# Theoretical Study of Some Chemical Kinetics and Physical Properties for some bnzohydrazide derivatives by using Semi-Empirical calculation (PM3 Method) 

# الاراسة النظرية للحركيات الكيميائية وبضض الخواص الفيزيائية لبحض مشتقات البنزو هيدرازيت باستخدام الحسابات الثبه الثقريبية (طريقة PM3) 

Nedaa A Jasim

Dep. Of chemistry, college of Education, university of Karbala, Iraq .


#### Abstract

A theoretical study in the present work involved program(Gaussian 03W) to use the method of calculating the total Semi-Empirical calculations(PM3 Method) of two benzohydrazide derivatives [ $\mathrm{N}^{\prime}$-(4-aminobenzylidene) benzohydrazide] ,[ $\mathrm{N}^{\prime}$-(4-(dimethylamino benzylidene) benzohydrazide], in order to elucidate the inhibition efficiencies and reactive sites of these compounds. They include calculations of dimensional geometry (lengths and bond angles) when the geometry of balanced. For these compounds some physical properties were calculated such as orbital energies ( $\mathrm{E}_{\text {LUмо }}, \mathrm{E}_{\text {номо }}$ ), energy gap $(\Delta E)$, dipole moment ( $\mu$ ), IP (ionization energy), hardness. Functions thermodynamic and IR intensity and absorption frequencies for tow compounds were also investigated. The purpose of this study to compare the possible sites for nucleophilic and electrophilic attacks and which compounds can be used anti-bacterial and antifungusthe. The results showed compound [ N '-(4-(dimethylamino benzyliden) benzohydrazide ] have lower electronegative, higher hardness $(\mathrm{n})$, lower energy gap $(\Delta E)$, and high vibrational frequencies of IR absorption compared with the other compound, which may be used as a highest anti- inflammatory. this difference in this results come according to the effect of subsisted groups of studied compounds .


الخلاصة
اعتمدت الار اسة النظرية في هذا العمل برنامج (Gaussian 09 W) ، وباستخدام طريقة الحساب شبه التجريبية
[N'-(4-aminobenzylidene bnzohydrazide] لأثنين من مشتقات البنزو هيبرازيت PM3
 الفعالة لهذين المركبين ، حيث تضمنت الاراسة الحسابات للابعاد الهندسية (اطوال وزو ايا التآصر ) عند الثكل الهندسي المتو ازن وتم حساب بعض الخصائص الفيز يائية لهذه المركبات مثل طاقة الاوربيتالات (ELUMO,
 لهذين المركبين . ان الغرض من الدر اسة هو المقارنة بالمو اقع الفعالة التي يككن مهاجمتها لللنيوكليوفيلات والالكترو فيلات [N'-(4-) benzohydrazide [ أي المركبات يمكن استخدامها كمضاد للابكتريا والفطريات. أظهرت النتائج ان المركب (dimethylamino benzylidene في منطقة امتصاص IR مع المركب الاخر ، لذا يككن ان يستخدم كاعلى مضاد حيوي . هذا التغاير في النتائج يعزى لتأثير المجمو عة المعوضة للمركبات التي تم دراستها.

## Introduction

Hydrazones have been demonstrated to possess antimicrobial, anticonvulsant, analgesic, antiinflammatory, antiplatelet, anti-tubercular, anticancer and antitumor activities ${ }^{[1,2]}$. Hydrazones possessing an azomethine $-\mathrm{NHN}=\mathrm{CH}-$ proton constitute an important class of compounds for new drug development. Many researchers have therefore synthesized these compounds as target structures and evaluated their biological activities. These observations have served as guides for the development of new hydrazones that possess various biological activities ${ }^{[3]}$. Hydrazide and their heterocyclised products display diverse biological activities including antibacterial, antifungicidal,

## Journal University of Kerbala, Vol. 14 No. 3 Scientific . 2016

analgesic, anti-inflammatory properties1-15. These heterocyclic systems find wide use in medicine, agriculture and industry.

