

# EFFECT OF THE ELECTRONIC STRUCTURE OF PARA-SUBSTITUTED BENZALDEHYDE BENZOHYDRAZONE ON ITS ANTIMICROBIAL ACTIVITY: A DFT ANALYSIS

## ABSTRACT

*Bacillus subtilis* a bacterium that has demonstrated its efficacy across various domains, including industry, agriculture, and commerce, owing to its protective, inhibitory, and biological mechanisms against specific microbes. However, at high concentrations, it can lead to food poisoning and severe infections, resulting in symptoms such as diarrhea and vomiting. Bacterial spores produced by *Bacillus subtilis* can induce conditions like gas gangrene and tetanus. In this context, benzohydrazones are recognized for their antimicrobial activity, particularly against *Bacillus subtilis*. This study aims to elucidate the relationship between the electronic structure of para-substituted benzaldehydebenzohydrazone derivatives and their antimicrobial activity. This leads to the proposal of a 2D pharmacophore for predicting the antibacterial activity of these derivatives. The quantitative structure-activity relationship (QSAR) approach employed is the KPG method. The electronic structures were optimized using the density functional theory (DFT) method with the B3LYP functional and the 6-31G (d,p) basis set. Charge and local molecular orbitals were considered in the optimization process. The resulting prediction equation ( $R=98.95\%$ ,  $R^2=97.91\%$ , Adjusted  $R^2=96.76\%$ ,  $F(5,9)=84.52$ ) derived from multiple linear regression provides the basis for the proposed 2D pharmacophore. This equation shows that antimicrobial activity of benzohydrazone derivative is on charge and orbital controlled. This pharmacophore holds potential utility in designing new molecular structures with enhanced activity against *Bacillus subtilis*.

Keywords: *Bacillus subtilis*, DFT, Hydrazone, KPG method, QSAR.

## 1. INTRODUCTION

*Bacillus subtilis*, commonly known as hay bacillus or grass bacillus, is a Gram-positive, catalase-positive bacterium. Not only is it the most extensively studied Gram-positive bacterium, but it also serves as a model organism for investigating bacterial chromosome replication and cell differentiation. Widely distributed in soil and the gastrointestinal tracts of ruminants and humans, *Bacillus subtilis* stands out as a champion bacterium in the production of secreted enzymes employed on an industrial scale by biotechnology companies [1–3]. Certain strains of *Bacillus subtilis* protect plants from fungal plant pathogens by producing an impressive array of antibiotics, including non-ribosomal lipopeptides (LPs) [4]. This bacterium holds considerable significance in agriculture, trade, and industry, being utilized for diverse purposes such as seed treatment, crop protection, turf foliation, enhancement of peanut seed germination, promotion of nodulation and root growth, and plant nutrition [5, 6]. There is potential for its development as a biological control agent against R.

solani in greenhouse cucumber and tomato crops [7]. Several *Bacillus* species are believed to possess the ability to degrade environmental pollutants, exemplified by *Bacillus subtilis* capability to degrade certain pesticides like carbendazim [8, 9] and chlorpyrifos [10]. Additionally, *Bacillus subtilis* contributes to an increase in the production of gibberellic acids (GAs) and indole acetic acid (IAA), fostering plant growth and enhancing the synthesis of defense molecules such as superoxide dismutase, peroxidase, and polyphenol oxidase [11]. Furthermore, *Bacillus* exhibits inhibitory effects on pathogens through the production of antibiotic lipopeptides [12, 13]. Despite its numerous benefits, *Bacillus subtilis* has adverse effects and poses a moderate potential danger to humans. It has been implicated in the etiology of food toxic-infections, with high concentrations leading to diarrhea and/or vomiting [14]. The production of extracellular enzymes and toxins by *Bacillus subtilis* contributes to its pathogenicity in humans. Moreover, resistance to several antimicrobial drugs presents a challenge for its treatment [15].

