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COMPUTATIONAL STUDY ON THE ELECTRONIC STRUCTURE OF PHENETHICILLIN-LACTIM ZWITTERIONS BY AUSTIN MODEL-1 (AM1) METHOD

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ABSTRACT

The geometry, conformation and electronic structure of phenethicillin-lactim and its zwitterions have been optimized and calculated by semi-empirical molecular orbital method (AM1), which includes experimental parameters and extensive simplification of the Schrodinger's equation (H Ψ =E Ψ) for calculation of various properties in the gas phase. The mechanism of formation of zwitterions has been studied. At this instant, the heats of formation (ΔH_f^o), dipole moment (μ), ionization potential (IP), full atomic charges and energies of frontier molecular orbitals (E_{HOMO} and E_{LUMO}) have been performed and discussed. The conformational changes and electronic properties have also been discussed for stable conformations.

KEYWORDS: Phenethicillin-lactim, zwitterions, HOMO, LUMO, frontier molecular orbitals.

INTRODUCTION

Phenethicillin (Broxil) is one of the penicillin derivatives and studied extensively due to their favourable absorption patterns and reduced undesirable side effects. [1] It is particularly used in the treatment of gonorrhoea [2] and the chemotherapy of bacterial infections compared with other semi-synthetic penicillin derivatives. [3] Austin Model-1 (AM1) is one of the semi-empirical methods with using experimental parameters and extensive simplification of the Schrodinger's equation ($H\Psi=E\Psi$) to optimize molecules for calculation of various properties

to solve chemical problems.^[4] Gas phase quantum calculations can reproduce the essential features of zwitterions for incorporating solvent effects into theoretical calculations on chemical systems to obtain agreement with experiments.^[5] In this way quantum chemistry simulates chemical structure and reactions numerically and allows studying chemical phenomena by running calculations on computer rather than examining reactions experimentally.^[6]

In continuation of research $work^{[7]}$ and based on these observations, the present study on molecular conformation and electronic properties of phenethicillin-lactim (1) and its zwitterions (2 and 3) in gas phase has been evaluated by AM1 method. The mechanism of formation of zwitterions RH^{\pm} (2 and 3) have been discussed.

Computational methods^[4]

Semi-empirical molecular orbital calculations were performed using Austin Model-1 (AM1). Geometry calculations in the ground state (key words: GNORM=5, MMOK, GEO-OK, CHARGE, and PRECISE) were completely optimized until get the lowest energy conformation. The initial molecular geometry was adopted as Pople's standard data^[8], and subsequently using fully optimized energy gradient method. The conformations were designated by Klyne-Prelog terms^[9] using s = syn, a = anti, $c = clinal (0\pm30^{\circ} \& 180\pm30^{\circ})$ and all other angles p = peri-planar.

RESULTS AND DISCUSSION

Electronic structure of phenethicillin-lactim (RH, 1) and its zwitterions (RH[±], 2&3)

The optimized electronic structure of Phenethicillin-lactim \mathbf{RH} (1) and its zwitterions $\mathbf{RH}^{\pm}(2\&3)$ are shown in Scheme-1. In this context, the numbering of phenethicillin-lactim is shown in Figure -1. The calculated heats of formation (ΔH_f^o), ionization potential (IP), dipole moment (μ), the energies of frontier molecular orbitals (E_{HOMO} and E_{LUMO}) and net charges on hetero atoms of the molecules (1 to 3) are presented in Table-I. It is observed that the net charges on N_7 - and N_{12} -atoms are -0.2249 and -0.2517 respectively in the case of phenethicillin-lactim (1). Usually, the sequence of protonation for nitrogen atoms of phenethicillin-lactim (1) is observed in the order of $N_7 < N_{12}$. It is also observed that ionization potential values are increased in the order of 3 < 1 < 2.

Figure - 1

The calculated values of frontier orbital energies (E_{HOMO} and E_{LUMO}) reveal the promotion of an electron from HOMO to LUMO, in a photochemical reaction, the supra-facial path way is allowed, due to the presence of same sign. The results revealed that the electronic properties and reactivity of molecule depend on its conformational structure. The dipole moments of molecules depend on the nature of the atoms and bonds comprising the molecules and on their arrangement. The dipole moment is increasing in the order of 1 < 2 < 3 and zwitterion (3) showed higher dipole moment. The electronegative hetero-atoms cause displacement of electrons that induces an additional dipole moment in the molecule. The magnitude of the induction effect [11] (μ_{ind}) of molecules can be estimated with respect to phenethicillin-lactim (1) by the equation (1).

