The plane trigonal NO_3^- anion contains 4 atoms so it presents $(4 \times 3) - 6 = 6$

vibrations modes. These six modes are decomposed according to a symmetrical elongation mode (ν_1, A_1 , Raman), a symmetrical deformation mode (ν_2, A_2 , IR), a doubly degenerated asymmetric elongation mode (ν_3, E' , IR et Raman) and a doubly degenerate asymmetric deformation mode (ν_4, E' , IR et Raman). The application of the correlation method between the free ion groups (D_{3h}), the site group (C_1) and the factor group ($C_{5_{2h}}$) of (PEAN) crystal, generates 24 internal vibration modes divided into two groups according to their activities, 12 modes active in Raman and 12 active in IR. The calculated and the experimental IR spectra of PEAN compound were found in the region 450-4000 cm^{-1} are presented graphically and listed respectively in **Fig. 9** and **Table 7**.

Vibrational modes of the anion NO_3^-

Based on the work reported above we have estimated the vibration domain of the nitrate group which is located between 700 and 1400 cm^{-1} . The intense bands around 1392, 1349 and 1311 are attributed to the asymmetry stretching modes ν_3 (NO_3^-), While the corresponding computed values are 1699, 1456 and 1269 cm^{-1} . Another stretching vibration of NO_3^- but symmetrical is provided at 1156/1031 and 1266/924 cm^{-1} respectively experimentally and by the DFT method. The band which is located experimentally towards 820 cm^{-1} is related to the outside plane deformation mode of NO_3 group, this mode it is theoretically calculated at 772 cm^{-1} . Finally the absorptions at 696 and 603 cm^{-1} are related to the deformation modes in the plane; these values are in good agreement with the calculated value (667 and 640 cm^{-1}).

Vibration modes of $-\text{NH}_3^+$

The $-\text{NH}_3^+$ groups of phenylethylammonium cation react with the nitrate anions via the N-H ... O type hydrogen bonds. The frequency of the band resulting from this type of coupling is observed experimentally at 2509 cm^{-1} and its theoretical peak located around 2428 cm^{-1} . Generally NH_3^+ stretching vibrations occur in the spectral range [3500-3000 cm^{-1}] [43]. This is already confirmed by the experiment and by the DFT method where the bands correspond to asymmetric stretching vibrations $-\text{NH}_3^+$ located respectively roughly 3447 cm^{-1} (experimentally) and 3492 cm^{-1} (theoretically). While the other symmetrical ones appear experimentally at 3385 cm^{-1} and theoretically to 3421 cm^{-1} . Concerning the bending modes, the $\delta_{\text{as}}(\text{NH}_3)$ bands are observed around 1640 and 1553 cm^{-1} , these modes are calculated at 1577 cm^{-1} . While the $\delta_{\text{s}}(\text{NH}_3)$ are located experimentally towards 1385, 1348, 1311, 1156 cm^{-1} and the calculated at 1357, 1297 and 1114 cm^{-1} .

Vibration modes of $-\text{CH}_2$ group

The stretching modes of the $-\text{CH}_2$ group (asymmetric and symmetric) generally appear at about 3100 and 3000 cm^{-1} [44]. The band observed at 3043 cm^{-1} in the infrared spectrum is attached to the symmetric stretching of methylene group ($-\text{CH}_2$), the matching theoretical mode appears at 3034 cm^{-1} . The asymmetric stretching modes of this group are computed at 3014, 2971 and 2963 cm^{-1} while the experimental modes are located at 3006 and 2963 cm^{-1} . The deformation modes of the $-\text{CH}_2$ groups are observed experimentally in the frequency region between 1553 and 1037 cm^{-1} , and their calculated by the DFT method appears in the spectral domain range between 1575 and 1007 cm^{-1} .

Ring vibration

The multiple weak bands in the frequency region 3100-3000 cm^{-1} , in the aromatic compounds, are principal due to the C-H stretching vibration in the aromatic cycle [45]. Absorptions observed at 3093, 3049 and 3018 cm^{-1} are attributed to C-H stretching vibrations (computed at 3141, 3129 and 3120 cm^{-1}). Focusing on the C-H in-plane bending vibrations, the frequencies of these vibrations are appear as medium strong to strong bands experimentally in the spectral range 1553-1037 cm^{-1} . In theoretical FT IR spectrum these vibrations are limited between 1575 and 1007 cm^{-1} . The stretching modes $\nu(\text{C}-\text{C})$ of the phenyl group are observed respectively around 1640, 1547, 1392, 1348, 1311 cm^{-1} . These vibrations are calculated at 1575, 1553, 1424, 1284 and 1173 cm^{-1} . The aromatic carbon-carbon ($\text{C}=\text{C}$) deformation vibration in the plane occurs experimentally in the region 1000-490 cm^{-1} and calculated in the frequency region between 1007-492 cm^{-1} . The peak around 485 cm^{-1} , affected to the deformations out of the plane of $\text{C}=\text{C}$ groups. The same vibrations are calculated at about 492 cm^{-1} .

3.7. Thermal behavior

To understand the behavior of the tile compound, thermal analysis DTA/TG (**Fig.10a**) of our compound carried out on a mass sample equal to 5.63 mg in air with a heating rate of 5 $\text{K}\cdot\text{min}^{-1}$ in an alumina crucible from 0 K up to 850 K shows the existence of three endothermic peaks located respectively at 341, 368 and 408 K on the DTA curve without a weight loss on the TG curve correspond to three successive phase transitions and an exothermic peak at 498 K accompanied by a significant weight loss (82.50%), during which our product undergoes total combustion which leads to volatile gases and a black carbonaceous residue.

The calorimetric thermal analysis (**Fig.10b**) carried out on this compound ($m = 6.8$ mg) in the temperature range between 298 and 573 K under an argon atmosphere and with a heating rate equal to 5 $\text{K}\cdot\text{min}^{-1}$ represents the same peaks corresponding to same phenomena

previously observed by DTA. In the subject to study the properties of the phase transitions which appear at about 341, 368 and 408 K, 9.4 mg of product are placed in an aluminum nacelle heated from 280 K to 428 K and then cooled to ambient temperature. The DSC curve (**Fig. 11**) shows during heating three endothermic peaks at 340, 373 and 423 K and three exothermic peaks during cooling at 318, 370 and 417 K. These peaks suggest the detection of three successive reversible phase transitions.

3.8. Electrical transport properties

Research in the field of ionic conductivity and understanding the mechanism of short-range interactions in materials has evolved very rapidly thanks to the explorer of many materials with a very high ionic conductivity, which is very important both in electronics field and in the energy field. From one material to another, the conductivity differs as well by the nature of responsible species for the conduction (Li^+ , Na^+ , Ag^+ , H^+ ...) than by the temperature range of use and the dimensionality of conductive phase (3D, tunnel, lamellar...). In order to discover a possible electrical conduction in the synthesized organic nitrate (PEAN) we have undertaken a dielectric studies to better understand the properties of electric transport in this crystal as a temperature function and in a wide range of frequencies (0.01-13000 kHz). The sample is presented in the pellet form of a 13 mm diameter obtained from polycrystalline powder compressed at a pressure of about 7 tons in a cylindrical stainless steel matrix attacking a thickness of 1.25 mm. In the subject to find a uniform electric contact, a Ag lacquer is deposited on both sides of the pellet.

Impedance spectroscopy

We have chosen to represent the electrical response of the (PEAN) material by Nyquist diagram (**Fig.12a**). The representative points of the complex impedance Z are arranged in an arc of approximately circular shape at the highest frequencies. The most important parameter for the subsequent study is the R resistance, whose value makes it possible to calculate the conductivity of (PEAN) material. To better describe the experimental Nyquist diagram, it therefore seems logical to model the electrical behavior by the series association of R and CPE cells, corresponding to the contributions of grains in the materials. In the circuit equivalent: R and CPE are the resistance and the constant phase element, respectively, used to reimburse the non-homogeneity of the system. These spectra (**Fig.12a**), passing close to the origin, are constituted by semicircles centered below the real axis suggesting the leaving from ideal Debye behavior [46]. We represent at diverse temperatures, as a function of the frequency, the variation of the real part Z' and imaginary part Z'' of the impedance. Indeed the **Fig.12b** ($Z' = f(\log f)$) shows that Z' decreases during the temperature and frequency

increasing, while we note in **Fig.12c**, which represents $Z'' = f(\log f)$ the appearance of peaks ($\omega_{\max}, Z''_{\max}$) whose increase in temperature causes a reduction in intensity. It is observed that everyone of these peak is centered at a maximum frequency (f_{\max}) and when the temperature increase these peak shifts towards the high frequencies. This explains the dependence of the relaxation phenomenon with the temperature [47, 48] and the position of these peaks follows the condition $\omega_m \cdot \tau_m = 1$ [47-49].