Benzohydrazide and their derivatives are poly functional molecules ${ }^{[4,5]}$ bearing $\mathrm{C}=\mathrm{O},-\mathrm{NH}-$ and $-\mathrm{NH}_{2}$ functionalities in their structures. Therefore, we can expect these compounds to behave as ambident nucleophiles in several reactions ${ }^{[6]}$. The scarce data about their kinetics and reaction mechanisms prompted us to perform experimental and theoretical studies in order to gain insight in the corresponding reaction mechanisms towards carbonylic electrophiles, p-nitrophenyl acetate in the present case. We have shown that experimental and theoretical integrated research may become a very useful methodology to describe reaction mechanisms in organic chemistry ${ }^{[7]}$.


Figure 1: Benzohydrazide structure
Hydrazide and their heterocyclised products display diverse biological activities including antibacterial, antifungicidal, analgesic, anti-inflammatory properties1-15. These heterocyclic systems find wide use in medicine, agriculture and industry ${ }^{[8,9]}$.

## Methods of calculation.

The present study is based on G09 program ${ }^{(14)}$ using PM3 (Modified Neglect of Diatomic Overlap Parametric Method Number 3 ) model which was developed and the treatment of organic molelcules ${ }^{(10,11)}$. Such treatment yields the equilibrium geometry energy values of the molecules in addition to their fundamental vibration frequencies (3N-6) and IR absorption intensities ${ }^{(12)}$. Solution of this equation yields vibration frequencies $\lambda=4 \pi^{2} v^{2} c^{2)(13)}$. Depending on this method some physical properties were calculated according to these equations $\mathrm{E}_{\mathrm{A}}=-\mathrm{E}_{\mathrm{LUMO}}, \eta=1 / 2$ ( $\left.\mathrm{E}_{\text {Hомо }}-\mathrm{E}_{\text {LUMO }}\right),\left(\mathrm{G}^{0}=\mathrm{H}^{\mathrm{O}}-\mathrm{ST}\right),\left(\mathrm{A}^{0}=\mathrm{G}^{0}-\mathrm{ST}\right)$.
G09 program of J. Frisch, G. W. Trucks, et al ${ }^{(14,15)}$ was applied throughout.

## Results and Discussion

The benzohydrazide derivatives investigated in the present work are:

## Compound $1 \quad \mathbf{N}^{\prime}$-(4-aminobenzylidene)benzohydrazide <br> Chemical Formula: $\mathrm{C}_{14} \mathrm{H}_{\mathbf{1 3}} \mathrm{N}_{3} \mathrm{O}$ <br> Molecular Weight: 239.27

## Comopound 2

$\mathbf{N}^{\prime}$-(4-(dimethylamino)benzylidene)benzohydrazide Chemical Formula: C16H17N3O
Exact Mass: 267.14


Figure 2: The geometric equilibrium for $\mathrm{N}^{\prime}$-(4-aminobenzylidene) benzohydrazide and $\mathrm{N}^{\prime}$-(4(dimethylamino benzylidene) benzohydrazide

In this research calculated the geometry (bond lengths ) of the two molecules o $\mathrm{N}^{\prime}$-(4aminobenzylidene)benzohydrazide and $\mathrm{N}^{\prime}$-(4-(dimethylamino benzylidene) benzohydrazide , using the semi-empirical PM3 methods. According to the results calculated and recorded in the (table 1 and Fig. 3)
Show that each th bonds $\mathrm{C}_{10}-\mathrm{N}_{13}$ compound 2] $\mathrm{N}^{\prime}$-(4-(dimethylamino benzylidene)] has high value compared to other compound 1 [ $\mathrm{N}^{\prime}$-(4-aminobenzylidene)benzohydrazide] may be caused by the less electron- negative of $\mathrm{N}\left(\mathrm{CH}_{3}\right)_{2}$. Also the change of the group substituted had effect on the value of th angles of the studied compound in this research have shown calculation in the (table 1 and Fig 3 ). That the angle ( $\angle \mathrm{C}_{9} \mathrm{C}_{10} \mathrm{~N}_{13}$ ) showed that in compound 2 has high value compared to compound 1. May be due to the big size of substitution groups on the angles.