In the laboratory, the chemistry of nitrogenous compounds has long been a subject of extensive study [16]. Hydrazones, a class of nitrogenous compounds, are particularly valuable in drug design due to the presence of an azomethine proton  $-NHN=CH-$  in their structure [17]. The physical, chemical, and biological properties that allow the understanding and prediction of the activity or behavior of molecules in the environment are embedded in the structure of these compounds. Molecular modeling, especially QSAR/QSPR (Quantitative Structure-Activity/Property Relationships), is a widely used technique for predicting the properties/activities of chemical systems from their molecular structures [18][18, 19]. Over the decades, QSAR has played a pivotal role in drug development, encouraging scientists in the pharmaceutical field to explore relationships between molecular parameters and properties beyond activity [20]. To comprehend a mechanism of action, researchers can examine the relationships between the descriptors of the QSAR model and toxicity or other activities [21]. The assumption that the molecular structure of a series of compounds contains essential information about the factors responsible for their physical, chemical, or biological properties has led chemoinformaticians to extensively employ QSAR/QSPR to study the biological activity of various compounds [19, 22–27]. Presently, one of the greatest challenges in QSAR/QSPR studies is assessing the reliability of the mechanistic interpretation of the identified relationships [28].

Given the adverse effects of *Bacillus subtilis* on humans, its resistance to antimicrobials, the pharmacological properties of hydrazones, and the significance of QSAR, there is an urgent need to identify hydrazone molecules that can more effectively control *Bacillus subtilis*. Therefore, the overarching objective of our project is to conduct QSAR on a series of hydrazone molecules and propose pharmacophores that can act more effectively on *Bacillus subtilis*,

with the aim of combating diarrhea, vomiting, and food poisoning caused by this pathogen.

## 2. MODEL, METHODS AND CALCULATIONS

### 2.1 Model

$$\log(C_{MIC}) = a + bM_{D_i} + c \log \left[ \sigma_{D_i} / (ABC)^{1/2} \right] + \sum_j \left[ e_j Q_j + f_j S_j^E + s_j S_j^N \right] + \sum_j \sum_m \left[ h_j(m) F_j(m) + x_j(m) S_j^E(m) \right] + \sum_j \sum_{m'} \left[ r_j(m') F_j(m') + t_j(m') S_j^N(m') \right] + \sum_j \left[ g_j \mu_j + k_j \eta_j + o_j \omega_j + z_j \zeta_j + w_j Q_j^{\max} \right] \quad (\text{Eq.1})$$

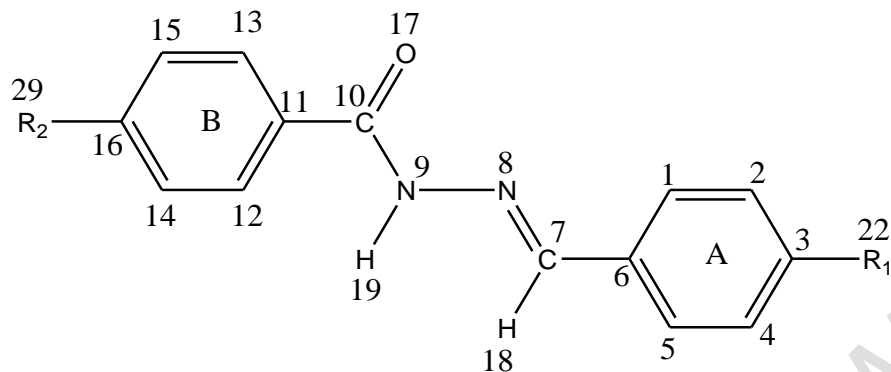
with  $a$ ,  $b$ ,  $c$ ,  $e_j$ ,  $f_j$ ,  $s_j$ ,  $h_j(m)$ ,  $x_j(m)$ ,  $r_j(m')$ ,  $t_j(m')$ ,  $g_j$ ,  $k_j$ ,  $o_j$ ,  $z_j$  and  $w_j$  are constants,  $M_{D_i}$  is the mass of the drug,  $\sigma_{D_i}$  is its symmetry number,  $ABC$  is the product of the moments of inertia of the drug around the three main axes of rotation,  $Q_j$  is the net charge of the atom  $j$ ,  $S_j^E$  and  $S_j^N$  are, respectively, the total electrophilic and nucleophilic atomic superdelocalizations of atom  $j$ ,  $F_j(m)$  and  $F_j(m')$  are respectively the electron populations (Fukui index) of the occupied ( $m$ ) and vacant ( $m'$ ) molecular orbitals (OMs) located on the atom  $j$ ,  $S_j^E(m)$  is the atomic electrophilic superdelocalizability of the OM ( $m$ ) localized on the atom  $j$ ,  $\mu_j$  is the local electronic chemical potential of the atom  $j$ ,  $\eta_j$  is the local atomic hardness of the atom  $j$ ,  $\omega_j$  is the local atomic electrophilicity of atom  $j$ ,  $\zeta_j$  is the local atomic softness of the atom  $j$ , and  $Q_j^{\max}$  is the maximum amount of electronic charge that atom  $j$  can accept from another site [29, 30].

