

DETERMINATION OF DESCRIPTORS WHICH INFLUENCE THE TOXICITY OF ORGANOCHLORINE COMPOUNDS USING QSAR METHOD

SUMMARY

Organochlorine Pesticides (OCP) are organic compounds obtained by the chlorination of various unsaturated hydrocarbons. They are very toxic and therefore belong to the family of persistent organic pollutants. If formerly these pesticides were used to improve the productivity and fight against certain vectors of diseases, today they are considered as "enemy" of the environment. To understand the origin of the toxicity of organochlorine compounds, we used 73 molecules (test set: 50 and validation: 23) containing at least one chlorine atom and for which the toxicity (LogLC50) against *Poecilia reticulata* is known to establish QSAR models. Firstly, we used principal component analysis (PCA) to identify the best descriptors. Then, the different models are established using the method of multiple linear regression (MLR). Models established with quantum and physicochemical descriptors only showed satisfactory results. But the best model is determined with the combination of both quantum and physicochemical descriptors. The criteria of this model are as follows:

$$R^2 = 0.939 ; R_{adjusted}^2 = 0.932 ; P_{value} < 0.0001 ; \alpha = 0.05$$

$$R_{CV}^2 = 0.935 ; R^2 - R_{CV}^2 = 0.004 ; MCE = 0.073 ; F = 134.701$$

These criteria show that the toxicity of organochlorine compounds is well described by the combination of quantum and physicochemical descriptors namely lipophilia (LogP), polarizability (pol), entropy (S), zero-point energy (ZPE) and the number of chlorine atoms (NCl).

Key words: organochlorine compounds, toxicity, QSAR, quantum descriptors and physicochemical descriptors

1- INTRODUCTION

Organochlorine pesticides use began in the 1940s with the advent of dichlorodiphenyl- trichloroethane (DDT) which was synthesized for the first time by Othmar Zeidler in 1873 [1] and whose insecticidal properties were discovered by Paul Müller in 1939 [2]. These pesticides have contributed to improve and increase agricultural yields and have led to progress in the control of food resources [3]. The use of organochlorine pesticides has not been limited to increasing agricultural yields. it has also spread to other sectors. Indeed, organochlorine pesticides have long time been used to fight against certain diseases vectors [4-7]. DDT is one of the insecticides recommended by WHO for indoor residual spraying for malaria control [8]. After two decades of intense use, research [7, 9] has begun to show the dangerousness of these chemical compounds to the environment.

Over time, several studies [10-13] will prove their presence in all ecosystems and elements which live there. The presence of organochlorine pesticides in all ecosystems is assumed to be due to both their persistence their volatility.

The persistence of organochlorine compounds in the environment is largely due to the stability of the carbon-chlorine bond that is resistant to degradation. But their presence throughout the food chain is the consequence of

41 their lipo-solubility. This property gives them the ability to cross the phospholipid structure of biological
42 membranes thereby reaching the adipose tissue in which they accumulate. The rate of these compounds increases
43 in the body of the species that live in these environments when we progress in the food chain in which humans
44 are at the top [14-16].

45 As for volatility, it is generally due to the relatively high vapor pressure. This property allows them to travel
46 great distances. Thus, according to studies [17-19], organochlorine pesticides have been found in environments
47 in which these pesticides have never been used and sometimes even far from the places where they were used. In
48 view of all the above-mentioned, the use of organochlorine pesticides will therefore be regressed from the 1970s
49 and even several will be banned in some countries. But it was until the year 2001 that the first Conference on
50 Persistent Organic Pollutants (POPs) was held. Indeed, in Sweden in 2001, the Stockholm Convention marked
51 the first convention on POPs. This convention has been signed by 151 countries. This high number of signatories
52 of the convention shows its importance. Since then, it was decided to reduce or even to eliminate the production
53 and the exploitation of persistent organic pollutants. Among the 12 organic pollutants covered by this
54 convention, 9 represent organochlorine pesticides.

55 The main objective of this study is to perform a QSAR study of organochlorine compounds to determine the
56 descriptors that influence their toxicity.

57 Crum-Brown and Frazer [20] were considered to be the precursors of the QSAR methodology. Indeed, already in
58 1868, they postulated that the biological activity of a molecule is a function of its chemical constitution. In 1893,
59 Richet [21] discovered that the toxicity of organic
60 compounds is inversely proportional to their solubility in water. But the era of QSAR really begins in the 1960s
61 with, on the one hand the publication of Hansch and Fujita [22] and on the other hand, the publication of Free
62 and Wilson [23]. And from then, the number of publications containing the word "QSAR" continues to increase
63 due its ability to predict the properties of chemical compounds.

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65 **2- MATERIALS AND METHODS**

66 **2.1- Materials**

67 **a- Selection of data set**

68 We investigated about 73 molecules as displayed in Table 2. These molecules were taken from Alan Katritzky et
69 al.'s article [24]. For each molecule, LogLC_{50} is calculated where LC_{50} stands for the concentration that causes
70 the death of 50% of the population of the test. Besides, the most toxic compound is assumed to display the
71 smallest value of LogLC_{50} . Furthermore, it should be noted that lethal dose (LD) and lethal concentration (LC)
72 are identical. But the difference is related to the mode of penetration of the substance into the body. When the
73 administered substance enters the body by inhalation, the notion of dose is replaced by that of concentration.
74 Thus, the lethal dose 50 becomes the lethal concentration 50 [25]. According to Hodge and Sterner [26]
75 chemicals products can be classified into 6 groups according to their toxicities.