Induction effect
$$(\mu_{ind}) = \mu(\mathbf{RH}^{\pm}) - \mu(\mathbf{RH})$$
 ----- (1)

It is found that the induction effect is increasing in the case of $\Delta\mu_{ind}$ 2.811D(2) $<\Delta\mu_{ind}$ 12.216D(3). According to the heat of formation (ΔH_f^o) data, the stability of compounds have been increased in the order of 3 < 2 < 1. It is investigated that the phenethicillin-lactim (1) is more stable than zwitterions (2 and 3). But geometry calculations in the ground state were completely optimized until the lowest energy conformation was found in the individual ions or molecules. It can be assumed that the electronic properties and reactivity of the molecule depend on its conformational structure. It is predicted that the protonation would take place preferably at N_{12} -atom than N_7 -atom in the case of phenethicillin-lactim (1). But, it is found that the stability of zwitterion $N_{12}H^{\pm}$ (2) (ΔH_f^o , -8.9408 kcal/mol) is more stable than N_7H^{\pm} (3) (ΔH_f^o , -43.3190 kcal/mol).

Table –I: Heat of formation (ΔH_f^0 in kcal/mol), ionization potential (eV), dipole moment (μ in Debye), energies of frontier molecular orbitals (in eV) and the atomic charges on hetero-atoms of phenethicillin-lactim(1) and its zwitterions (2&3) from AM1 calculations.

Parameters	1	$2(N_{12}H^{\pm})$	$3(N_7H^{\pm})$			
ΔH _f ° (kcal/mol)	-109.6539	-100.7131	-66.3349			
Ionization potential (eV)	9.1089	9.2531	8.9348			
μ (Debye)	2.327	5.138	14.543			
E _{HOMO} (eV)	-9.109	-9.253	-8.935			
E_{LUMO} (eV)	-0.081	-0.009	-0.793			
Electron excitation energies (eV)	9.028	9.244	8.142			
S ₂ (atomic charge)	+0.0281	+0.0823	+0.0568			
N ₇ (atomic charge)	-0.2249	-0.1825	-0.0570			
N ₁₂ (atomic charge)	-0.2517	-0.2334	-0.2528			
O ₁₀ (atomic charge)	-0.2807	-0.2538	-0.5038			
O ₁₅ (atomic charge)	-0.2751	-0.2934	-0.3025			
O ₁₆ (atomic charge)	-0.2020	-0.2276	-0.2350			
O ₃₂ (atomic charge)	-0.3561	-0.2047	-0.4635			
O ₃₃ (atomic charge)	-0.2280	-0.2756	-0.0956			
Bold values indicates	Higher values					

In the case of formation of zwitterions (2 and 3) is considered by the removal of a proton from O_{10} -atom of phenethicillin-lactim (1) and the protonation at N_{12} - atom in the case of $N_{12}H^{\pm}$ (2) is considered by decreasing net atomic charges at N_7 -, N_{12} -, O_{10} - and O_{32} -atoms and increasing at O_{15} -, O_{16} - and O_{33} - atoms. The protonation site of phenethicillin-lactim (1) at N_7 -atom is considered in the case of N_7H^{\pm} (3) by increasing net atomic charges at N_{12} -, O_{10} -, O_{15} -, O_{16} - and O_{32} - atoms and decreasing at N_7 - and O_{33} - atoms.

Equilibrium of phenethicillin-lactim(RH, 1) and its zwitterions(RH $^{\pm}$, 2&3) with their conformational analysis

Equilibrium is typically found in polar solvents by rapid inter- or intra-molecular proton transfer from O_{10} - atom to N_7 - or N_{12} - atoms of phenethicillin-lactim (1) in the formation of zwitterions \mathbf{RH}^{\pm} (2 and 3) and it is established as per Scheme-1. The exact proton-migration in phenethicillin-lactim (1) has been observed in accordance with the negative charge distribution on N-atoms (Table-1). Thus, formed zwitterions \mathbf{RH}^{\pm} (2 and 3) with the protonation at N_7 - or N_{12} - atoms of phenethicillin-lactim (1) can be assigned by *anti*- or *syn*-conformations with the comparison of their geometry and electronic structure. The proton affinity $(PA)^{12}$ values for the different nitrogen atoms of phenethicillin-lactim \mathbf{RH} (1) were calculated as per the equation (2).

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$$PA = \Delta H_f^o(H^+) + \Delta H_f^o(B) - \Delta H_f^o(BH^+) - - - - (2).$$

Where PA is the proton affinity, $\Delta H_f^o(B)$ is the heat of formation for phenethicillin-lactim, $\Delta H_f^o(BH^+)$ is the heat of formation for the cation, and $\Delta H_f^o(H^+)$ is heat of formation for the proton (367.2kcal/mol). In the formation of zwitterions, it can be assumed that $\Delta H_f^o(H^+)$ is to be neglected in the inter- or intra-molecular proton transfer in the equilibrium as per equation (3).

Thus, the proton affinity (PA)^[13] becomes

$$PA = \Delta H_f^{o}(\mathbf{R}\mathbf{H}) - \Delta H_f^{o}(\mathbf{R}\mathbf{H}^{\pm}) \qquad \dots (4).$$

Where $\Delta H_f^o(\mathbf{R}\mathbf{H})$ is the heat of formation of phenethicillin-lactim $\mathbf{R}\mathbf{H}$ (1) and $H_f^o(\mathbf{R}\mathbf{H}^{\pm})$ is the heat of formation of zwitterions $\mathbf{R}\mathbf{H}^{\pm}$ (2 and 3). The proton affinity is found 8.9408 kcal/mol and 43.3190 kcal/mol respectively in the case of zwitterions $\mathbf{N}_{12}\mathbf{H}^{\pm}$ (2) and $\mathbf{N}_7\mathbf{H}^{\pm}$ (3).

