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# Environmental toxicity prediction using computational tools: prediction of potential hazardous effects of chemicals in *Lactuca sativa* seed germination

Predicción de la toxicidad ambiental usando herramientas computacionales: predicción de efectos tóxicos de químicos en la germinación de semillas de *Lactuca sativa* 

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#### **ABSTRACT**

The main aim of the study was to develop quantitative structure-activity relationship (QSAR) models for the prediction of phytotoxicity effects of chemical compounds on the *Lactuca sativa* seeds germination. A database of 73 compounds, assayed against *L. sativa* and Dragon's molecular descriptors are used to obtain a QSAR model for the prediction of the phytotoxicity. The model is carried out with QSARINS software and validated according to OECD principles. The best model showed good value for the determination coefficient ( $R^2 = 0.917$ ) and others parameters appropriate for fitting (s = 0.256 and RMSE<sub>tr</sub> = 0.236). The validation results confirmed that the model has good robustness and stability ( $Q^2_{LOO} = 0.874$  and  $Q^2_{LMO} = 0.875$ ), an excellent predictive power ( $R^2_{ext} = 0.896$ ) and was product of a non-random correlation ( $R^2_{Y-scr} = 0.130$  and  $Q^2_{Y-scr} = -0.265$ ). Finally, we can say that this model is a good predictor tool to predict the toxicity over *L. sativa* of chemical compounds.

**KEYWORDS:** ecotoxicity, Lactuca sativa, phytotoxicity, QSARINS software.

#### **RESUMEN**

El objetivo principal del estudio fue desarrollar modelos cuantitativos de relación estructuraactividad (QSAR) para la predicción de los efectos fitotóxicos de compuestos químicos, en la germinación de las semillas de *Lactuca sativa*. Se utiliza una base de datos de 73 compuestos, ensayados contra *L. sativa* y los descriptores moleculares del programa Dragon para obtener un modelo QSAR para la predicción de la fitotoxicidad. El modelo se lleva a cabo con el software QSARINS y se valida de acuerdo con los principios de la OCDE. El mejor modelo mostró buen valor para el coeficiente de determinación ( $R^2 = 0.917$ ) y otros parámetros apropiados para el ajuste (s = 0.256 and RMSE<sub>tr</sub>=0.236). Los resultados de la validación confirmaron que el modelo tiene una buena robustez y estabilidad ( $Q^2_{LOO} = 0.874$  and  $Q^2_{LMO} = 0.875$ ), un excelente poder predictivo ( $R^2_{ext} = 0.896$ ) y que no fue producto de una correlación casual ( $R^2_{Y-scr} = 0.130$  and  $Q^2_{Y-scr} = -0.265$ ). Finalmente, podemos decir que el modelo es una buena herramienta de predicción para predecir la toxicidad de compuestos químicos sobre *L. sativa*.

PALABRAS CLAVE: ecotoxicidad, fitotoxicidad, Lactuca sativa, programa QSARINS.

#### INTRODUCTION

As a consequence of progress in the chemical and pharmaceutical industries in the last decades, the quantity and the number of xenobiotics have considerably increased. The U.S. agencies, such as the Environmental Protection Agency (U.S. EPA) and the U. S. Food and Drug Administration (U.S. FDA), reported that in recent years people have been exposed to about 63000 chemicals, many of which show a high toxic activity [1]. The detrimental effects caused by these compounds act on both people's health and on the balance of natural and manipulated ecosystems [2].

The environment is regularly exposed to organic chemicals (e. g. phenols, anilines, benzenes derivatives, etc.) as well as mixtures of several chemical elements; through their use in industrial processes, chemical assays in research institutions and domestic use of different substances. Wastewaters usually contain a variety of organic and inorganic chemicals, and the joint actions of the individual components can lead to complicated integrated toxic effects to the environment [3]. Several ecological perturbations are associated with wastewater discharge into fluvial ecosystems, such as the introduction of micropollutants to stream flow or the loss of invertebrate biodiversity [4, 5]. Consequently, the development of tools able to assess potential hazardous effects of chemicals on living organisms needs to receive attention. Therefore, information about the toxicity of industrial organic chemicals to the environment is of interest [6].

The experimental tests provide most reliable data on the effects of chemicals, but they involve much time consumption and extensive resources, which makes it difficult to research great numbers of potential toxic compounds [7]. Therefore, quantitative structure-activity/toxicity relationships (QSAR/QSTR) studies provide an invaluable tool in the prediction of environmental toxicity directly from the molecular structure of compounds [8]. The QSAR/QSTR studies offer the advantages of higher speed and lower cost, especially when compared to experimental testing [9], and, in recent years, the predictions from computer models have been widely used in modern toxicological research, as they are an important alternative for obtaining experimental evidence and play an important role in evaluating the toxicity of chemicals [10].