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82 Table 1: Class of toxicity according to the scale Hodge and Sterner [27]

Index or class of toxicity	Term commonly used	Toxicological parameter (DL ₅₀)
1	Extremely toxic	DL ₅₀ ≤ 1 mg/kg
2	Highly toxic	1 mg/kg ≤ DL ₅₀ ≤ 50 mg/kg
3	Moderately toxic	50 mg/kg ≤ DL ₅₀ ≤ 500mg/kg
4	Slightly toxic	500 mg/kg ≤ DL ₅₀ ≤ 5000mg/kg
5	almost not toxic	5000 mg/kg ≤ DL ₅₀ ≤ 15000 mg/kg
6	relatively harmless	DL ₅₀ ≥ 15000 mg/kg

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84 The set of all 73 molecules are registered in Table 2.

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UNDER PEER REVIEW

87 Table 2 : Names and LogLC50 of 73 organochlorine molecules

N°	Compounds	Log LC50			
Test set					
1	3-chloroaniline	2.02	38	2-chlorophenol	1.94
2	3-chlorophenol	1.70	39	3,4-dichloroaniline	1.61
3	4,5-dichloro-2-methoxyphenol	1.40	40	3,4-dichlorotoluène	1.40
4	4-chloro-3,5-dimethylphenol	1.34	41	3,5-dichloroaniline	1.38
5	4-chloroaniline	2.33	42	3-chlorotoluène	2.16
6	4-chlorophenol	1.82	43	chlorobenzene	2.23
7	4-chlorotoluène	1.67	44	lindane	-0.69
8	Chloroform	2.93	45	trichloroethene	2.58
9	dichloromethane	3.54	46	1,2,3-trichlorobenzene	1.11
10	hexachlorobutadiene	-0.20	47	1,3-dichlorobenzene	1.72
11	pentachlorobenzene	-0.15	48	2,4-dichlorophenol	1.41
12	pentachloroethane	1.74	49	2,5-dichlorophenol	1.42
13	pentachlorophenol	0.22	50	3,4,5,6-tetrachloro-2-hydroxyphenol	1.00
14	tetrachloroethene	1.98	Validation set		
15	tetrachloromethane	2.64	51	1,1-dichloroéthane	3.31
16	α,α -dichloro-m-xylene	-0.16	52	1,2,3,4-tetrachlorobenzene	0.65
17	1,1,1-trichloroéthane	3.00	53	1,2,3,5-tetrachlorobenzene	0.57
18	1,1,2-trichloroéthane	2.82	54	1,2,4,5-tetrachlorobenzene	0.15
19	1,2-dichlorobenzene	1.60	55	1,2,4-trichlorobenzene	1.17
20	2,3,4,5-tetrachloroaniline	0.19	56	1,2-dichloroethane	3.06
21	2,3,5-trichlorophenol	1.08	57	1,2-dichloropropane	3.01
22	2,4-dichloroacetophenone	1.80	58	1,3-dichloropropane	2.87
23	2,4-dichloroaniline	1.59	59	1,4-dichlorobenzene	1.44
24	2-chloro-4-methylphenol	2.40	60	2,2,2-trichloroethanol	3.31
25	Dieldrin	-1.78	61	2,3,4,6-tétrachlorophenol	0.67
26	hexachloroethane	0.81	62	2,3,6-trichloroaniline	1.27
27	1,1,2,2-tetrachloroethane	2.23	63	2,4,5-trichlorophenol	0.80
28	1,2,3-trichloropropane	2.45	64	2,4,6-trichlorophenol	1.06
29	1,3,5-trichlorobenzene	1.26	65	2,4, α -trichlorotoluène	0.08
30	1-chlorobutane	3.02	66	2,4-dichlorotoluène	1.46
31	2,3,4,5-tetrachlorophenol	0.48	67	2,6-dichlorophenol	1.68
32	2,3,5,6-tetrachloroaniline	0.07	68	2-chloroaniline	1.69
33	2,3,5,6-tetrachlorophenol	0.74	69	3,4,5-trichloro-2,6-dimethoxyphenol	1.12
34	2,3,6-trichlorophenol	1.44	70	3,4,5-trichloro-2-methoxyphenol	1.03
35	2,4,5-trichloroaniline	1.08	71	3,4,5-trichlorophenol	0.92
36	2,4,5-trichlorotoluène	0.94	72	3,5-dichlorophenol	1.22
37	2,5-dichloroaniline	1.01	73	4-chloro-3-methylphenol	1.67

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91 **b- Molecular descriptors**

92 We have calculated several quantum and physicochemical descriptors to carry out the QSAR model.

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94 ➤ **Quantum descriptors**

95 For quantum descriptors we determined highest occupied molecular orbital energy (E_{HOMO}), lowest unoccupied
96 molecular orbital energy (E_{LUMO}), total energy (E_T), dipole moment (DM), constant volume heat capacity (C_v),
97 entropy (s), thermal energy (E_{th}), highest Mulliken electronic charge (CAE), lowest Mulliken electronic charge
98 (CAF), ionization potential (PI) ; electronic affinity (AE), energy gap between E_{HOMO} and E_{LUMO} (ΔE), absolute
99 electronegativity (χ), Chemical potential (μ), absolute hardness (η), mollesse (\S) and reactivity index (ω). we
100 also considered the number of chlorine (N_{Cl}) as a descriptor. These descriptors were generated or calculated by
101 Gaussian 03 [27]. The calculations were performed thanks to DFT method with B3LYP as the functional and 6-
102 311++g(d,p) as basis set. Moreover, they were determined on reference to equations below.