Figure 3: The geometric equilibrium (3D) the $\mathrm{N}^{\prime}$-(4-aminobenzylidene)benzohydrazide (Compound 1) and $\mathrm{N}^{-}$-(4-(dimethylamino benzylidene) benzohydrazide(Compound 2)

## Journal University of Kerbala, Vol. 14 No. 3 Scientific . 2016

Table 1 : Calculated geometric parameters (bond lengths in Angstrom length angles in degree) of the N'-(4-aminobenzylidene)benzohydrazide and $\mathrm{N}^{\prime}$-(4-(dimethylamino benzylidene) benzohydrazide


| $\mathrm{A}(10-11-12)$ | 128.6 | $\mathrm{~A}(6-14-16)$ | 124.2 |  |  | $\mathrm{~A}(13-20-36)$ | 110.4 |
| :---: | :---: | :---: | :---: | :--- | :--- | :---: | :---: |
| $\mathrm{~A}(10-11-25)$ | 114.2 | $\mathrm{~A}(8-7-12)$ | 119.4 |  |  | $\mathrm{~A}(13-20-37)$ | 113.2 |
|  |  |  |  |  |  |  |  |
| $\mathrm{~A}(12-11-25)$ | 117.2 | $\mathrm{~A}(8-7-18)$ | 118.7 |  |  | $\mathrm{~A}(15-14-16)$ | 121.7 |
| $\mathrm{~A}(11-12-13)$ | 121.6 | $\mathrm{~A}(7-8-9)$ | 120.4 |  |  | $\mathrm{~A}(14-15-17)$ | 117.9 |
| $\mathrm{~A}(11-12-17)$ | 118.7 | $\mathrm{~A}(7-8-26)$ | 119.7 |  |  | $\mathrm{~A}(14-15-30)$ | 114.0 |
| $\mathrm{~A}(13-12-17)$ | 119.8 | $\mathrm{~A}(12-7-18)$ | 121.8 |  |  | $\mathrm{~A}(17-15-30)$ | 114.7 |
| $\mathrm{~A}(12-13-14)$ | 120.3 | $\mathrm{~A}(7-12-11)$ | 120.3 |  |  | $\mathrm{~A}(15-17-18)$ | 123.1 |
| $\mathrm{~A}(12-13-26)$ | 120.1 | $\mathrm{~A}(7-12-29)$ | 120.3 |  |  | $\mathrm{~A}(17-18-31)$ | 114.7 |
| $\mathrm{~A}(12-17-16)$ | 120.4 | $\mathrm{~A}(7-18-17)$ | 127.6 |  |  | $\mathrm{~A}(32-19-33)$ | 108.4 |
| $\mathrm{~A}(12-17-29)$ | 119.8 | $\mathrm{~A}(7-18-31)$ | 117.7 |  |  | $\mathrm{~A}(32-19-34)$ | 108.2 |
| $\mathrm{~A}(14-13-26)$ | 119.6 | $\mathrm{~A}(9-8-26)$ | 119.8 |  |  | $\mathrm{~A}(33-19-34)$ | 108.1 |
|  |  |  |  |  |  | $\mathrm{~A}(35-20-36)$ | 108.2 |
|  |  |  |  |  |  | $\mathrm{~A}(35-20-37)$ | 108.3 |
|  |  |  |  |  |  | $\mathrm{~A}(36-20-37)$ | 108.1 |
|  |  |  |  |  |  |  |  |

## Physical properties

Depending on the Semi-empirical method of calculation according to the (PM3) is calculate some physical properties of the molecules studied in this research; Dipole moments ( $\mu$ in Debye) , energies(e V)of the high Occupied Molecular Orbital ( $\mathrm{E}_{\text {номо }}$ ) and the Lower Unoccupied Molecular Orbital ( $\mathrm{E}_{\mathrm{LUMO}}$ ) IP (ionization energy ). The higher HOMO energy values show the molecule is a good electron donor, in other hand, the lower HOMO energy values indicate that, a weaker ability of the molecules for donating electron. LUMO energy presents the ability of a molecule for receiving electron ${ }^{(17-19}$ ) . the negative $\mathrm{E}_{\text {Номо }}$ is equal to the ionization potential) the calculation has been ionization energies (e V), Also calculated the energy difference ( $\Delta \mathrm{E}$, e V), And finally calculated (Molecular Hardness)Hardness $(\eta)=1 / 2\left(E_{\text {номо }}-E_{\text {Lumо }}\right)$, (Electron Affinity) $\mathrm{E}_{\mathrm{A}}=-\mathrm{E}_{\text {LUMO }}$ according Koopmans theorem for N system of electrons ${ }^{(20)}$. The HOMO and LUMO of a molecule play important roles in intermolecular interactions, through the interaction between the HOMO of the drug with the LUMO of the receptor and vice versa. The interactions stabilized inversely with energy gap between the interacting orbitals. Increasing HOMO energy and decreasing LUMO energy in the drug molecule lead to enhancement stabilizing interactions, and hence, binding with the receptor enhancement stabilizing interactions, and hence, binding with the receptor ${ }^{(20,21)}$. ). The HOMO and LUMO of a molecule play important roles in intermolecular interactions, through the interaction between the HOMO of the drug with the LUMO of the receptor and vice versa. The interactions stabilized inversely with energy gap between the interacting orbitals ${ }^{(22,23)}$.