Electrical conductivity

The knowledge of geometry of the sample (surface [S] and thickness [e]) makes it possible to calculate the conductivity σ of the sample at a given temperature according to the equation:

$$\sigma = \frac{e}{SR}$$

Conductivity measurements being performed at different temperatures, it is then possible to plot the $\log(\sigma \cdot T) = f(1000 / T)$ curve for our sample (PEAN) (**Fig.12d**). These measurements revealed a slope break at $T = 378$ K, hence our representative curve consists of two regions, each region in the form of an affine branch that obeys the Arrhenius law:

$$\sigma T = A \exp\left(-\frac{E_a}{KT}\right)$$

In the last equation A , K and E_a represent, respectively, the pre-exponential factor, the Boltzman constant and the activation energy.

Beyond the temperature 378 K the activation energy increases considerably and goes from $E_{a1} = 0.08$ eV to $E_{a2} = 0.2$ eV. In the vicinity of this temperature a phase transition has been detected, this thermal phenomenon is already confirmed by DTA (341K), TG and DSC up (340K) and down (318K) temperature. The abrupt change in conductivity marks the transition to a more chaotic state that could be correlated with the changes in the orientation of phenylethylammonium or NO_3^- ions and the high dynamic disorder of the $-\text{NH}_3^+$ groups in the hydrogen bonds $\text{N-H} \cdots \text{O}$. For $E_{a2} = 0.20$ eV, the conductivity increases to a value equal to $1.21 \cdot 10^{-4} \Omega^{-1}\text{cm}^{-1}$, this important variation probably suggests the high mobility of H^+ protons under the effect of thermal agitation, also the appearance of new charge dipole-dipole type carriers which improves the conductivity within our material.

Correlation between electrical and structural properties

The augmentation of the temperature has a very interesting effect on the architecture and the network of our crystal building, it favors the vibration of layers formed by the organic groups and the nitrate anions can lead to the rapid mobility of H^+ proton in the hydrogen bond network and to the reorientation of the ions NO_3^- . In addition, it's worth noting that the X-ray study shows that the H-bonding system in the PEAN compound is three-dimensional taking

into account the interactions of type C-H ... π which own a strong conductivity wherever the motion of the protons H^+ takes place in the 3D directions [50]. Besides, the conductivity in the studied structure is superior to those in the lamellar structures where the movement of H^+ happens in only 2D [51].

4. Conclusions

Single crystals of a novel proton transfer compound phenylethylammonium trioxonitrate (PEAN) was crystallized at 300K by slow evaporation-solution growth technique and investigated by X-ray diffraction, thermal analyses, IR spectroscopy and ionic conductivity in the solid state phase. XRD reveals the arrangement of NO_3^- anions in a corrugated behavior in the $y = 1/4$ and $3/4$ positions between which the 2-phenylethylammonium groups are interleaved, which are interconnected by C-H ... π interactions. These latter interactions allow the reinforcement of the 3D network. The Hirshfeld surface study was used to evaluate the intermolecular contacts in the crystal arrangement namely H...H, H...O/O...H, C...H/H...C, C...C. The experimental vibrational frequencies of the fundamental modes of PEAN have been properly analyzed, assigned, and compared the theoretical results. NBO, AIM analysis has been used to study the stability of our compound. The thermal analyses show three successive phase transitions, one of them is confirmed by ionic conductivity. In addition a correlation between structural and electrical properties is also occurs.

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Figure Captions

Fig.1: ORTEP drawing of (PEAN) with the atom-labeling scheme. Displacement ellipsoids are drawn at the 30 % probability level. H atoms are represented as small spheres of arbitrary radii (a) and the optimized molecular structure (b).

Fig.2 : Projection along the \vec{a} axis of atomic arrangement of PEAN (a). Hydrogen bond motifs in (PEAN) (b). Interactions between organic cations C-H... π stacking (c).

Fig.3 : Hirshfeld surfaces mapped with d_{norm} (a) (dotted lines “red” represent hydrogen bonds), *shape index* (b) and *curvedness* (c) for the PEAN.

Fig.4 : Fingerprint plots of the full and the import intercontacts and percentage of various intermolecular contacts contributed to the Hirshfeld surface in the PEAN compound.

Fig.5 : Graphical representation of the AIM analysis of PEAN: bond critical points (small red spheres), ring critical points (small yellow sphere), bond paths (pink lines).

Fig.6 : Reduced density gradient (a) and isosurface density (b) plot along with the color filled scale bar defining interaction limits for the PEAN compound.

Fig.7 : Molecular electrostatic potential MEP (a), Surfaces local minima and maxima of MEP are represented red and blue spheres of (PEAN).

Fig.8 : The map for electron density in the XY plane of PEAN. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

Fig.9 : Theoretical and experimental FT-IR spectrum of PEAN.

Fig.10 : DTA, TG (a) and DSC (b) curves c of (PEAN) at rising temperature.

Fig.11 : DSC curves rising and falling temperature of (PEAN).

Fig.12 : Cole-Cole plots (a), plots of the real and imaginary parts of impedance Z' (b) and Z'' (c) vs $\log(f)$, Arrhenius plots ionic conductivity of $\ln(\sigma.T)$ versus $1000/T$ of (PEAN) at various temperatures.

Tables captions

Table 1 : Crystal data and experimental parameters used for the intensity data collection strategy and final results of the structure determination.

Table 2 : Hydrogen-bond geometry (Å, °) of PEAN.

Table 3 : Bond length (Å), bond angles (°) and torsional angles (°) by X-ray data (with estimated standard deviation in parentheses) and by theoretical calculations at different levels of theory.

Table 4 : Topological parameters for Hydrogen bond interactions.

Table 5: Second order perturbation theory analysis of Fock matrix on NBO basis for the (PEAN) using B3LYP/ 6-311++G(d,p).

Table 6 : Analysis of molecular surface based on blue and red point numbers (Fig. 7).

Table 7: Observed and calculated wavenumbers (cm⁻¹) and assignments for (PEAN) compound.