103 $PI = -E_{HOMO}; \quad AE = -E_{LUMO}; \quad \Delta E = E_{LUMO} - E_{HOMO}$
104 $\chi = \frac{E_{LUMO} + E_{HOMO}}{2} = \frac{AE + PI}{2} = -\mu; \quad \eta = \frac{E_{LUMO} - E_{HOMO}}{2} = \frac{PI - AE}{2}; \quad \S = \frac{1}{\eta}; \quad \omega = \frac{\mu^2}{2\eta}$

105
106 ➤ **Physicochemical descriptors**

107 For physico-chemical descriptors, we used ChemSketch [28] to determine lipophilicity (LogP), formula weight
108 (M), molar refractivity (Rm), molar volume (Vm), parachlor (Pc), index of refraction (Ir), surface tension (γ),
109 density (d) and polarizability (Pol).

110 **2.2- Methods**

111 **a- Descriptive analysis**

112 XLSTAT [29] software and the principal component analysis (PCA) method were used to realize the matrix of
113 correlation. Principal component analysis (PCA) permits to examine descriptors set and to select the good ones
114 that give the best model at prediction [30]. It allows to identify the descriptors which correlate well with the
115 biological activity Log (LC50).

116
117 **b- Statistical analysis**

118 The establishment of QSAR model consists of making out mathematical relationship between biological activity
119 and chemical descriptors. Thus, QSAR model is considered satisfied when the following conditions are satisfied.
120 The choice of the best statistical model has to satisfy the following criteria that encompass the highest coefficient
121 of determination (R^2), the highest of adjusted determination coefficient ($R^2_{adjusted}$), the highest coefficient of
122 Fischer (F), the highest coefficient of cross validation (R^2_{CV}), the lowest values of Mean Square Error (MSE) and
123 the difference $R^2 - R^2_{CV} < 0.3$.
124 The calculation of these parameters requires a statistical analysis. The most available methods allowing that
125 calculation are Simple Linear Regression (SLR), Multiple Linear Regression (MLR), neurons networks and
126 Partial Least Squares (PLS). In this article, MLR method of XLSTAT is used to perform the prediction [29].

127 3- RESULTS AND DISCUSSIONS

128 3.1- Matrix of correlation

129 The principal component analysis permits to perform the matrix of correlation. Any descriptor having a partial
130 correlation coefficient with the biological activity Log (LC50) less than 0.5 is removed from the descriptors set.
131 For 2 descriptors having partial correlation coefficients with biological activity greater than 0.5 and whose
132 partial correlation coefficient between both descriptors is greater than 0.95 then the one with the smallest partial
133 correlation coefficient with the biological activity is also removed from the descriptors set. And we obtain the
134 matrix of correlation presented by Tables 3 and 4. These tables show that the selected descriptors correlate well
135 with toxicity. Because all of correlation's coefficients are higher than 0.5.

136

137 Table 3: Matrix of correlation (Pearson (n)) of quantum descriptors used for model 1

Variables	Log LC50	E _{LUMO}	S	ZPE	N _{Cl}	ΔE
Log LC50	1					
E _{LUMO}	0,522	1				
S	-0,848	-0,425	1			
ZPE	0,52	0,668	-0,542	1		
N _{Cl}	-0,569	-0,772	0,536	-0,9	1	
ΔE	0,68	0,336	-0,581	0,182	-0,091	1

138

139 Table 4 : Matrix of correlation (Pearson (n)) of Physico-chemical descriptors used for model 2

Variables	Log LC50	LogP	M	Pc	Ir	γ	d	Pol
Log LC50	1							
LogP	-0,847	1						
M	-0,871	0,826	1					
Pc	-0,777	0,668	0,779	1				
Ir	-0,728	0,622	0,628	0,576	1			
γ	-0,679	0,59	0,682	0,55	0,941	1		
D	-0,574	0,603	0,797	0,413	0,527	0,656	1	
Pol	-0,925	0,832	0,912	0,853	0,774	0,749	0,552	1

140

141 3.2- Quantum descriptors model

142 > Equation of model 1

143 $Log LC50 = 3.11 - 12.66 * E_{LUMO} - 0.049 * S - 4.92E-04 * ZPE - 0.51 * NCl + 13.05 * \Delta E$

144 $N = 50 ; R^2 = 0.88 ; R^2_{adjusted} = 0.86 ; R^2_{CV} = 0.86 ; R^2 - R^2_{CV} = 0.02 ;$

145 $MCE = 0.15 ; F = 63.40 ; P_{value} < 0.0001 ; \alpha = 0.05$

146 Here, N is the number of molecule, MCE is Mean Square Error, R^2 is the coefficient of determination,

147 $R^2_{adjusted}$ is the adjusted coefficient of determination, R^2_{CV} is the coefficient of cross validation and F is the