The results showed that compound 1 [ N '-(4-aminobenzylidene)benzohydrazide ] has high value for each of ( Dipole moment, IP, Electron Affinity $\mathrm{E}_{\mathrm{A}}$ ), And less value ( $\mathrm{E}_{\text {LUMo }}$ ), this means that this compound has more ability to loss electrons and easier ionization compared to compound 2 [ N '-(4-(dimethylamino benzylidene) benzohydrazide]. Also the results showed compound 2 has higher hardness $(\boldsymbol{\eta})$, lower energy gap $(\mathbf{\Delta E})$ ( Table 2), which may be explained the highest antiinflammatory compound (Table 2 )

Table 2:Calculated, $\mu$ (in Debye), orbital energies ( $\mathrm{E}_{\text {Hомо }}, \mathrm{E}_{\mathrm{LUOM}}, \Delta \mathrm{E}$ in eV ), $\operatorname{IP}\left(\mathrm{in} \mathrm{eV}\right.$ ), $\mathrm{E}_{\mathrm{A}}$ (in
$\mathrm{eV})$, and $\eta(\mathrm{in} \mathrm{eV})$ for the derivatives of bnzohydrazide.

| Comp. | $\mu$ Debye | $\mathrm{E}_{\text {Номо }}$ | $\mathrm{E}_{\text {Lомо }}$ | $\Delta \mathrm{E}$ | IP | $\mathrm{E}_{\mathrm{A}}$ | $\eta$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{~N}_{3} \mathrm{O}(1)$ | 3.3475 | -8.8563 | -0.3282 | 8.5281 | 8.8563 | 0.3282 | -4.26405 |
| $\mathrm{C}_{16} \mathrm{H}_{17} \mathrm{~N}_{3} \mathrm{O}(2)$ | 5.8520 | -8.6582 | -0.2166 | 8.4416 | 8.6582 | 0.2166 | -4.2208 |

## Thermodynamics functions

Thermodynamics functions for the studied molecules were listed in Table 3. Compared with $\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{~N}_{3} \mathrm{O}$ (1) $\mathrm{C}_{16} \mathrm{H}_{17} \mathrm{~N}_{3} \mathrm{O}$ (2), molecules along with the rotational constants, obtained in this study, Both $\mathrm{NH}_{2}, \mathrm{~N}\left(\mathrm{CH}_{3}\right)_{2}$ substituent's in same compound causes an increase in the all thermodynamics functions $\left[\mathrm{E}^{0}, \mathrm{H}^{0}, \mathrm{G}^{\mathrm{o}}, \mathrm{A}^{\mathrm{o}}, \mathrm{C}_{\mathrm{V}}, \mathrm{C}_{\mathrm{P}}, \mathrm{S}^{\mathrm{o}}\right]$ due to its stabilization by resonance effect.
$\mathrm{N}\left(\mathrm{CH}_{3}\right)_{2}>\mathrm{NH}_{2}$
The deferent thermodynamics functions values due to deferent substituted groups.Future studies must aim to computationally dock these conformers to the active site of the adrenergic receptor in order to determine which conformer has the greatest biological effect.where used to calculate the vibration and rotation contributions to the thermodynamic functions according to the statistical thermodynamic equations