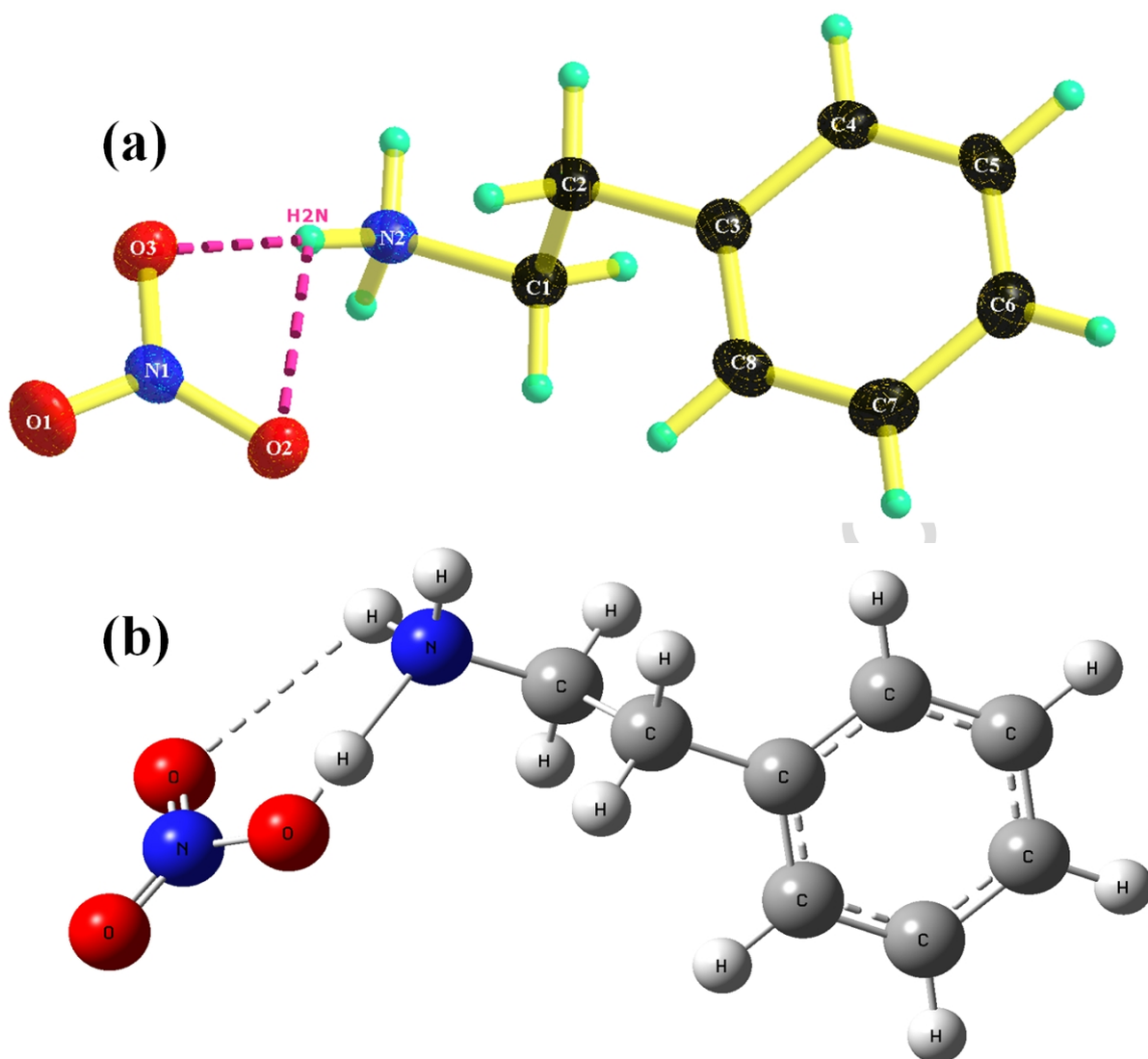


Fig. 1

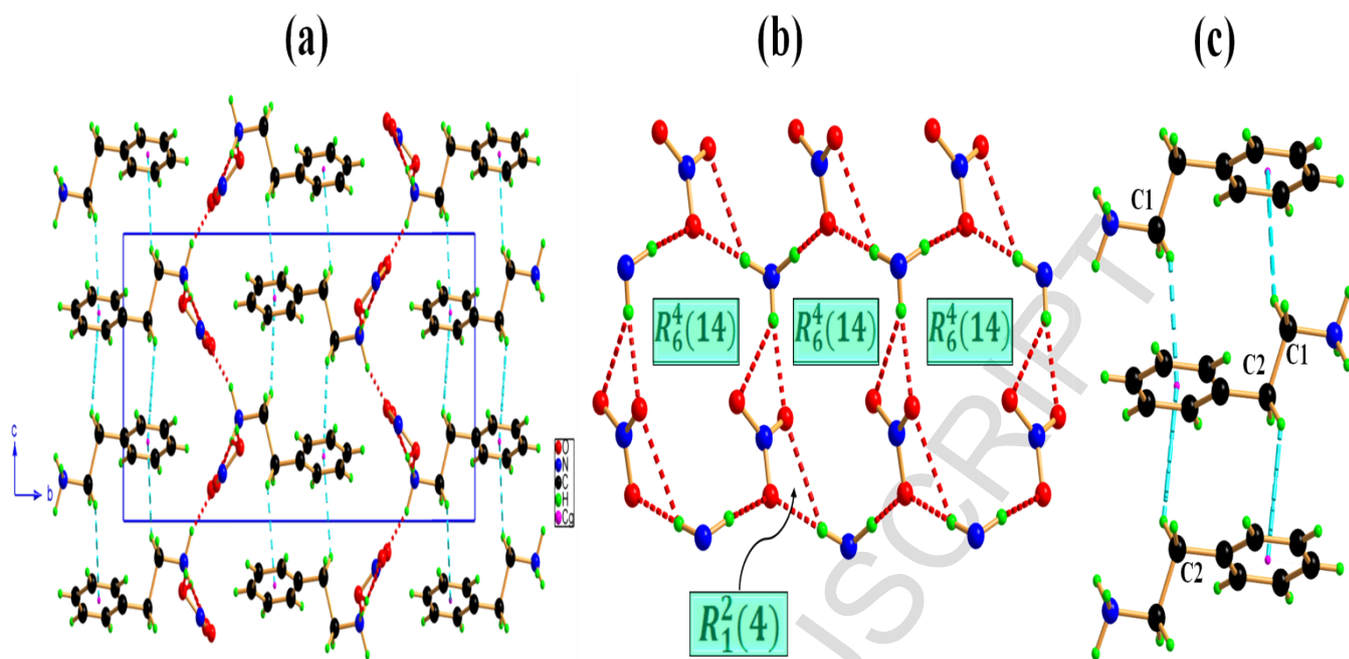


Fig. 2

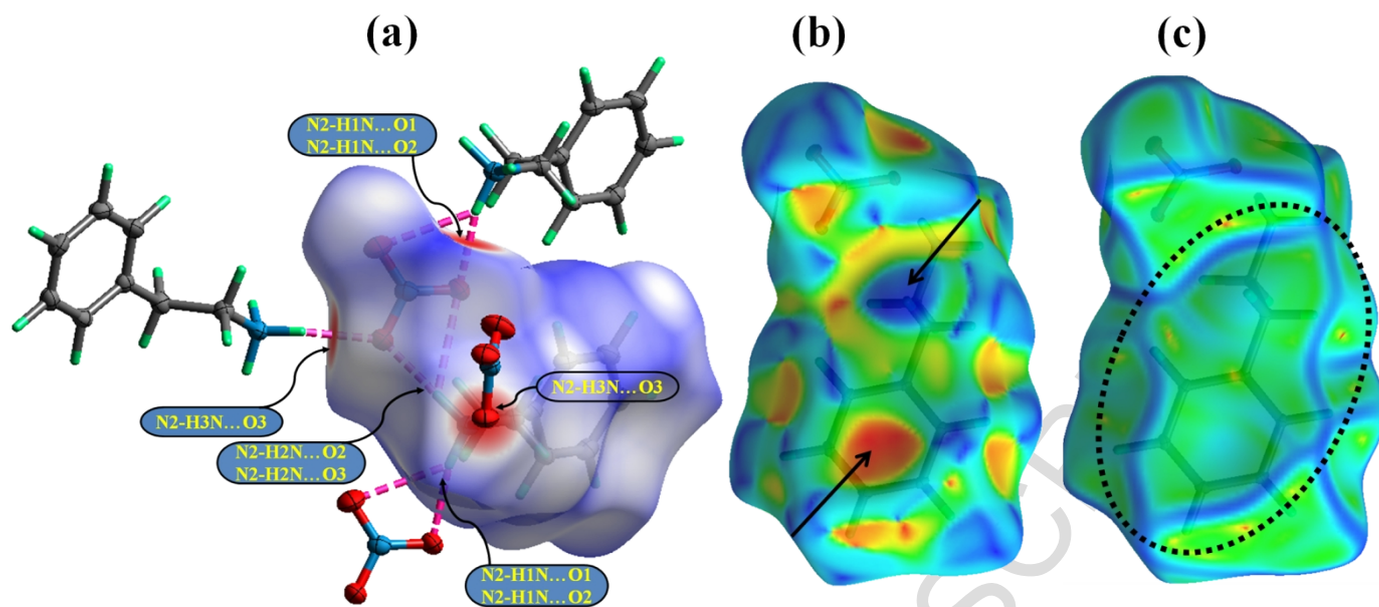


Fig. 3

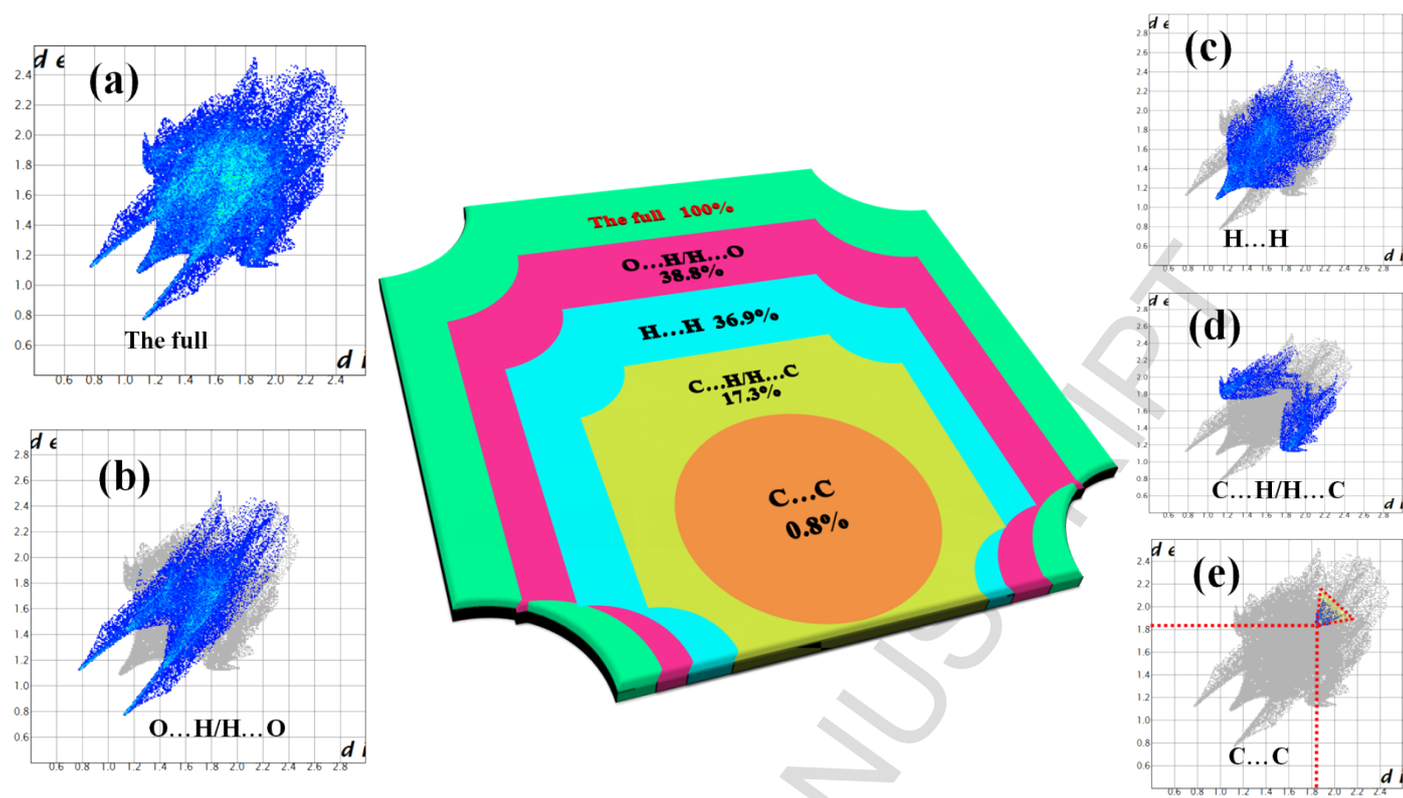


Fig. 4

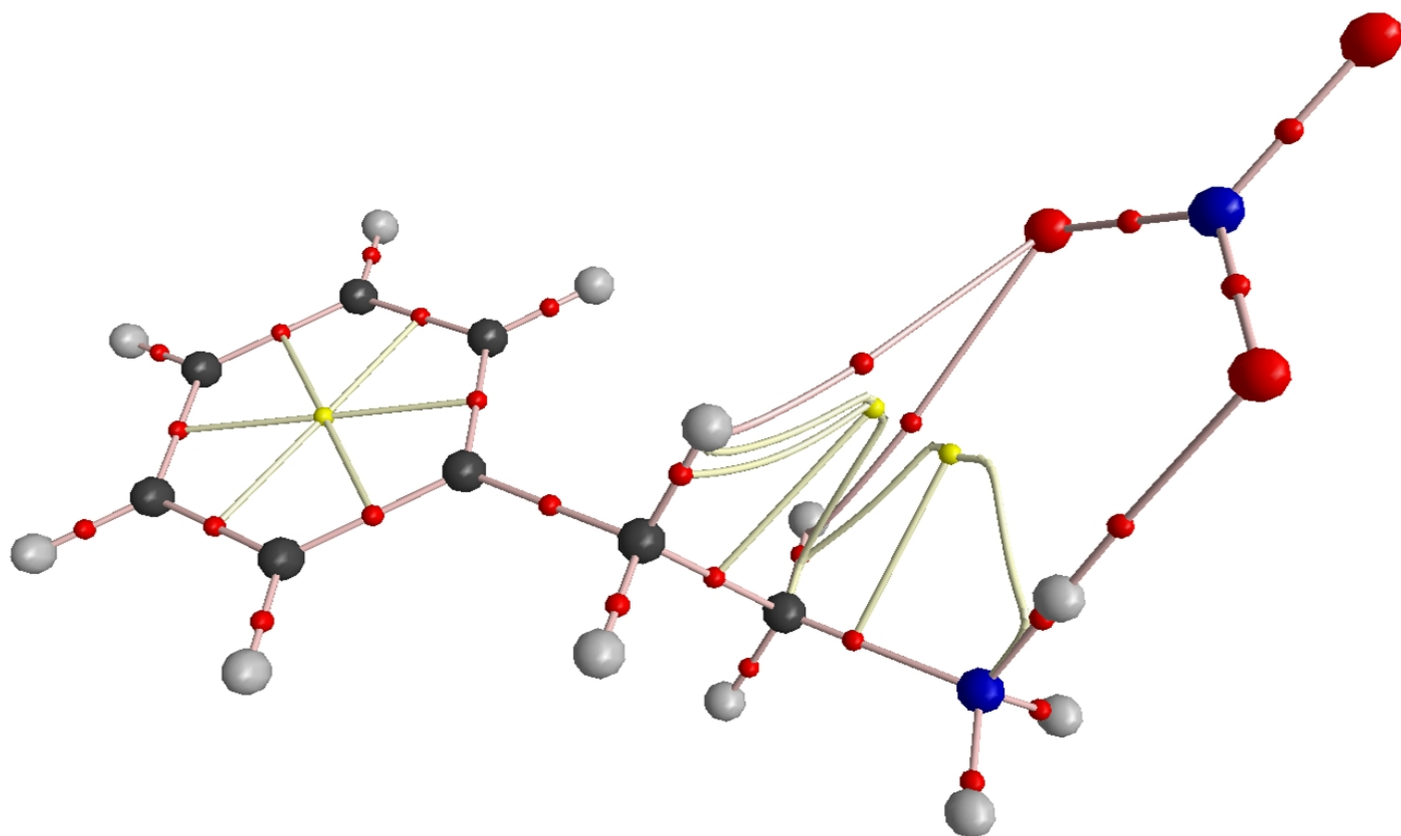


Fig. 5

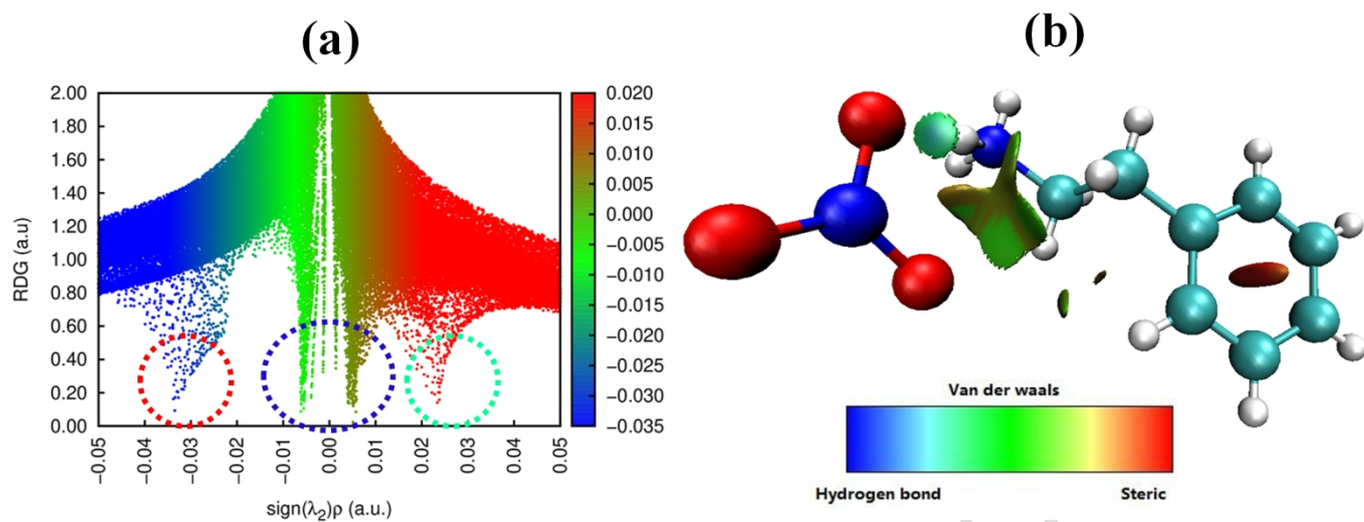


Fig. 6

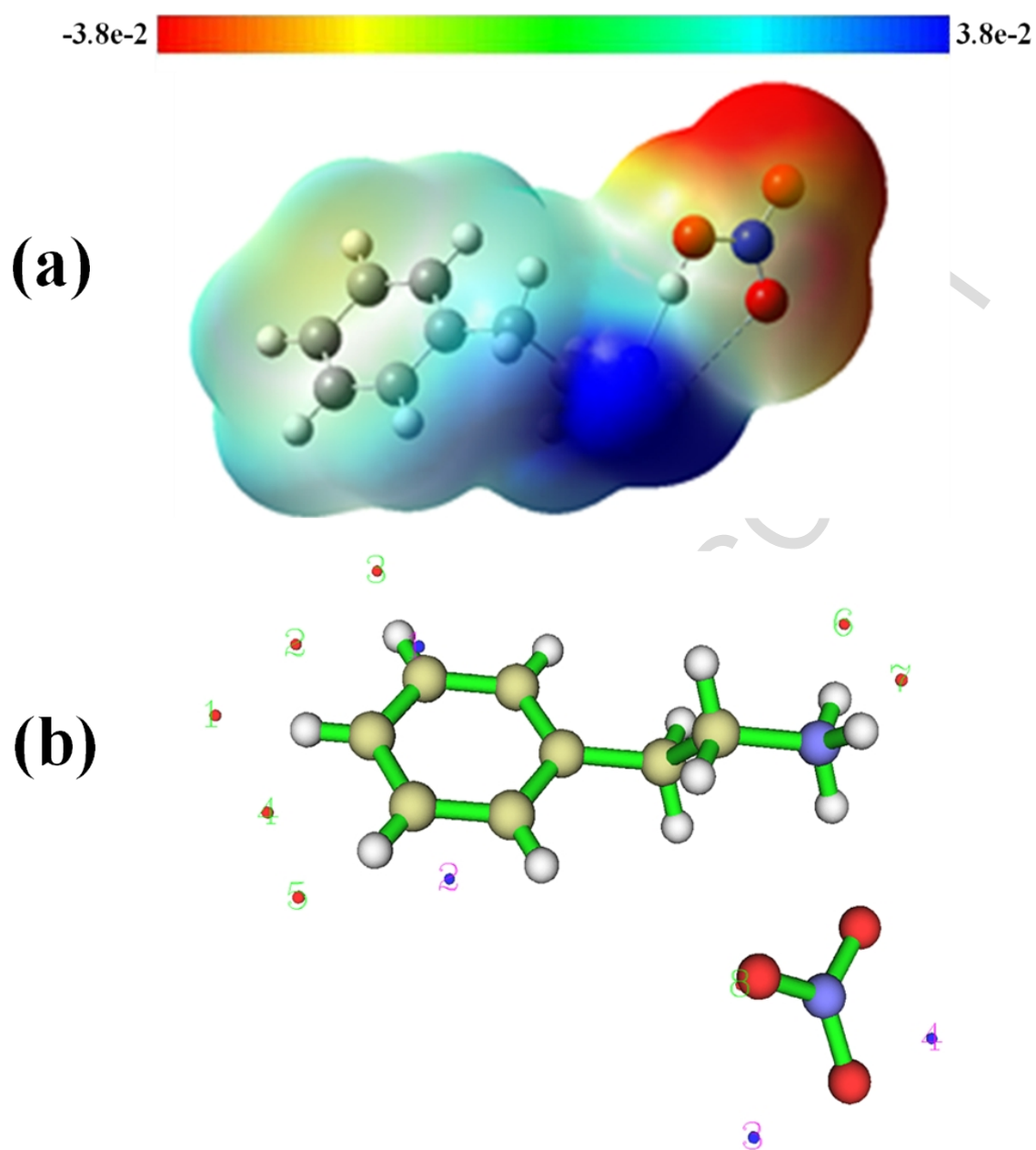


Fig. 7

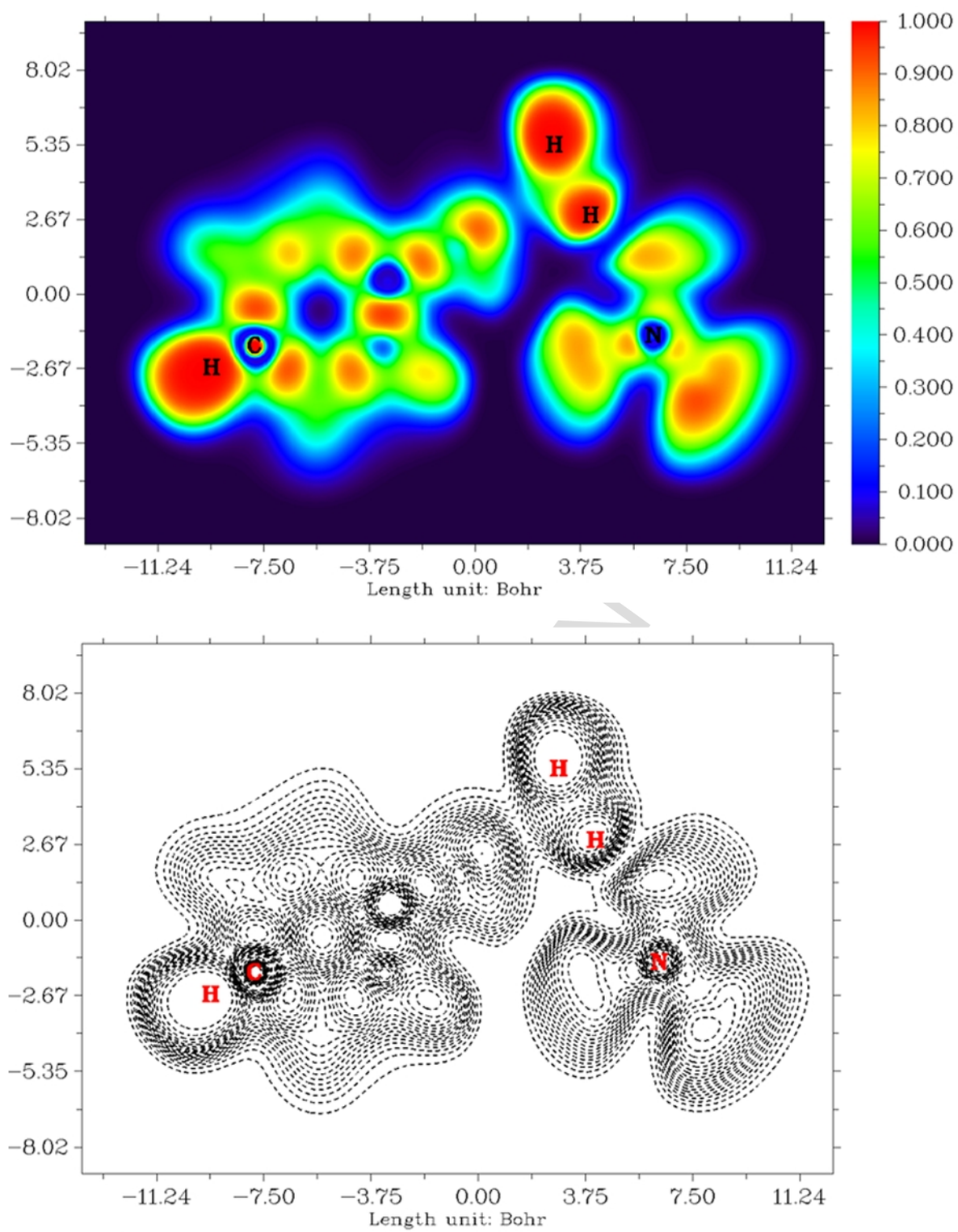


Fig. 8

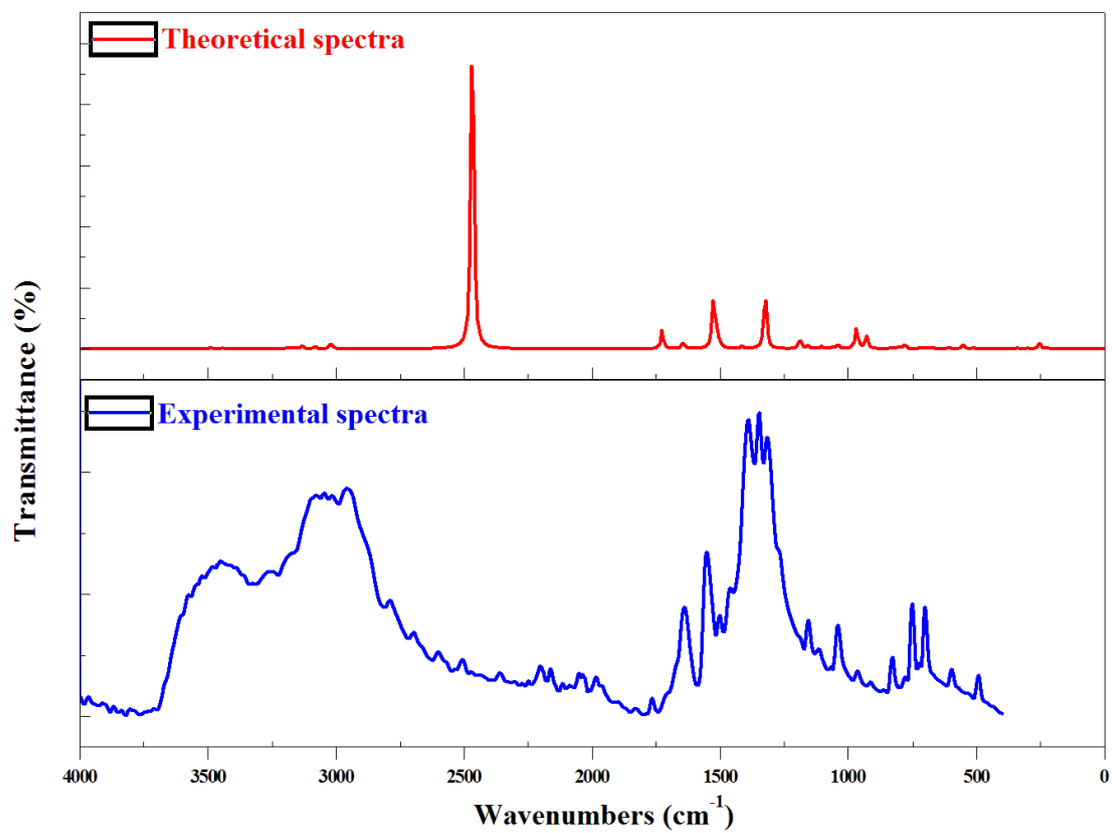


Fig. 9

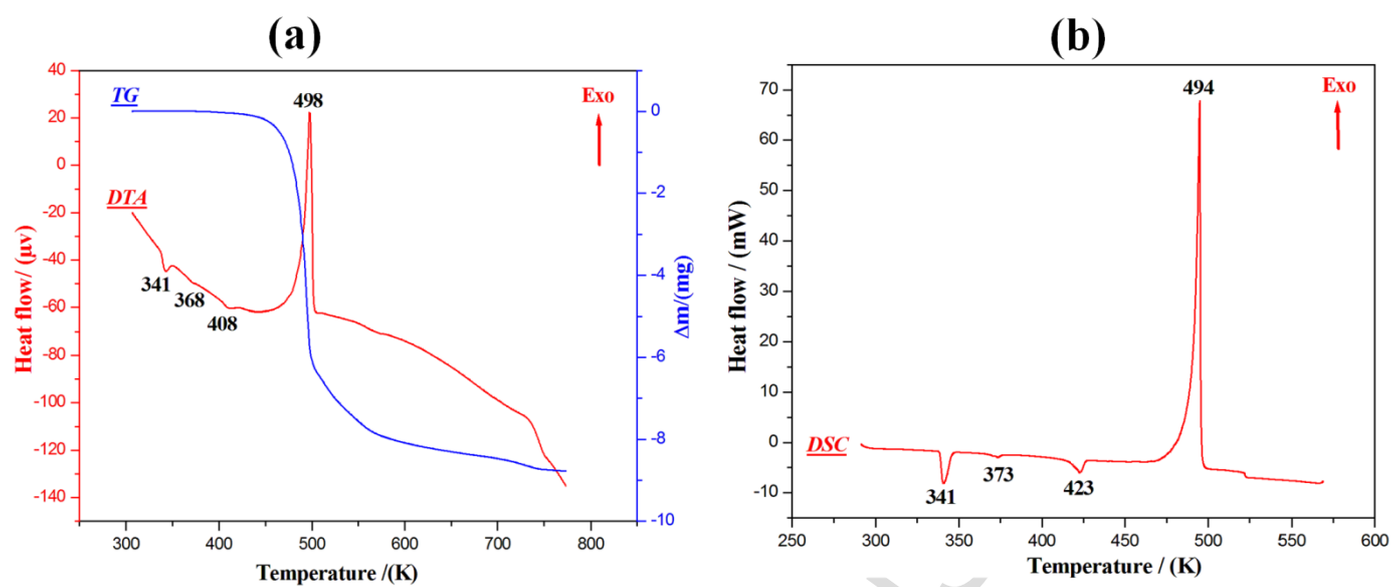


Fig. 10

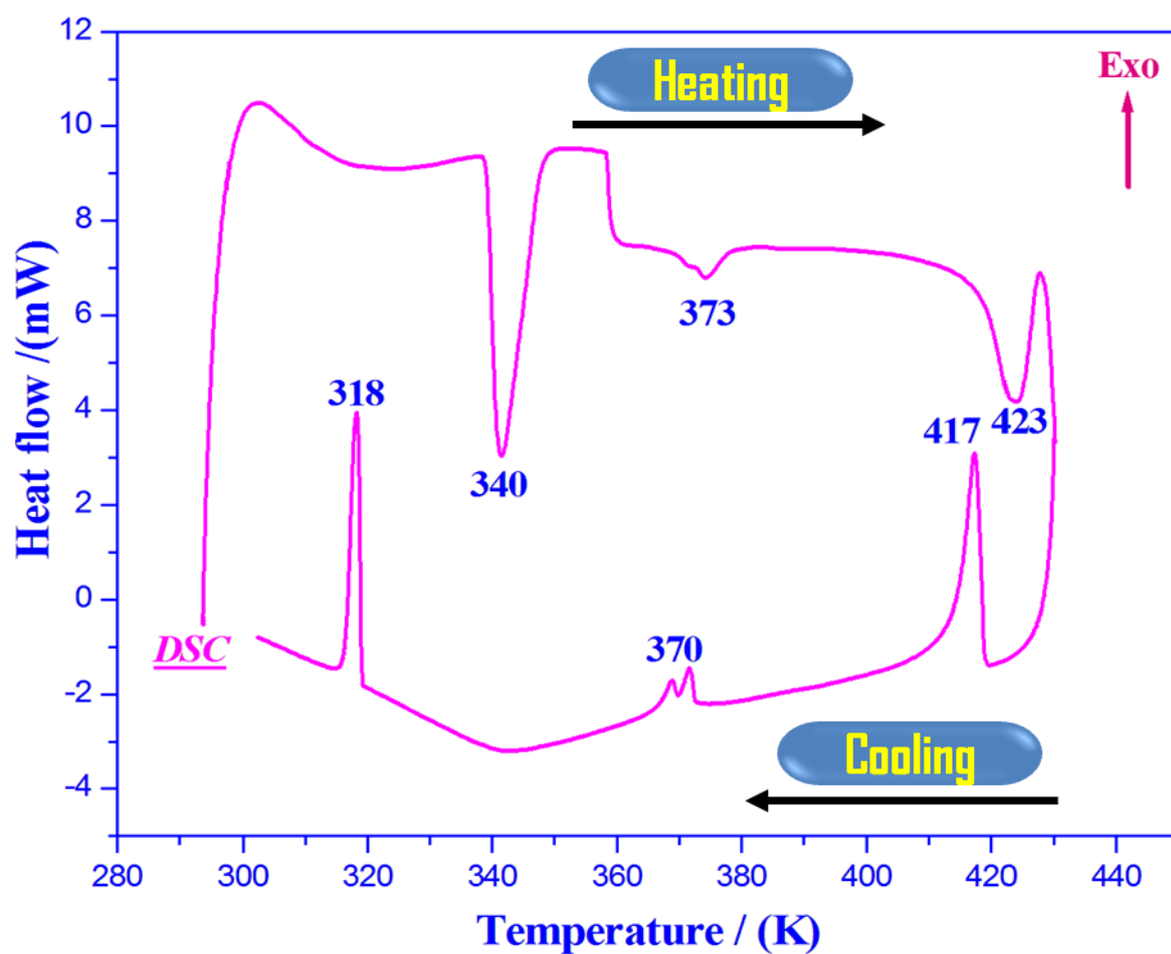


Fig. 11

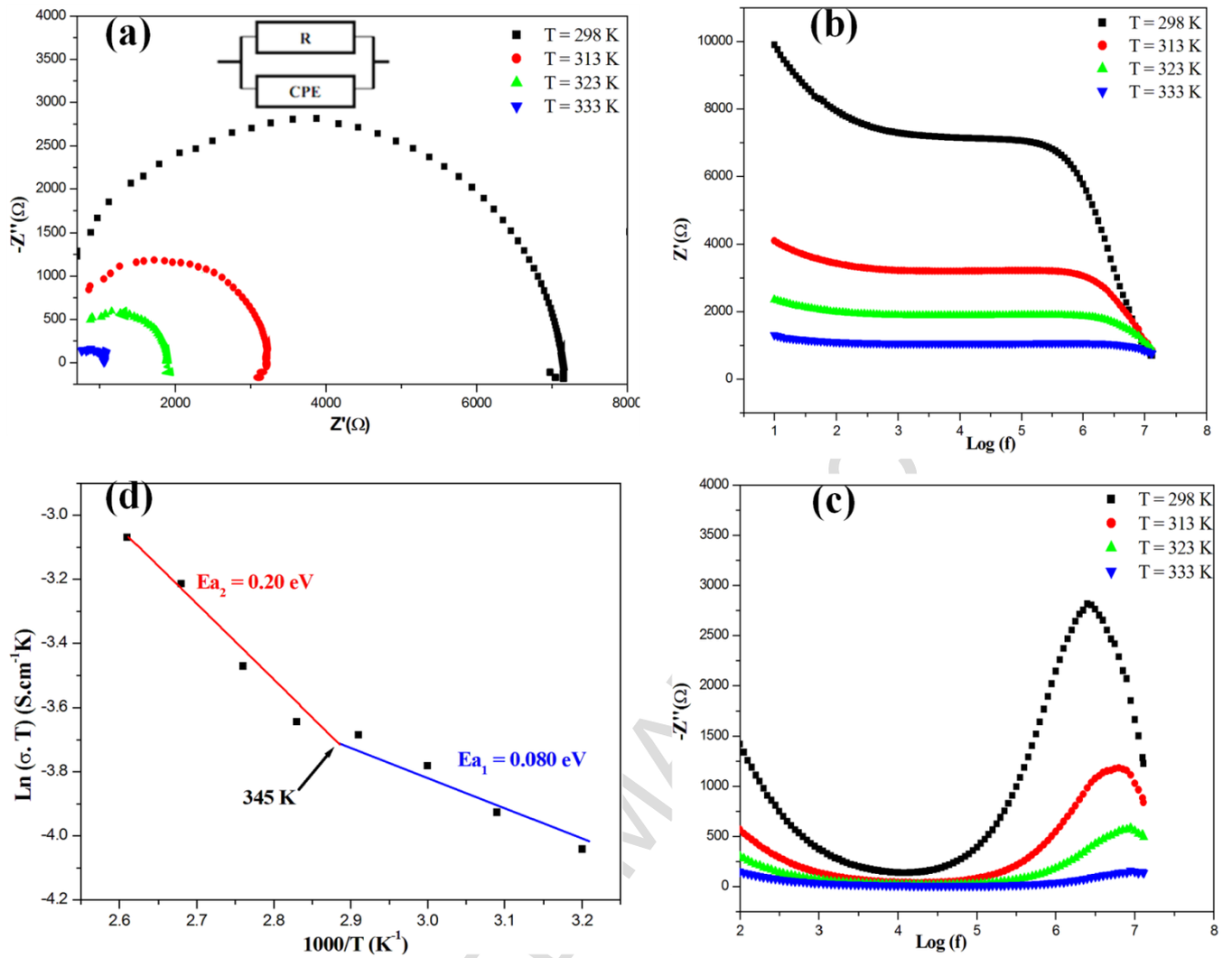


Fig. 12

Highlights

- Single crystals of PEAN were grown by slow evaporation-solution growth technique.
- Structural analysis was done by single crystal X-ray diffraction analysis.
- Thermal stability was investigated by thermal analyses.
- FT-IR spectra in the solid phase are recorded and analyzed.
- The optimized geometry and vibrational wavenumbers are computed using DFT method.
- Ionic conductivity was also investigated.

Table 1

CCDC	1839301
Temperature	150 K
Empirical formula	C ₈ H ₁₂ N ₂ O ₃