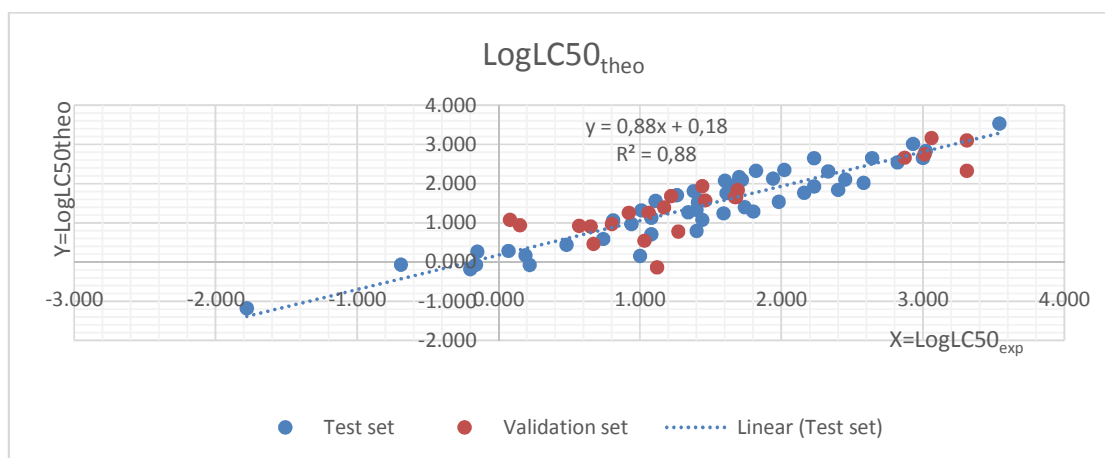
148 coefficient of Fischer.

149 Furthermore, the small value of P_{value} means that the selected variables do bring a significant amount of
 150 information. The high values of R^2 (0.88) and $R_{adjusted}^2$ (0.86) and the low value of MSE (0.15) including
 151 $0.5 < R_{CV}^2 < 0.9$ and $R^2 - R_{CV}^2 < 0.3$ indicate that the proposed model is satisfactory and can predict our
 152 biological activity (LC50).

153 $R_{adjusted}^2 = 0.86$ indicates that the toxicity of 86% of our compounds are described with reliability by the
 154 selected descriptors. The high value of the coefficient of Fischer ($F = 63.40$) shows the strong relation which
 155 exists between the toxicity and the selected descriptors.

156 We can notice from the analysis of the equation of model 1 that Log (LC50) increases when energy of the lowest
 157 unoccupied molecular orbital (E_{LUMO}), the entropy (s), the zero-point energy (ZPE) and the number of chlorine
 158 atoms (N_{Cl}) decrease while the gap energy (ΔE) increases. Thus, the toxicity of organochlorine compounds
 159 evolves in the same direction as ΔE and in the opposite direction as E_{LUMO} , S , ZPE and N_{Cl} .

160 The regression line between the experimental and theoretical LogLC50 of the test and the validation set is
 161 illustrated in Figure 1. The high value of determination coefficient ($R^2 = 0.88$) and the low value of mean
 162 square error $MCE = 0.15$ prove a good similarity between the predicted and experimental values. This good
 163 similarity is highlighted also into Figure 2.

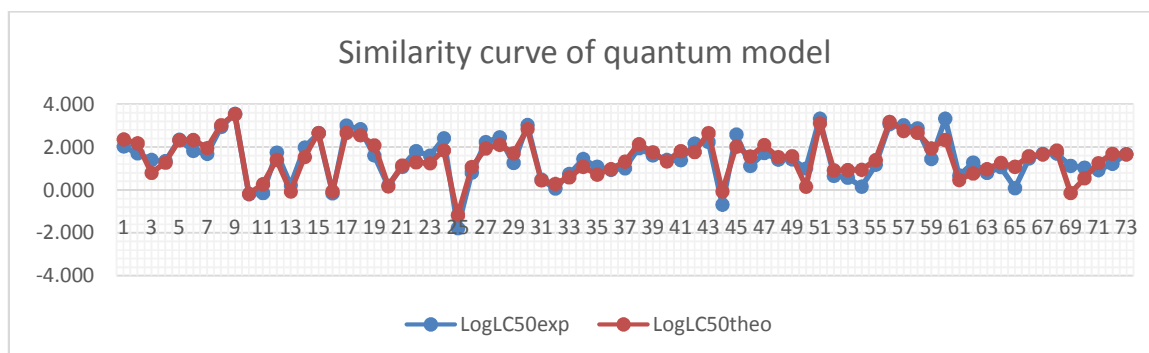


164

165 Figure 1: Regression line of the test and validation set

166 The regression line of the test set indicates that: $LogLC_{50theo} = 0.88 * LogLC_{50exp} + 0.18$

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168

169 Figure 2: Similarity curve of the experimental and predicted values of the quantum descriptors model

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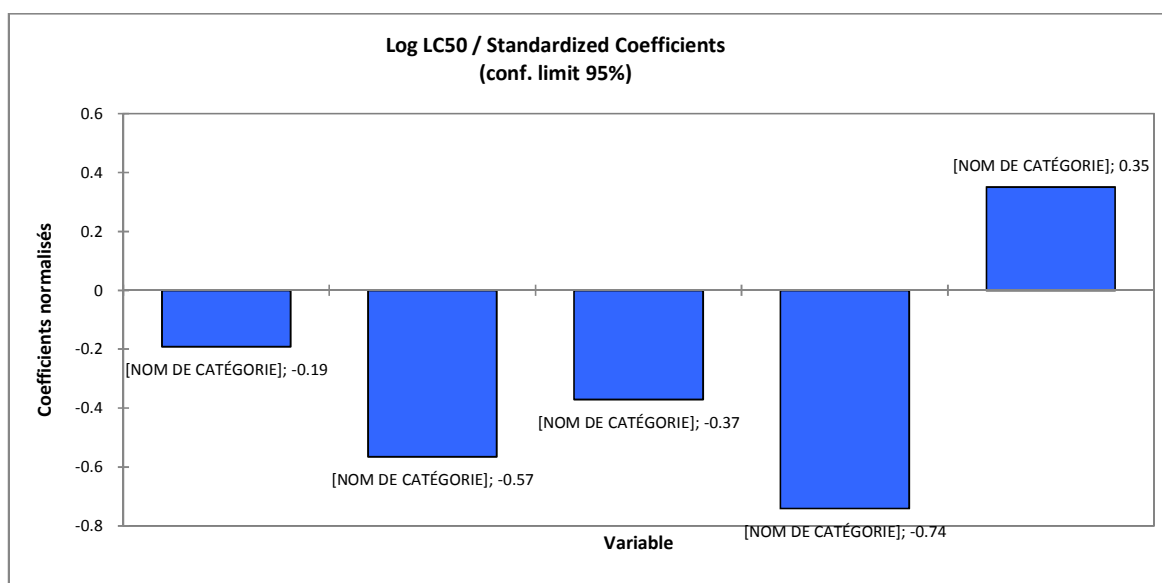
171 ➤ Contribution of descriptors