Among the different endpoints used to evaluate the potential toxicological impact of chemical exposure to terrestrial organisms, *Lactuca sativa* was chosen because is a very used test species; due to its sensitivity to different organic compounds, its high-frequency use in phytotoxicological testing and its handiness [11, 12]. Also, lettuce (*L. sativa*) has agricultural importance and is one of the species recommended by the EPA, FDA and OECD (Organization for Economic Cooperation and Development) [11].



Furthermore, our research group have a large experience, of more than a decade, developing QSAR models to predict different physicochemical, chemical, pharmacokinetical, toxicological as well as biological properties of different compounds [13-19]. The main objective of the present study was to propose a new and better mathematical model for prediction of potential hazardous effects of chemicals in *L. sativa* seed germination process using QSAR INSubria software (QSARINS) [20-22]. This approach offers an alternative to conventional methods used to predict the toxicity by identifying the relationship between the chemical structure and its toxicity.

#### **MATERIALS AND METHOD**

# **Chemical Database and Descriptor Calculation**

The general dataset used in this study assay was obtained from previously published papers [23, 24]; it consists of 73 compounds, assayed against *L. sativa*. The dataset was divided in training set (56 compounds) to fit the model the external prediction set (18 compounds), also known as test set. The experimental values of the molecules of the entire dataset are shown in Table 1.

For this study, we employed Dragon software [25] to calculate the molecular descriptors (MDs). This program allows calculating several families (0-2) of molecular descriptors such as: constitutional indices, functional group counts, 2D-autocorrelations, topological indices and molecular properties among others. After that, those MDs with values constant or near to constant were removed, thus they were not used in further analysis. Consequently, we finally used 313 descriptors to perform the genetic algorithm (implemented in QSARINS software) for variable selection.

# **QSAR-MLR Model Development**

For the development of the multiple linear regression (MLR) model QSARINS software version 2.2.2 was used, which has been developed in the QSAR Research Unit, University of Insubria, Italy. It allows obtaining MLR models by ordinary least squares (OLS) method [21]. The model was developed based on criteria such as fitting (highest  $R^2$ ), robustness (greatest  $Q^2_{LOO}$ ), stability (lowest  $R^2-Q^2_{LOO}$ ), and low correlation of descriptors (lowest  $K_{xx}$ ), amongst others, so that it has a small difference between fitting, cross-validation and external validation parameters.