Formula weight (g mol ⁻¹)	184.20
Crystal system	Monoclinic
Space group	P2 ₁ /c
a	5.9497(8) Å
b	20.494(3) Å
c	8.4528(11) Å
β	113,812(8)°
Z	4
V	942.9(2) Å ³
F(000)	392
μ(Mo Kα)	0.100 mm ⁻¹
Index ranges	-7 ≤ h ≤ 5, -26 ≤ k ≤ 26, -9 ≤ l ≤ 10
Reflections collected	6632
Independent reflections	2195
Reflections with I > 2σ(I)	1539
R _{int}	0.043
Absorption correction: multi-scan	
Refined parameters	130
R[F ² > 2σ(F ²)]	0.049
wR(F ²)	0.125
Goodness of fit	1.09
Δρ _{max} 0.24 e Å ⁻³	Δρ _{min} -0.21 e Å ⁻³

Table 2

$D-H\cdots A$	$D-H$ (Å)	$H\cdots A$ (Å)	$D\cdots A$ (Å)	$D-H\cdots A$ (°)
N2—H1N \cdots O2 ⁱ	0.90(2)	2.01(2)	2.890(2)	166.4(18)
N2—H1N \cdots O1 ⁱ	0.90(2)	2.49(2)	3.208(3)	137.0(17)
N2—H2N \cdots O3	0.94(2)	1.88(2)	2.815(2)	178(2)
N2—H2N \cdots O2	0.94(2)	2.62(2)	3.254(2)	125.3(17)
N2—H3N \cdots O3 ⁱⁱⁱ	0.97(3)	1.93(3)	2.865(2)	159(2)
C1—H1A \cdots Cg ⁱⁱⁱ	0.97	2.735	3.554(2)	142.5
C2—H2B \cdots Cg ^{iv}	0.97	2.952	3.704(2)	135.2

Symmetry code: : (i) $x-1, y, z$; (ii) $x, -y+1/2, z+1/2$; (iii) $x-1, -y+1/2, z+1/2$; (iv) $x-1, -y+1/2, z-1/2$; Cg (gravity center of the ring C3-C4-C5-C6-C7-C8)

Table 3