172 The contributions of the quantum descriptors in the prediction of the organochlorine
173 molecules toxicity were illustrated by the Figure 3. The classification of the contribution of
174 the descriptors in the model is as follows:

$$175 \quad N_{Cl}(-0.74) > S(-0.57) > ZPE(-0.37) > \Delta E(0.35) > ELUMO(-0.19)$$

176 According to the contribution of these descriptors, the number of chlorine molecules (N_{Cl}) is
177 the most important descriptor and lowest unoccupied molecular orbital energy (E_{LUMO}) is the
178 least important descriptor for the prediction of toxicity of organochlorine molecules.

179



180

181 Figure 3: Contribution of descriptors in model 1

182 3.3- Physico-chemical descriptors model

183 ➤ Equation of model 2

$$184 \quad \text{Log LC50} = 9,95 - 0,11 * \text{LogP} - 3,31 * \text{Ir} + 0,03 * \gamma - 0,73 * d - 0,22 * \text{pol}; \quad N = 50; R^2 = 0.90; MCE = 0.12;$$

$$185 \quad F = 80.29; P_{value} < 0.0001; \alpha = 0.05 \quad R_{adjusted}^2 = 0.89; R_{cv}^2 = 0.89; R^2 - R_{cv}^2 = 0.01$$

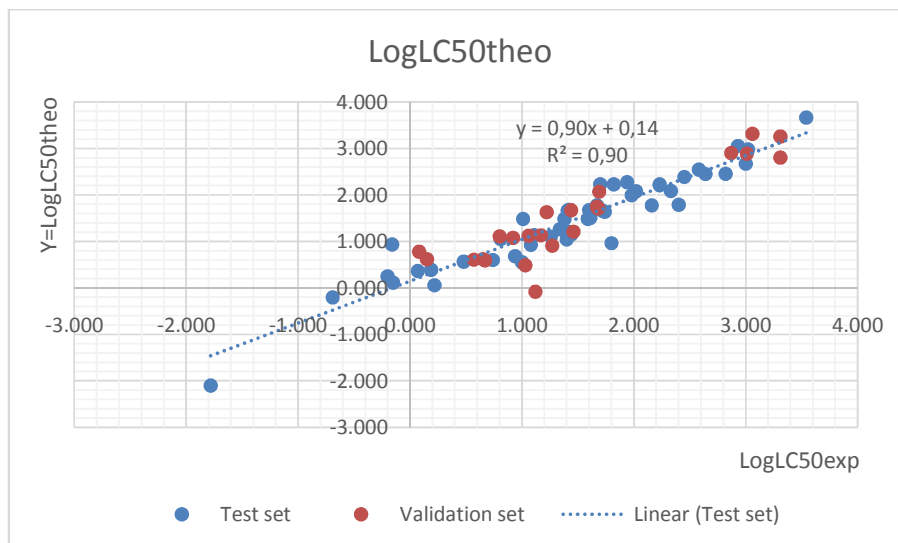
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187 The various calculated parameters prove that the model established with the physicochemical descriptors is
188 predictive and reliable. In this model 89% of our compounds are described with reliability by the selected
189 descriptors. Therefore, this model is assumed to be satisfactory.

190 The toxicity of organochlorine compounds increases when lipophilicity (LogP), density (d), polarisability (pol)
191 and index of refraction decrease and surface tension (γ) increases.

192 The regression line between the experimental and theoretical LogLC50 of the test and the validation set is
193 illustrated in Figure 4. Here, the high value of determination coefficient ($R^2 = 0.90$) and the low value of mean

194 square error $MCE = 0.12$ prove a good similarity between the predicted and experimental values. This good
 195 similarity is emphasized also into Figure 5.
 196