### 2.2 Methods

To find the relationship between the electronic structure and the inhibitory concentration of the series of para-substituted benzaldehydebenzohydrazone derivatives, the used methodology widely explained in the articles [31–41]. The results obtained were presented following a routine methodology. So this article contains only the results and discussion because the method and calculations have been discussed in several articles [31–41].

### 2.3 Selection of the molecules

The para-substituted benzaldehydebenzohydrazone derivatives used are from reference [42]. The general formula and the antibacterial activities (mean inhibitory concentration: CMIC) of these molecules are represented respectively in Figure 1 and Table 1.



**Fig. 1.** General structure of para-substituted benzaldehydebzohydrazone derivatives

**Table 1.** Molecules of para-substituted benzaldehydebzohydrazone derivatives with the decimal logarithm of their average inhibitory concentrations [ $\log(C_{MIC})$ ].

N°	R <sub>1</sub>	R <sub>2</sub>	$\log(C_{MIC})$
A <sub>1</sub>	-H	-H	-5.35
A <sub>2</sub>	-H	-CH <sub>3</sub>	-5.37
A <sub>3</sub>	-H	-OCH <sub>3</sub>	-5.40
A <sub>4</sub>	-H	-Cl	-5.41
A <sub>5</sub>	-H	-OH	-5.38
A <sub>6</sub>	-OCH <sub>3</sub>	-H	-5.40
A <sub>7</sub>	-OCH <sub>3</sub>	-CH <sub>3</sub>	-5.42
A <sub>8</sub>	-OCH <sub>3</sub>	-OCH <sub>3</sub>	-5.45
A <sub>9</sub>	-OCH <sub>3</sub>	-Cl	-5.45
A <sub>10</sub>	-OCH <sub>3</sub>	-OH	-5.41
A <sub>11</sub>	-NO <sub>2</sub>	-H	-5.42
A <sub>12</sub>	-NO <sub>2</sub>	-CH <sub>3</sub>	-5.45
A <sub>13</sub>	-NO <sub>2</sub>	-OCH <sub>3</sub>	-5.47
A <sub>14</sub>	-NO <sub>2</sub>	-Cl	-5.48
A <sub>15</sub>	-NO <sub>2</sub>	-OH	-5.45

We observed that the amplitude is low. This could be a limitation of this study. To provide a comprehensive example of how to implement the KPG method, a study with a higher amplitude should be conducted.

## 2.4 Calculations

The Gaussian program was employed for the geometrical optimizations [43] of the fifteen structures of para-substituted benzaldehydebzohydrazone derivatives using the density functional theory (DFT) and the 6-31G(d,p)/B3LYP basis. Regarding the D-CENT QSAR program [44, 45] it was utilized to calculate the local atomic reactivity indices for the various atoms in the shared skeleton of the distinct para-substituted benzaldehydebzohydrazone molecules. To exclude atoms with weak coefficients, the Statistica 10 program was applied, enabling the execution of multiple linear regression [46]. The common skeleton

of the fifteen antibacterial molecules under investigation is illustrated in Figure 2, including the numbering of its different atoms.