The spatial arrangement of atoms in a molecule is considered to study the conformations of phenethicillin-lactim (1), and its zwitterions (2 & 3). In this context, it is investigated that *anti*- or *syn*- conformations are existed according to the position of atoms. The change in energy content may depend upon the changes in the dihedral angles. The atomic numbering of phenethicillin-lactim (1) is revealed as per Figure-1 and incorporated the most important data of dihedral angles (Table - II) of molecules (1 to 3) for the sake of discussion.

Scheme-1

Dihedral angle	1		$2(N_{12}H^{\pm})$		$3(N_7H^{\pm})$	
(°)	Angle	(*)	Angle	(*)	Angle	(*)
$C_4C_3S_2C_1$	-21.14	-SC	+8.64	+sc	-28.72	-sc
$C_8C_4C_3S_2$	+163.15	+ap	+103.34	+ap	+166.21	+ <i>ac</i>
$O_{10}C_8C_4C_3$	-168.67	-ac	+66.09	+ <i>sp</i>	-161.22	-ac
$C_{13}N_{12}C_{11}C_{9}$	+160.93	+ac	-15.00	-sc	+127.76	+ <i>ap</i>
$C_{14}C_{13}N_{12}C_{11}$	+177.46	+ac	+141.01	+ <i>ap</i>	+178.61	+ac
$O_{15}C_{13}N_{12}C_{11}$	-0.07	-sc	-90.06	-ap	-2.27	-SC
$O_{16}C_{14}C_{13}N_{12}$	+146.27	+ap	+164.39	+ <i>ac</i>	-167.79	-ac
$C_{17}C_{14}C_{13}N_{12}$	+29.57	+sc	+48.25	+s p	+75.71	+ <i>sp</i>
$C_{18}O_{16}C_{14}C_{13}$	+85.90	+sp	+119.84	+ <i>ap</i>	+98.59	+ <i>ap</i>
$O_{32}C_8C_4C_3$	+17.01	+sc	+66.09	+ <i>sp</i>	+21.92	+sc
$H_{45}O_{15}C_{13}N_{12}$	-3.49	-sc	-179.89	-ac	+176.34	+ <i>ac</i>

Table – II: Dihedral angle (°) of phenethicillin-lactim (1) and its zwitterions (2&3) from AM1 calculations.

+102.51

+ap

- -

-159.25

- -

-ac

- -

-179.99

 $H_{34}O_{10}C_8C_4$

 $HN_{12}C_{11}C_9$

 $HN_7C_4C_3$

Bold values indicates – change of conformation / dihedral angle from phenethicillin-lactim (1) to zwitterions (2 & 3).

From the Table-II and Scheme-1, the zwitterion $N_{12}H^{\pm}$ (2) is formed by the transfer of a proton from O_{10} -atom to N_{12} - atom of phenethicillin (1). It is observed that conformation *-sc* of $C_4C_3S_2C_1$, *-ac* of $O_{10}C_8C_4C_3$, *+ac* of $C_{13}N_{12}C_{11}C_9$, *+ac* of $C_{14}C_{13}N_{12}C_{11}$ and *+sc* of $C_{17}C_{14}C_{13}N_{12}$ are changed to +sc, +sp, -sc, +ap and +sp conformations respectively. The dihedral angle of +ap of $O_{16}C_{14}C_{13}N_{12}$, *-sc* of $O_{15}C_{13}N_{12}C_{11}$, +sp of $C_{18}O_{16}C_{14}C_{13}$, +sc of $O_{32}C_8C_4C_3$, and *-ac* of $H_{45}O_{15}C_{13}N_{12}$ are changed to +ac, -ap, +ap, +sp and -ac conformation and found the rest of positions have a little changes. It is also investigated that the protonation at N_{12} - atom is shown +ap conformation in the case of $HN_{12}C_{10}C_9$.

If the phenethicillin zwitterion N_7H^{\pm} (3) is formed by the transfer of a proton from O_{10} -atom to N_7 - atom of phenethicillin-lactim (1), with the conformation +ap of $C_8C_4C_3S_2$, +ac of $C_{13}N_{12}C_{11}C_9$ and +sc of $C_{17}C_{14}C_{13}N_{12}$ are changed to +ac, +ap and +sp conformations respectively. The dihedral angle of +ap of $O_{16}C_{14}C_{13}N_{12}$, +sp of $C_{18}O_{16}C_{14}C_{13}$ and -sc of $H_{45}O_{15}C_{13}N_{12}$ are changed to -ac, +ap and +ac conformations and observed the rest of positions have moderate changes. It is found that the protonation at N_7 -atom is shown -ac conformation in the case of $HN_7C_4C_3$.

^{*} Conformational analysis using prefixes a = anti, s = syn, p = peri-planar, c = clinal, and + & - signs. [9]

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