	B3LYP/6-311++G(d,p)	B3LYP/6-311++G(3df,3pd)	Exp.
O3-N1	1.3683	1.3636	1.2639
O3-H2N	1.044	1.0386	1.8766
O2-N1	1.2237	1.2196	1.2514
O1-N1	1.2043	1.2014	1.2309
N2-C1	1.4808	1.4764	1.4884
N2-H1N	1.0174	1.0149	0.9005
N2-H2N	1.5996	1.6157	0.9383
N2-H3N	1.0175	1.0149	0.9757
C3-C4	1.3991	1.3953	1.3925
C3-C2	1.5129	1.5093	1.5085
C3-C8	1.3993	1.3955	1.3883
C4-H4	1.0859	1.0834	0.9305
C4-C5	1.3937	1.39	1.3857
C7-H7	1.0843	1.0818	0.9294
C7-C6	1.3939	1.3902	1.3863
C7-C8	1.3936	1.3899	1.3871
C2-H2A	1.0963	1.0935	0.9694
C2-H2B	1.0941	1.0912	0.9696
C2-C1	1.5349	1.5317	1.5191
C1-H1A	1.0964	1.0938	0.9701
C1-H1B	1.0929	1.0902	0.9702
C5-H5	1.0844	1.0818	0.9294
C5-C6	1.3937	1.39	1.3736
C6-H6	1.0841	1.0815	0.9296
C8-H8	1.0856	1.0831	0.9299
N1-O3-H2N	107.118	106.8928	109.9016
O3-N1-O2	117.1404	117.0397	118.8534
O3-N1-O1	115.4747	115.4672	119.9664
O2-N1-O1	127.3849	127.493	121.1771
C1-N2-H1N	110.5172	110.6249	110.5345

C1-N2-H2N	114.4809	114.3752	111.0567
C1-N2-H3N	110.6499	110.8219	111.2474
H1N-N2-H2N	112.9704	113.2256	109.1759
H1N-N2-H3N	106.9991	107.148	106.8079
H2N-N2-H3N	100.5516	99.9347	107.8794
C4-C3-C2	120.8862	120.9056	121.1216