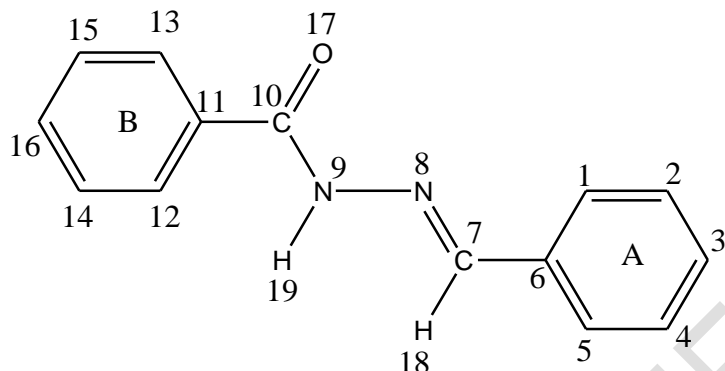


Fig. 2. Common skeleton of para-substituted benzaldehydebenzohydrazone derivatives.

### 3. RESULTS AND DISCUSSION

#### 3.1 Results

After multiple linear regression (MLR), the following model was obtained:

$$\log(C_{MIC}) = 2.17 + 6.13Q_5 - 12.0Q_{10} + 0.38Q_{13}^{max} + 0.69F_{19}(HOMO) * 0.27Q_{14} \text{ (Eq.2)}$$

With  $R = 98.95\%$ ;  $R^2 = 97.91\%$ ; Adjusted  $R^2 = 96.76\%$ ;  $F(5.9) = 84.517$ ; ( $p = .0000003$ ) and  $SD = 0.007$ . No outliers were detected and no residuals outside the 2 limit.

According to this prediction model, the activity of para-substituted benzaldehydebenzohydrazone derivatives on *Bacillus subtilis* potentially depends on five (05) local atomic reactivity indices:  $Q_5$  is the net charge of the  $C_5$  carbon atom of the aromatic ring A,  $Q_{10}$  is the net charge of the carbon atom  $C_{10}$  of the carbonyl group,  $Q_{13}^{max}$  is the maximum value of electronic charge that atom  $C_{13}$  of the aromatic ring B may receive,  $F_{19}(HOMO)^*$  is the Fukui index of the first highest occupied molecular orbital (HOMO)\* located on the hydrogen atom  $H_{19}$ ,  $Q_{14}$  is the net charge of the carbon atom  $C_{14}$  of the aromatic ring B.

Tables 2 and 3 group respectively the beta coefficients and the correlation matrix between the five explanatory variables of the model.