$$
\begin{align*}
& U_{v i b}^{0}=\sum_{i=1}^{3 N-6} \frac{R T X_{i}}{e^{X_{i}}-1} \\
& X_{i}=\frac{1.4 \bar{\nu}}{T}  \tag{2}\\
& S_{v i b}^{0}=R \sum_{i=1}^{3 N-6}\left[\frac{X_{i}}{e^{X_{i}}-1}-\ln \left(1-e^{-X_{i}}\right)\right] \\
& S_{\text {rot }}^{0}=R\left[\frac{3}{2}+\ln \frac{8 \pi^{2}\left(8 \pi^{2} I_{X} I_{Y} I_{Z}\right)^{\frac{1}{2}}(K T)^{\frac{3}{2}}}{\sigma \hbar^{3}}\right]  \tag{4}\\
& \text { I : Moment of inertia } \\
& \text { Symmetry Coefficient } \sigma \\
& \mathrm{h}: \text { Planck's constant } \quad \mathrm{N}_{\mathrm{o}} \text { : Avocado's number } \\
& \mathrm{K} \text { : Boltzmann constant } \quad \mathrm{R} \text { : gas constant }
\end{align*}
$$

Table 3: The calculated standard thermodynamics functions at 298.15 oK of the derivatives of bnzohydrazide.

| Comp. | $\mathrm{E}^{0}$ <br> $\mathrm{~kJ} / \mathrm{Mol}$ | $\mathrm{H}^{0}$ <br> $\mathrm{~kJ} / \mathrm{mol}$ | $\mathrm{G}^{0}$ <br> $\mathrm{~kJ} / \mathrm{mol}$ | $\mathrm{S}^{0} \mathrm{~kJ} / \mathrm{mol} . \mathrm{deg}$ | $\mathrm{A}^{0} \mathrm{~kJ} / \mathrm{mol}$ | CV <br> $\mathrm{kJ} / \mathrm{mol} . \mathrm{deg}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Compound (1) <br> $C_{14} H_{13} N_{3} \mathrm{O}$ | 683.954 | 686.431 | 525.511 | 0.5408 | 364.5915 | 0.2507 |
| Compound $(2)$ <br> $C_{16} H_{17} N_{3} O$ | 832.431 | 834.909 | 647.467 | 0.6290 | 460.0250 | 0.2955 |

Frequencies IR Intensity of Absorption .
Table 4: Frequencies of Absorption of IR Intensity of Compound (1) $\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{~N}_{3} \mathrm{O}$ and Compound (2) $\mathrm{C}_{16} \mathrm{H}_{17} \mathrm{~N}_{3} \mathrm{O}$

| Type of <br> bond | Compound 1 <br> $C_{14} H_{13} N_{3} \mathrm{O}$ |  | Compound <br> $C_{16} H_{17} N_{3} O$ |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Frequency cm |  |  |  |
|  | 1869.6667 | IR intensity | Frequency <br> $\mathrm{cm}^{-1}$ | IR intensity |
| C=O <br> St. | 1945.8476 | 439.8104 | 1856.2778 | 115.6890 |
| C-H <br> St. | 3002.1298 | 15.1314 | 1954.1520 | 400.8234 |
| N-H <br> St. | 3368.1215 | 13.3575 | 3016.3847 | 12.3456 |