C4-C3-C8	118.4454	118.3939	118.1313
C2-C3-C8	120.6496	120.6793	120.7466
C3-C4-H4	119.5501	119.5182	119.5249
C3-C4-C5	120.9011	120.9423	120.948
H4-C4-C5	119.5468	119.5377	119.5271
H7-C7-C6	120.0874	120.106	119.9963
H7-C7-C8	119.7711	119.7709	119.895
C6-C7-C8	120.1408	120.1224	120.1086
C3-C2-H2A	109.6094	109.6432	109.329
C3-C2-H2B	110.1261	110.135	109.3446
C3-C2-C1	112.1862	112.2326	111.3027
H2A-C2-H2B	106.874	106.93	108.047
H2A-C2-C1	109.0084	108.9109	109.3788
H2B-C2-C1	108.8857	108.8371	109.3765
N2-C1-C2	110.8169	110.8017	110.1046
N2-C1-H1A	111.7173	111.8248	109.6634
N2-C1-H1B	107.5261	107.6006	109.6678
C2-C1-H1A	109.7122	109.6023	109.6221
C2-C1-H1B	109.7502	109.7158	109.6153
H1A-C1-H1B	107.2166	107.1957	108.1373
C4-C5-H5	119.8184	119.8172	119.8078
C4-C5-C6	120.0893	120.0724	120.335
H5-C5-C6	120.0914	120.1096	119.8572
C7-C6-C5	119.5764	119.5801	119.5697
C7-C6-H6	120.2121	120.2099	120.2206
C5-C6-H6	120.2106	120.209	120.2096
C3-C8-C7	120.8465	120.8884	120.905
C3-C8-H8	119.5336	119.5019	119.5365
C7-C8-H8	119.618	119.6081	119.5586

Table 4

	O3...H1A	O3...H2A	O2...H2N	RCP	NRCP1	NRCP2