Table 2. Statistical parameters of the different indices of the regression model

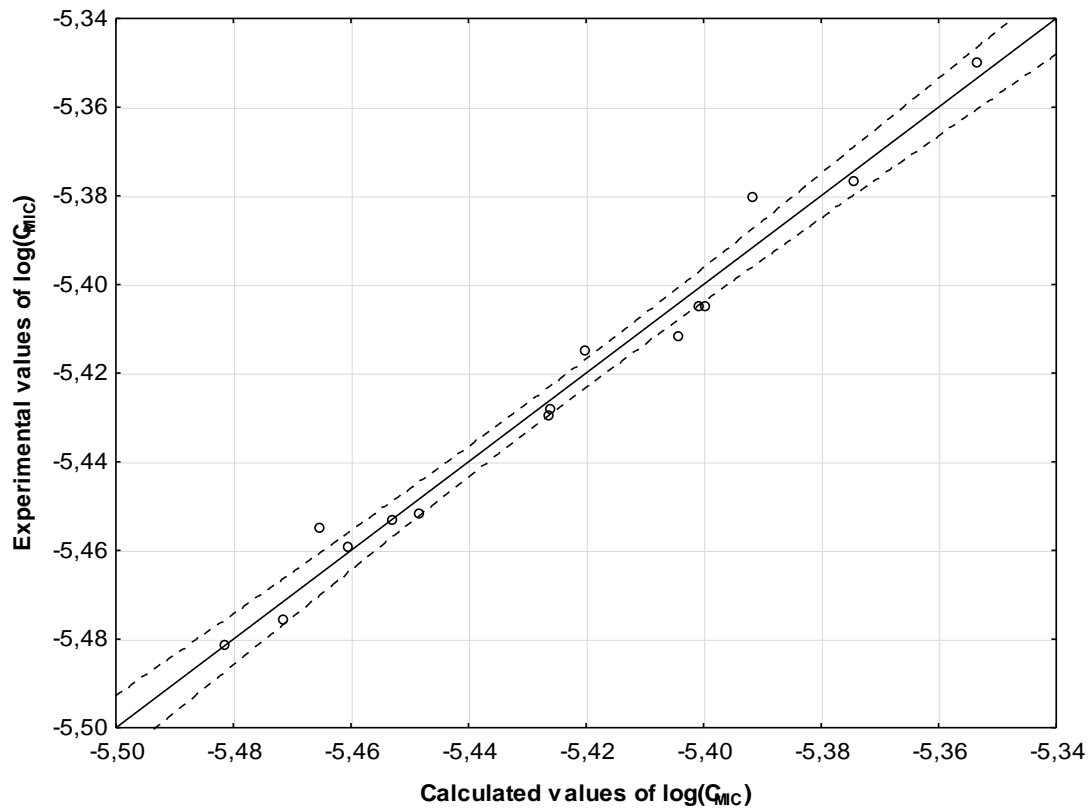
Variables	p-value	Beta coefficients
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$Q_5$	0.0000001	1.041
$Q_{10}$	0.000002	-0.695
$Q_{13}^{max}$	.002	0.277
$F_{19}(\text{HOMO})^*$	.02	0.139
$Q_{14}$	.05	0.139

**Table 3. Correlation matrix of the different variables of the model**

	$Q_5$	$Q_{10}$	$Q_{13}^{max}$	$F_{19}(\text{HOMO})^*$	$Q_{14}$
$Q_5$	1.00				
$Q_{10}$	0,058	1.00			
$Q_{13}^{max}$	0,056	0,210	1.00		
$F_{19}(\text{HOMO})^*$	0,012	0,069	0,005	1.00	
$Q_{14}$	0,0001	0,228	0,256	0,002	1.00

Figure 3 shows that there is a good correlation between observed and calculated values because almost all points are inside the 95% confidence interval.



**Fig. 3. Prediction plot of the experimental values of  $\log(C_{MIC})$  against the values estimated from Eq. 1. The dashed lines indicate the 95% confidence interval.**

Table 4 shows the local MO structure of atoms with reactivity indices appearing in Eq. 1 (see Fig. 2). Nomenclature: Molecule (HOMO)\* / (HOMO-2)\* (HOMO-1)\* (HOMO)\* - (LUMO)\* (LUMO+1)\* (LUMO+2)\*.