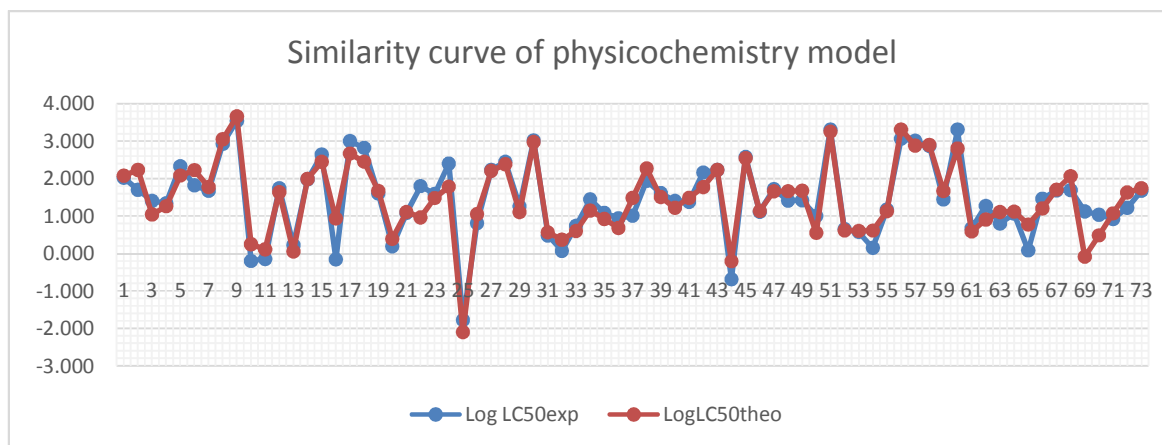


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198 Figure 4: Regression line of the test and validation set

199 The regression line of the test set indicates that: $LogLC_{50theo} = 0.90 * LogLC_{50exp} + 0.14$

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202 Figure 5: Similarity curve of the experimental and predicted values of the physicochemical descriptors model

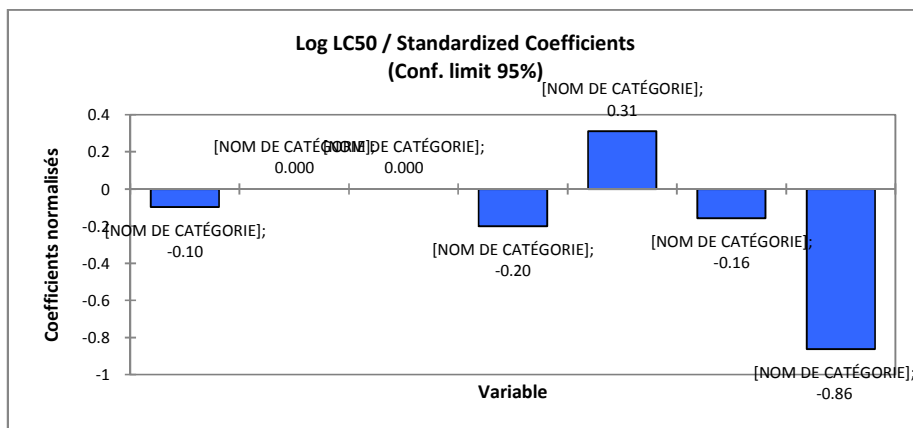
203

204 ➤ Contribution of descriptors

205 The contributions of the physicochemical descriptors in the prediction of the organochlorine molecules toxicity
 206 were illustrated by the Figure 6. The classification of the contribution of the descriptors in the model is as
 207 follows:

208 $Pol(-0.86) > \gamma(0.31) > Ir(-0.20) > d(-0.16) > LogP(-0.10)$

209 According to this classification, polarizability is the most important descriptor and lipophilicity is the least
 210 important descriptor to prediction of toxicity of organochlorine molecules.



211

212 Figure 6: Contribution of descriptors in model 2

213 3.4- Quantum and Physicochemical descriptors model

214 ➤ Equation of model

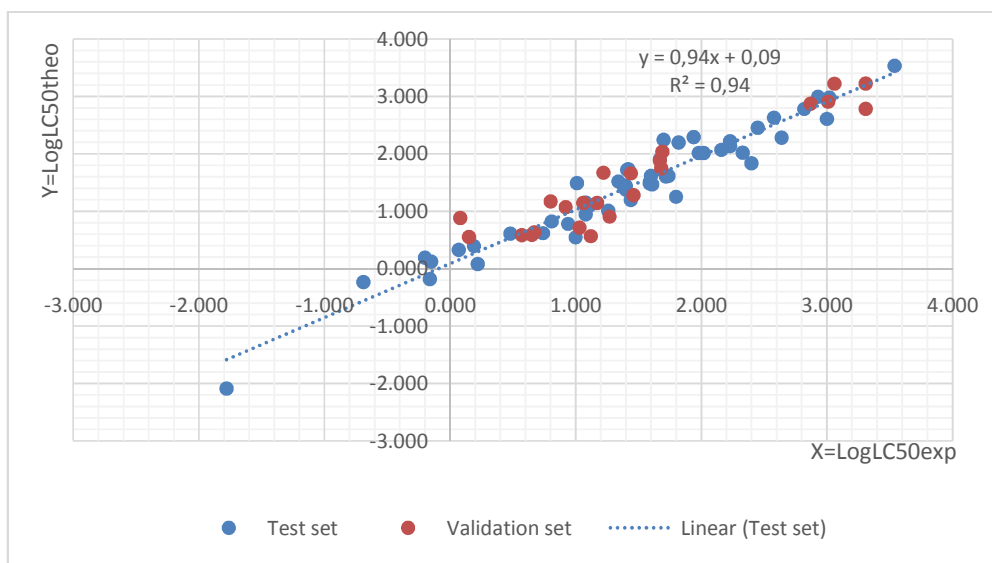
215 $\text{Log LC50} = 3,72 - 0,14*\text{LogP} - 0,29*\text{pol} + 3,17\text{E-}02*\text{S} - 5,70\text{E-}04*\text{ZPE} - 0,36*\text{N}_{\text{Cl}}$

216 $R^2 = 0,94; R^2_{\text{adjusted}} = 0,93; R^2_{\text{CV}} = 0,93; R^2 - R^2_{\text{CV}} = 0,004; MCE = 0,07; F = 134,70; P_{\text{value}} < 0,0001;$

217 $\alpha = 0,05$

218 The greatest values of R^2 ; R^2_{adjusted} and F and the lowest values of MCE and P_{value} shows the strong relation
 219 which exists between the toxicity and the selected descriptors. In this model 93% of our compounds are
 220 described with reliability by the selected descriptors. Then $R^2_{\text{CV}} (= 0,93) > 0,9$ and $R^2 - R^2_{\text{CV}} (= 0,004) <$
 221 $0,3$ prove that this model is excellent. This equation shows that LogLC50 of organochlorine molecules increases
 222 when LogP, Pol, ZPE and NCl decrease and S increases.