Density of electrons $\rho(\mathbf{r})$	0.00619	0.00556	0.03365	0.02356	0.0052	0.0057
Laplacian of electron density $\nabla^2 \rho(\mathbf{r})$	0.02270	0.01860	0.08946	0.16501	0.02130	0.02518
Lagrangian kinetic energy G(r)	2.96942	2.49820	16.02369	21.712721	2.65792795	3.15495108
Energy density H(r)	0.60600	0.43177	-1.93379	4.1828072	0.68325221	0.7946060
Potential energy density V(r)	-2.36343	-2.06643	-17.95747	-17.52991	-1.974676	-2.360345
Interaction energy (E_{int})	-1.18171	-1.03322	-8.97874			
Electron localization function (ELF)	0.01587	0.01562	0.13543	0.025115	0.010432	0.01065
Localized orbital locator (LOL)	0.11290	0.11213	0.28363	0.13844	0.09331	0.09418
Local information entropy	0.00061	0.00055	0.00274	0.002003	0.000514	0.00057
Reduced density gradient (RDG)	$0.12208 \cdot 10^{-14}$	$0.97634 \cdot 10^{-15}$	$0.11651 \cdot 10^{-14}$	$0.798467 \cdot 10^{-15}$	$0.23067 \cdot 10^{-14}$	$0.20652 \cdot 10^{-15}$
Average local ionization energy	0.45668	0.42441	0.50993	0.70658	0.46858	0.50783
Ellipticity (ϵ)	0.25625	0.39951	0.06388	-1.2073	-1.8530	-1.7866
Eta index (ζ)	0.13971	0.16475	0.27771	0.19040	0.14145	0.14131

Table 5.