**Table 4. The local molecular orbitals of atoms 5, 10, 13, 19 and 14 of the developed model**

Molecules	Atom (C <sub>5</sub> )	Atom (C <sub>10</sub> )	Atom (C <sub>13</sub> )	Atom (H <sub>19</sub> )	Atom (C <sub>14</sub> )
A <sub>1</sub> (59)	54π55π5π9- 60π61π62π	57σ58π59π - 60π61σ64π	52π57π58- 60π61π62π	45σ53σ54σ- 62σ63σ65σ	56π57π58π- 60π61π62π
A <sub>2</sub> (63)	58π59π63π- 64π65π66π	61π62π63π- 64σ65π68π	60π61π62- 64π65π67π	48σ57σ58- 66σ67σ69σ	60π61π62π- 64π65π67π
A <sub>3</sub> (67)	63π66π67- 68π69π70π	62π65π66π - 68π69π72π	65π66π67- 68π69π70π	51σ61σ62- 69σ70σ71σ	65π66π67π- 69π70π71π
A <sub>4</sub> (67)	62π64π67π- 68π69π70π	65σ66π67π- 68σ69π72π	63π65π66- 68π69π70π	51σ61σ62- 70σ74σ75σ	63π65π66π- 68π69π70π
A <sub>5</sub> (63)	59π62π63π- 64π65π66π	61π62σ63- 64π65σ68π	61π62π63- 64π65π66π	49σ57σ58- 65σ66σ69σ	61π62π63π- 66π67π68π
A <sub>6</sub> (67)	62π65π67π- 68π69π70π	64σ66π 67σ- 68π69σ72σ	63π64π66- 68π69π70π	39σ52σ61- 69σ70σ71σ	60π64π66π- 68π69π70π
A <sub>7</sub> (71)	65π66π71π- 72π74π76π	69σ70σ71π- 72π73σ74π	68π69π70- 72π73π74π	41σ55σ65- 73σ74σ75σ	68π69π70π- 72π75π76π
A <sub>8</sub> (75)	70π72π75π- 76π77π78π	73σ74σ75π- 76σ77π78π	73π74π75- 76π78π79π	43σ58σ69- 77σ79σ81σ	73π74π75π- 78π79π80π
A <sub>9</sub> (75)	71π73π75π- 76π77π78π	73σ74π75π- 76π77σ81π	71π72π74- 76π77π78π	57σ58σ69- 77σ78σ80σ	72π73π74π- 76π78π79π
A <sub>10</sub> (71)	66π68π71π- 72π73π74π	69σ70σ71π- 72π 73π74σ	69π70π71- 72π74π75π	40σ56σ65- 73σ77σ78σ	68π69π70π- 73π75π76π
A <sub>11</sub> (70)	65π66π70π- 71π72π73π	68σ69π70π- 71σ72π74π	67π68π69- 72π74π75π	54σ65σ66- 73σ75σ78σ	67π68π69π- 72π75π76π
A <sub>12</sub> (74)	69π70π74π- 75π76π77π	72σ73π74π- 75σ76σ77π	72π73π74- 76π78π79π	57σ69σ70- 77σ79σ81σ	71π72π73π- 76π79π80π
A <sub>13</sub> (78)	74π77π78π- 79π80π81π	71σ76π77π- 79π80σ81σ	76π77π78- 80π81π82π	60σ73σ74- 81σ83σ85σ	76π77π78π- 83π84π85π
A <sub>14</sub> (78)	73π74π78π- 79π80π81π	76π77σ78σ- 79σ80π82π	76π77π78- 80π83π85π	71σ73σ74- 81σ82σ83σ	75π76π77π- 80π83π84π
A <sub>15</sub> (74)	70π73π74π- 75π76π77π	67σ72π 73π- 75π76σ77π	72π73π74- 76π77π78π	58σ69σ70- 77σ78σ79σ	72π73π74π- 78π79π80π

### 3.2 Discussion

#### *Global statistical parameters of the developed model*

- Obtaining an adjusted R<sup>2</sup> coefficient = 96.76% proves that the antimicrobial activity of para-substituted benzaldehydebenzohydrazone molecules on *Bacillus subtilis* is strongly correlated with the five local atomic reactivity indices[47].

- Obtaining a Fischer probability  $F(5.9) = 84.517 > \text{tabulated } F(5.9) = 3.48$ , shows that the developed model is statistically significant and that there is a 95% chance that there is a real relationship between the antimicrobial activity of the para-substituted benzaldehydebenzohydrazone molecules and the local atomic reactivity indices[47].
- The p-value obtained ( $p=.0000003 < .05$ ) also confirms that the developed multiple linear regression model is statistically significant.

#### *Statistical parameters of the five indices of the established model*

The values of the beta coefficients show that the most important index in the antimicrobial activity of *Bacillus Subtilis* is the net charge  $Q_5$  of atom 5 and the order of priority of the different indices in the model is as follows:  $Q_5 > Q_{10} > Q_{13}^{max} > F_{19}(HOMO) > Q_{14}$

The negative value of the beta coefficient of  $Q_{10}$  variable shows that there is an inverse relationship between the antimicrobial activity and the variable.

All p-values for the five local atomic reactivity indices are less than 5%. Therefore, all these indices are statistically significant and can be used for variable-by-variable analysis.