# Journal University of Kerbala , Vol. 14 No. 3 Scientific . 2016 

## References

[1] Allen, F.H., Kennard O.,Watson D.G., Brammer L., Orpen A.G.,Taylor R., Tables of bond lengths determined by X-ray and Neutron Diffraction. Part 1. Bond lengths in Organic compounds. J. Chem. Soc. Perkin Trans. II 1987,pp. S1-S19.
[2] Bedia K K, Elçin O, Seda U, Fatma K, Nathaly S, Sevim R \& Dimoglo A, Synthesis and characterization of novel hydrazide-hydrazones and the study of their structure-antituberculosis activity , Eur J Med Chem. ,41(11), pp.1253-1261, 2006.
[3] Titov A P, Grekov V I, Rybachenko V V, Shevchenko, Correlation of IR parameters with reactivity characteristics for the hydrazides of carboxylic acids, Theo. Exp. Chem., , 4, 6, pp 476-481, 1971.
[4] John D. K., Derrik L. W., Julianne M.L., Andrei S. and Charles F. B. , Characterization of New and Some Known Organic Compounds: Phenyl Carbazate and its Transformation into Hydrazones, Chem. Educator, 14, pp. 1-6, 2009.
[5] Byrkit G.D., Michalek G.A., Hydrazine in Organic Chemistry, Ind. Eng. Chem., 42 (9), pp. 1862-1875, 1950.
[6] CampodÓnico . P. R. , Aliaga M. E. , José G. Santos J. G., Reactivity of benzohydrazide derivatives towards acetylation reaction.Experimental and theoretical studies, Chem. Phys. Lett, , 488 ,pp. 8689, 2010.
[7] Castro E.A ., Bessolo J., Aguayo R., Santos J.G., Kinetic investigation of the reactions of S-4nitrophenyl 4-substituted thiobenzoates with secondary alicyclic amines in aqueous ethanol . J. Org Chem ., 68(21) :pp.8157-8161, 2003.
[8] Dharmesh R. D. and MukeshC. P., Synthesis, characterization and biological activity of novel pyrole compounds, Rasayan J. Chem., 4, 3, pp.580-584, 2011.
[9] Maheta S. , Patel S.J., Synthesis and biological activity of 4-chloro-2-hydroxy-N-(5-methylene-4-oxo-2-aryl-thiazolidin-3-yl) benzamide, Bulg. Chem. Commun., 43, 3, pp. 411-418 2011.
[10] James J. P. S., Optimization of parameters for semiempirical methods , J. Comp. Chem.,10, pp. 209 216, 1989.
[11] Stewart. J.J .P, Csaszar, P., Pulay, P. "Fast Semiempirical Calculations, J. Comp. Chem . , 4, 3, pp. 227228, 1982.
[12] Bingham R C, Dewar M J S and Lo H C The Chemistry of Heterocyclic Compounds, Thiophene and Its Derivatives .J. Am.Chem.Soc., 97 (6) , pp. 1285 -1293, 1975.
[13 ] Michael J. S. Dewar, Walter T. , Ground states of molecules. 38. The MNDO method. Approximations and parameters J. Am. Chem. Soc., 99 ,pp. 4899-4907, 1977.
[14 ] Frisch M. J., Trucks G. W, Schlegel H. B., Scuseria G. E.,. Robb M. A.. Cheeseman J. R, Scalmani G., Barone V., Mennucci B., Petersson G. A., Gaussian 09, Revision E.01, 2009.
[15] Young, D. 'Computational Chemistry: A Practical Guide for Applying Techniques to Real World Problems" ; John Wiley \& Sons, inc: USA, 2001.
[16 ] Jensen, F. 'Introduction to computational Chemistry", 2nd ed.; John Wiley \&Sons, inc: England, 2007
[17] Ogretir C, Mihci B, Bereket G. Quantum chemical studies of some pyridine derivatives as corrosion inhibitors. J.Mol.Struct: Theochem.; 488:pp.223-231, 1999.
[18] Bereket G, Ogretir C, Ozsahin C. Quantum chemical studies on the inhibition efficiencies of some piperazine derivatives for the corrosion of steel in acidic medium. J Mol Struct: Theochem ;663, PP. 39-46, 2003.
[19] Obot IB.,Obi-Egbedi NO.HSAB descriptors of thiadiazole derivatives calculated by DFT: possible relationship as mild steel corrosion inhibitors.Der Pharma Chemica.1, pp.106-123, 2009.
[20] Fukui K., "Role of frontier orbitals in chemical reactions," Sci. J., 218, pp. 747-754, 1982.
[21] Chattaraj, P. K.; and Parr, R. G.; "Principle of maximum hardness," J. Am. Chem. Soc., 113,pp. 18541855, 1991.
[22] Zhan C. G., J. A. Nicholsand ,D. A. Dixon, J. Phys. Chem A, 107,PP. 4184-4195, 2003.
[23] Amado A.M., Sónia M. Fiuza, Luis A. E. Batista de Carvalho, On the Effects of Changing Gaussian Program Version and SCRF Defining Parameters: Isopropylamine as a Case Study, Bull. Chem. Soc. Jpn. J. 85, 9, pp. 962-975, 2012.