223 The regression line between the experimental and theoretical LogLC50 of the test and the validation set is
 224 illustrated in Figure 7. The high value of determination coefficient ($R^2 = 0,94$) and the low value of mean
 225 square error $MCE = 0,073$ prove a good similarity between the predicted and experimental values. This good
 226 similarity is demonstrated also through Figure 8.

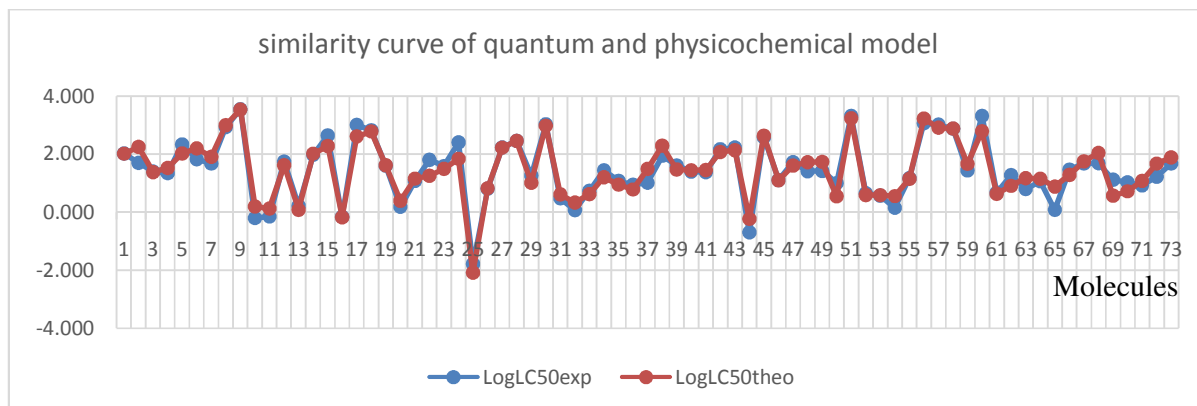


227

228 Figure 7: Regression line of the test and validation set

229 The regression line of the test set indicates that: $LogLC_{50theo} = 0.94 * LogLC_{50exp} + 0.09$

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231

232 Figure 8: Similarity curve of the experimental and predicted values of the quantum and physicochemical
233 descriptors model

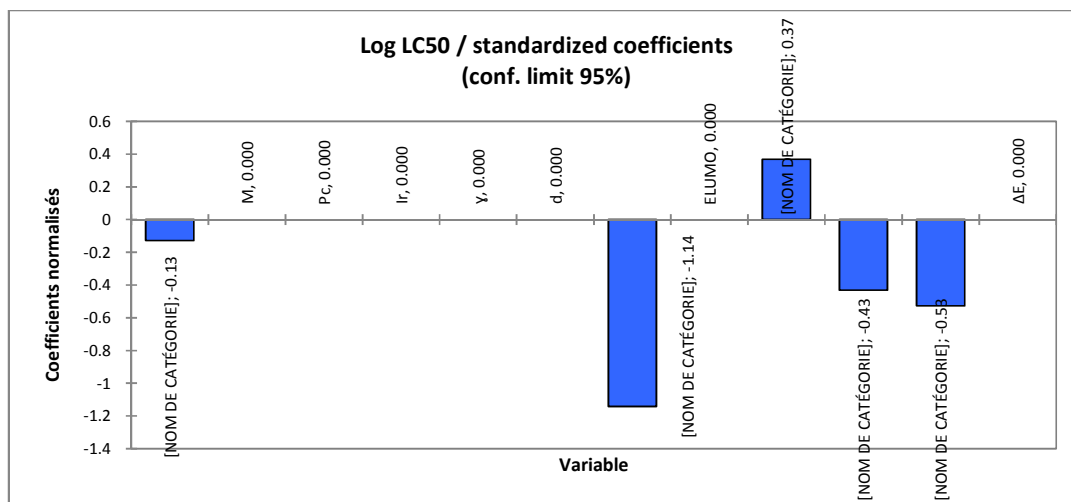
234

235 > Contribution of descriptors

236 The contributions of the five descriptors in the prediction of the organochlorine molecules toxicity are illustrated
237 by the Figure 9. The classification of the contribution of the descriptors in the model is as follows:

238 $Pol(1.14) > N_{cl}(0.53) > ZPE(0.43) > S(0.37) > LogP(0.13)$

239 According to the contribution of these descriptors, polarizability is the most important descriptor and
240 lipophilicity is the least important descriptor for prediction of toxicity of organochlorine molecules.



241

242 Figure 9: Contribution of descriptors in model

243

244 3.5- Comparison of different models

245 The comparison of the criteria of validation enables us to choose the best model for the toxicity of the
246 organochlorine molecules prediction. These criteria are summarized in the Table 5.