#### *Correlation between the different indices of the multiple linear regression model*

The analysis of the values in this table of correlation shows that there is a low correlation (practically less than 50%) between the different indices. This proves the interdependence between these indices in predicting the antimicrobial activity of para-substituted benzaldehydebenzohydrazone molecules.

#### *Orbital analysis*

All the implicated orbital of atoms C(5), C(13) and C(14) are  $\pi$ -type because they belong to aromatic fragment. But for atom C(10) it is about a mixture of  $\sigma$  and  $\pi$ -type. It is also confirm the  $\sigma$ -type of hydrogen orbitals.

#### *Variable-by-variable analysis of the prediction model indices:*

The lower  $C_{MIC}$ , the better the antimicrobial activity. Therefore, better antimicrobial activity of para-substituted benzaldehydebenzohydrazone derivatives would be associated with low value of negative net charges  $Q_5$  and  $Q_{14}$  with positive coefficients, low positive value of  $Q_{13}^{max}$  with a positive coefficient, to a low value of the Fukui index  $F_{19}(HOMO)^*$  positive with a positive coefficient and a high value of the net load  $Q_{10}$  positive with a negative coefficient.

Atoms 5, 10, and 14 are carbon atoms. However, the net charges of atoms  $C_5$  and  $C_{14}$  are negative, while the net charge of atom  $C_{10}$  is positive. This discrepancy can be attributed to the fact that atoms 5 and 14 are carbons

belonging to aromatic rings, thus rich in electrons. In contrast, atom 5 is a carbon doubly bound to an oxygen atom, depleting it in electrons. The net charge  $Q_{14}$  of the atom is influenced by the nature of the substituent  $R_2$  attached to the  $C_{16}$  carbon of the aromatic ring B (fig. 2). Analyses suggest that both atom 5 and atom 14 could contribute to antimicrobial activity through cation-anion or cation- $\pi$  interactions, including the  $\pi$ - $\pi$  interactions involving the  $C_{14}$ - $C_{16}$  and  $C_5$ - $C_6$  atoms. On the other hand, the  $C_{10}$  carbon atom would establish anion-cation or anion- $\pi$  interactions.

Atom 13 is a carbon atom. low value of  $Q_{13}^{max}$  suggest that this atom is not prone to receive extra charge. Therefore atom 13 is probably facing an electron-deficient center. Atom 13 must interact with an electron acceptor. This interaction may be of the  $\pi$ -alkyl or alkyl-alkyl type.

Atom 19 is a hydrogen atom bound to the nitrogen atom  $N_9$ . The local molecular orbitals (MOs) of this atom are all sigma ( $\sigma$ ) in nature. For better inhibitory activity of the compounds a low positive value of  $F_{19}(\text{HOMO})^*$  is necessary. For this, the orbital of the  $H_{19}$  atom must interact with an electron-deficient center through its highest occupied sigma local molecular orbital.

Based on these analyses, a 2D pharmacophore was proposed in figure 4, indicating the sites where substituents should be attached to enhance the antimicrobial activity of the compounds in general and their inhibitory activity on *Bacillus subtilis* in particular.

**Fig.4. Pharmacophore 2D pour l'inhibition de *Bacillus subtilis***

#### **4. CONCLUSION**

The utilization of the Klopman-Peradejordi-Gómez (KPG) method has allowed us to establish a prediction model for the antimicrobial activity of para-substituted benzaldehydebenzohydrazone derivatives. This predictive model demonstrates a 98.95% likelihood of a correlation between the antimicrobial activity of these para-substituted benzaldehydebenzohydrazone molecules and the five potential local atomic reactivity indices. The pharmacophore, derived from a variable-by-variable analysis of this model, highlights specific atoms in the common backbone of para-substituted benzaldehydebenzohydrazone derivatives where bioisosteres could be strategically grafted to enhance their antimicrobial activity. Leveraging this 2D pharmacophore, we can propose potential structures for antimicrobially active derivatives of para-substituted benzaldehydebenzohydrazone.

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