Donor (i)	Acceptor (j)	^a E(2)	E(i)-E(j) ^b	F(i,j) ^c
$\pi(\text{O}_1\text{-N}_1)$	$\pi^*(\text{O}_1\text{-N}_1)$	10.05	0.55	0.078
LP(O ₃)	$\sigma^*(\text{O}_2\text{-N}_1)$	9.82	0.71	0.075
LP(O ₃)	$\sigma^*(\text{O}_1\text{-N}_1)$	15.20	0.64	0.090
LP(O ₃)	$\pi^*(\text{O}_1\text{-N}_1)$	47.66	0.27	0.108
LP(O ₂)	$\sigma^*(\text{O}_3\text{-N}_1)$	15.36	0.65	0.089
LP(O ₂)	$\sigma^*(\text{O}_1\text{-N}_1)$	14.40	0.60	0.086
LP(O ₂)	$\pi^*(\text{O}_1\text{-N}_1)$	55.10	0.26	0.112
LP(O ₁)	$\sigma^*(\text{O}_3\text{-N}_1)$	18.59	0.63	0.097
LP(O ₁)	$\sigma^*(\text{O}_2\text{-N}_1)$	17.51	0.66	0.096
$\pi^*(\text{O}_1\text{-N}_1)$	$\sigma^*(\text{O}_1\text{-N}_1)$	71.79	0.33	0.223
LP(O ₃)	$\sigma^*(\text{N}_2\text{-H}_2\text{N})$	14.04	0.78	0.094
$\pi(\text{C}_3\text{-C}_4)$	$\pi^*(\text{C}_7\text{-C}_8)$	19.99	0.29	0.068
$\pi(\text{C}_3\text{-C}_4)$	$\pi^*(\text{C}_5\text{-C}_6)$	20.17	0.29	0.069
$\pi(\text{C}_7\text{-C}_8)$	$\pi^*(\text{C}_3\text{-C}_4)$	22.22	0.28	0.071
$\pi(\text{C}_7\text{-C}_8)$	$\pi^*(\text{C}_5\text{-C}_6)$	20.03	0.29	0.068
$\pi(\text{C}_5\text{-C}_6)$	$\pi^*(\text{C}_3\text{-C}_4)$	20.36	0.28	0.068
$\pi(\text{C}_5\text{-C}_6)$	$\pi^*(\text{C}_7\text{-C}_8)$	20.20	0.29	0.068

^aE(2) means energy of hyper conjugate interactions(stabilization energy in Kcal/mol).

^bEnergy difference between donor and acceptor i and j NBO orbitals in a.u.

^cF(i,j) is the Fock matrix element between i and j NBO orbitals in a.u.

Table 6.

Number of surface minima	a.u	eV	Kcal/mol	X/Y/Z coordinate(Angstrom)		
1	-0.00522338	-0.142135	-3.277721	-3.740211	0.175737	1.827884
2	-0.01406535	-0.382738	-8.826148	-2.343773	-2.038873	-1.372020
3	-0.09900615	-2.694094	-62.127347	3.343670	-2.947025	0.592511
4	-0.09691886	-2.637296	-60.817551	5.162916	-0.343579	-1.602288
Number of surface maxima	a.u	eV	Kcal/mol	X/Y/Z coordinate(Angstrom)		
1	0.03026541	0.823564	18.991849	-6.168387	-1.885104	0.326353
2	0.02967241	0.807427	18.619735	-5.960049	-0.346093	-0.497468
3	0.03326189	0.905102	20.872169	-5.757719	1.376247	-1.446149
4	0.02701755	0.735185	16.953785	-4.574122	-2.672136	1.096269
5	0.02703892	0.735766	16.967192	-3.441940	-3.399866	1.781505
6	0.13218519	3.596942	82.947528	1.087223	3.935644	-0.082410
7	0.12700403	3.455955	79.696296	2.147701	2.857330	2.561221
8	-0.03031125	-0.824811	-19.020615	2.273719	-1.126966	-1.541740

Table 7.

Exp.	B3LYP/6-311++G(d,p) Method			Vibrational assignments (% PED)
	Unscaled	Scaled	I ¹	
3453	3553	3492	14.14	vNH 100
3267	3480	3421	5.29	vNH 99
3200	3195	3141	13.33	vCH 92
3100	3184	3129	25.84	vCH 85
3056	3174	3120	4.27	vCH 91
3018	3162	3108	5.64	vCH 84
2936	3158	3104	9.59	vCH 85
2789	3086	3034	25.3	vCH 87
2696	3066	3014	6.2	vCH 90
2602	3022	2971	39.28	vCH 78
2509	3014	2963	12.48	vCH 96
2373	2470	2428	2790.85	vOH 97
2205	1728	1699	123.75	vON 47 β HON 30
2162	1646	1577	52.94	δ HNHC 55 β HNH 40
2050	1644	1575	4.58	vCC 65 β HCC 20
1982	1622	1553	1.02	vCC 65 β HCC 11
1758	1530	1466	501.19	β HCC 62 β CCC 16
1640	1525	1461	9.75	vON 32 β HON 19 β HCH 21 γ OCNH 10
1547	1520	1456	28.6	vON 12 β HCH 61
1497	1495	1432	7.92	β HCH 87
1392	1486	1424	6.94	β HCC 52 vCC 16
1348	1416	1357	20.25	δ HCHN 50 β HCC 22
1317	1366	1309	0.4	β HCC 86
1156	1354	1297	0.64	δ HCCC 49 δ HNHC 10
1106	1341	1284	0.11	vCC 57 β HCC 25
1037	1324	1269	28.81	vON 52 β HCC 13
963	1322	1266	455.26	vON 30 β HCC 31
827	1269	1215	1.31	δ HCCC 28
746	1224	1173	0.3	β HCC 14 vCC 40
702	1205	1155	1.38	vCC 22 β HCC 70
597	1188	1138	114	β HNH 47 γ OCNH 11
491	1184	1134	0.57	vCC 12 β HCC 71
	1162	1114	24.46	δ HNHC 26 β HCC 15 β HNH 13
	1118	1071	17.34	β HCC 33 vCC 13
	1067	1022	9.31	β HCC 22 vCC 38
	1051	1007	6.11	β HCC 17 vCC 41 β CCC 24
	1041	997	30.05	vCC 10 vNC 56
	1014	971	0.62	vCC 19 β CCC 24 δ HCCC 36
	1012	969	0.02	vCC 14 β CCC 18 δ HCCC 45
	995	953	0.03	δ HCCC 83
	973	932	9.34	vCC 29 δ HCCC 13
	965	924	159.18	vON 46 β ONO 22
	930	891	119.72	δ HNHC 13 δ HCCC 13 β HNH 26
	928	889	6.42	δ HCCC 66
	863	827	0.06	δ HCCC 98
	832	798	4.42	vCC 19 β CCC 45

806	772	11.68	γ OOON 99
791	758	2.52	δ HCHN 11 δ HCCC 62
766	734	33.52	δ HCCC 57
711	681	42.91	δ HCCC 24 δ CCCC 49
697	667	6.14	ν ON 33 β ONO 55
668	640	10.15	β ONO 69
636	609	0.02	ν CC 11 β CCC 79
607	582	11.05	β CCN 49
514	492	19.06	γ CCCC 57 β CCC 11
465	446	15.16	δ HNHC 14 β HNH 43
415	397	0	δ HCCC 17 δ CCCC 72
349	334	1.94	β CCC 67
315	301	5.07	ν CC 10 β CCN 10 β HNH 40
288	276	20.98	β CCC 30 ν NH 18 γ CCCC 18
254	244	53.75	ν NH 56
150	144	5.28	β HON 10 δ CCCN 38
101	97	0.92	β HNH 11 β CCC 20 γ CCCC 19 γ OCNH 18
98	94	1.8	β CCC 12 ν NH 12 γ CCCC 13 γ OCNH 16
76	72	0.53	δ HONO 70
54	52	0.19	δ HNCC 63
31	30	0.36	δ CCCN 20 β HNC 37 δ CHON 26
23	22	0.99	δ HNCC 76
11	10	2.01	β HNC 16 δ CHON 66

I^i : infrared intensity ($\text{km}\cdot\text{mol}^{-1}$), ν , β , δ and γ denote stretching, in-plane bending, torsion and Out-of-plane bending modes, respectively. PED: potential energy distribution data are taken from VEDA4.

*Scaling factor: from 4000 to 1700 cm^{-1} are scaled with 0.983 and lower than 1700 cm^{-1} are scaled with 0.958 ,