Nature of descriptors	Quantum	Physicochemical	Quantum and Physicochemical
Criteria			
<i>N</i>	50	50	50
<i>R</i>²	0.88	0.90	0.94
<i>R</i>²_{adjusted}	0.86	0.89	0.93
<i>MCE</i>	0.15	0.12	0.07
<i>F</i>	63.40	80.29	134.70
<i>R</i>²_{CV}	0.86	0.89	0.93
<i>R</i>² - <i>R</i>²_{CV}	0.02	0.01	0.004
<i>P</i> value	< 0.0001	< 0.0001	< 0.0001
<i>α</i>	0.05	0.05	0.05

249

250 After comparing the validation criteria contained in Table 5, we notice that the best model is that obtained with
 251 the union of both quantum and physicochemical descriptors. This is proven by the highest value of
 252 *R*², *R*²_{adjusted}, *F* and *R*²_{CV} and the lowest values of *MCE* and *R*² - *R*²_{CV}. The values of *R*²_{CV} superior to 0,9 and
 253 *R*² - *R*²_{CV} lower than 0,3 show that the established model
 254 is excellent [31]. Besides, The table below summarizes the different molecules, the experimental values of
 255 toxicity (Log (LC50))_{exp}, the theoretical values of toxicity (Log (LC50))_{theo} and the ratio (Log (LC50))_{exp} / (Log
 256 (LC50))_{theo}.

257 Table 6 : Ratio (Log (LC50))_{exp} / (Log (LC50))_{theo} of mixed model

N°	Compounds	Log(LC50 _{exp})	Log(LC50 _{theo})	Log(LC50 _{exp}) / Log(LC50 _{theo})
Validation set				
51	1,1-dichloroéthane	3.31	3.225	1.026
52	1,2,3,4-tetrachlorobenzene	0.65	0.590	1.101
53	1,2,3,5-tetrachlorobenzene	0.57	0.583	0.978
54	1,2,4,5-tetrachlorobenzene	0.15	0.552	0.272
55	1,2,4-trichlorobenzene	1.17	1.145	1.022
56	1,2-dichloroéthane	3.06	3.221	0.950
57	1,2-dichloropropane	3.01	2.908	1.035
58	1,3-dichloropropane	2.87	2.875	0.998
59	1,4-dichlorobenzene	1.44	1.660	0.867
60	2,2,2-trichloroethanol	3.31	2.783	1.189
61	2,3,4,6-tétrachlorophenol	0.67	0.632	1.060
62	2,3,6-trichloroaniline	1.27	0.905	1.403
63	2,4,5-trichlorophenol	0.8	1.170	0.684
64	2,4,6-trichlorophenol	1.06	1.145	0.926
65	2,4,α-trichlorotoluène	0.08	0.882	0.091
66	2,4-dichlorotoluène	1.46	1.282	1.139
67	2,6-dichlorophenol	1.68	1.746	0.962
68	2-chloroaniline	1.69	2.038	0.829
69	3,4,5-trichloro-2,6-diméthoxyphenol	1.12	0.565	1.982

70	3,4,5-trichloro-2-methoxyphenol	1.03	0.717	1.437
71	3,4,5-trichlorophenol	0.92	1.073	0.858
72	3,5-dichlorophenol	1.22	1.670	0.730
73	4-chloro-3-methylphenol	1.67	1.884	0.887

258

259 The ratio $(\text{Log}(\text{LC50}))_{\text{exp}} / (\text{Log}(\text{LC50}))_{\text{theo}}$ varies around 1 for most compounds, which justifies that our model
260 is excellent.

261

262 **4- CONCLUSION**

263 The organochlorine pesticides constitute a subgroup of the organochlorine compounds. These compounds are
264 well-known for their toxicity. What led us to determine, by QSAR method, the theoretical descriptors which
265 could better explain this toxicity. Firstly, we determined on the one hand the quantum descriptors and on the
266 other hand the physicochemical descriptors. Then, a Principal component analysis enabled us to select the best
267 descriptors. Finally, three QSAR models were established. If the models established with the quantum
268 descriptors only and the physico-chemical descriptors only gave good results with respectively $R^2 =$

269 0.88 ; $R^2_{adjusted} = 0.86$; $R^2_{CV} = 0.86$; $R^2 - R^2_{CV} = 0.02$; $MCE = 0.15$; $F = 63.40$ and $R^2 = 0.90$;
270 $R^2_{adjusted} = 0.89$; $R^2_{CV} = 0.89$; $R^2 - R^2_{CV} = 0.01$; $MCE = 0.12$; $F = 80.29$, the best model was obtained with the
271 combination of the two types of descriptors whose equation and criteria of validation are:

272 $\text{Log LC50} = 3,72 - 0,14 * \text{LogP} - 0,29 * \text{pol} + 3,17E-02 * \text{S} - 5,70E-04 * \text{ZPE} - 0,36 * \text{N}_{Cl}$
273 $R^2 = 0.94$; $R^2_{adjusted} = 0.93$; $R^2_{CV} = 0.94$; $R^2 - R^2_{CV} = 0.004$; $MCE = 0.07$; $F = 134.70$; $P_{value} <$
274 0.0001 ; $\alpha = 0.05$.

275 In perspective, we will determine the descriptors that influence the half-life time and the bioaccumulation factor
276 of organochlorine compounds and then propose less toxic, less bio-accumulative and less persistent
277 organochlorine pesticides.

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