Table 1. Structures and experimental values<sup>a</sup> of the data set

1	No	Smile	Log CL <sub>50</sub>	No	Smile	Log CL <sub>50</sub>
CICI=CC=C(O)C(C)=C1	1	ClC1=CC=CC=C1O	1.818	45	CC(O)C	2.826
CICI=CC(CI)=CC(O)=CI	2	ClC1=CC(O)=CC=C1	1.650	46	CC(C)CO	2.485
Cici-cc(Ci)-c(Ci)-c(Ci)-cici   1.002	3	ClC1=CC=C(O)C(Cl)=C1	1.381	47	O=C(O)C1=CC=C(CC)C(CC)=C1C(O)=O	1.664
S	4	ClC1=CC(Cl)=CC(O)=C1	1.112	48		0.594
7         CCI=CC(CI)=CC=CIO         1.516         (SCN2C(CI=CC=CC=CIN=N2)=0)=S         1.530           8         CCI=CC(O)=CC=CIO         1.362         50         CCOP(OCC)(OCI=NN(C2=CC=CC=CC)         1.146           9         OCI=CC=CC=CIO         2.428         C=N1)=S         1.146           10         OC2=CC=CICC=CC=CIO         1.914         52         CCCNCCC         2.048           11         CCI=CC=CC=CI         1.909         53         CCCCNCCC         1.492           13         CICI=CC=C(C)=CI         1.160         55         CC(NCI=CC=CC=CI)         2.293           14         CICI=CC=C(CI)C(CI)=CI         0.863         56         CI=NC=CC=CI         2.538           16         CICI=CC=C(CI)C(CI)=CI         0.920         57         NCI=CC=CC=CIN         2.778           17         CICI=CC(CI)C(CI)=CI         0.858         58         0-CCI=CC=CC=CI         2.140           18         CICI=CC(CI)C(CI)C(CI)=CI         0.623         59         0-C(O)CCCCC         2.020           20         CICI=CC(CI)C(CI)C(CI)=CI         0.636         61         CI(C2=CC=C)CICC=CC=CI         2.287           21         CICI=CC(C)C(C)C(CI)C(CI)=CI         0.536         60         0 ICC=CC=CC=CI         0.887 <td>5</td> <td>ClC1=CC(Cl)=C(Cl)C(O)=C1</td> <td>1.002</td> <td></td> <td>C(CCCC)=C1C(O)=O</td> <td>0.564</td>	5	ClC1=CC(Cl)=C(Cl)C(O)=C1	1.002		C(CCCC)=C1C(O)=O	0.564
Second Color   Seco	6	ClC1=C(O)C(Cl)=CC(Cl)=C1	1.079	49		1.520
1.146	7	CC1=CC(Cl)=CC=C1O	1.516		(SCN2C(C1=CC=CC=C1N=N2)=O)=S	1.550
9 OCI=CC=CC=C1O	8	CC1=CC(O)=CC=C1Cl	1.362	50	, , , , , , , , , , , , , , , , , , , ,	1 146
11	9	OC1=CC=CC=C1O	2.428		C=N1)=S	1.140
12	10	OC2=CC=C1C=CC=CC1=C2	1.554	51	NCCN	3.829
13	11	CC1=CC=CC=C1O	1.914	52	CCCNCCC	2.048
CICI=CC=C(CI)C=CI	12	CC1=CC(O)=CC=C1	1.909	53	CCCCNCCCC	1.492
15	13	ClC1=CC=CC=C1	1.487	54	C=CC(N)=O	3.172
16	14	ClC1=CC=C(Cl)C=C1	1.160	55	CC(NC1=CC=CC=C1)=O	2.293
17	15	ClC1=C(Cl)C(Cl)=CC=C1	0.863	56	C1=NC=CC=C1	2.538
18	16	C1C1=CC=C(C1)C(C1)=C1	0.920	57	NC1=CC=CC=C1N	2.778
C C C C C C C C C C C C C C C C C C C	17	ClC1=CC(Cl)=CC(Cl)=C1	0.858	58	O=CC1=CC=CC=C1	2.140
CICI=CC(CI)=C(CI)C(CI)=C1CI   0.368   61   C1(C2=CC=CC=C2)=CC=CC=C1   0.887	18	ClC1=CC=C(Cl)C(Cl)=C1Cl	0.623	59	O=C(O)CCCCC	2.020
21         CIC1=CC=CC=CI[N+]([O-])=O         1.775         62         C1=CC=CS1         1.981           22         CIC1=CC=CC([N+]([O-])=O)=C1         1.693         63         CC3(O)C2CCIC(N(C)C)         1.881           23         CIC1=C(CI)C([N+]([O-])=O)=CC=C1         1.386         C(C(C(N)=O)=C(O)C(O)IC(C2=CC(O)         1.158           24         NC1=CC=CC=C1         2.418         C4=C3C=CC=C4O)=O)=O         1.158           25         NC1=CC=CCCC C1         1.938         64         CC1=CC(C)=NC(NS(=O)(C2=CC=C(N))         2.196           26         NC1=CC=C(CI)C=C1         1.948         C=C2)=O)=N1         2.196           27         NC1=CC=C(CI)C=C1C1         1.516         65         CCN2C=C(C(O)=O)         1.694           28         NC1=CC=C(CI)C=C(CI)=C1         1.458         66         CCC1C(C)(C)C(C)C(C)C(C)C(C)C(C)         1.694           29         NC1=CC(CI)=CC(CI)=C1         1.458         66         CCC1C(C)(O)C(O)C(C)C(C)C(C)C(C)C(C)         C(O)C(C)C(C)C(C)C(C)C(C)         1.694           31         NC1=CC(CI)=C(CI)C=C1C1         1.160         C(O22CC(C)(OC)C(O)C(C)C(C)C(C)C(C)         C(O1)=O)=O         1.838           32         NC1=CC(CI)=C(CI)C=C1C1         0.882         67         OCC(NC(C(CI)CI)=O)=O(O)         2.310	19	ClC1=C(Cl)C=C(Cl)C(Cl)=C1	0.536	60	O1C=CC=C1	2.207
22         CICI=CC=CC([N+]([O-])=O)=C1         1.693         63         CC3(O)C2CCIC(N(C)C)         1.158           23         CICI=C(CI)C([N+]([O-])=O)=CC=C1         1.386         C(C(C(N)=O)=C(O)C(O)IC(C2=C(O)         1.158           24         NC1=CC=CC=C1         2.418         C4=C3C=CC=C4O)=O)=O         1.158           25         NC1=CC=CCC[CI]         1.938         64         CC1=CC(C)=NC(NS(=O)(C2=CC=C(N))         2.196           26         NC1=CC=CC(CI)=C1         1.948         C=C2)=O)=N1         2.196           27         NC1=CC=C(CI)C=C1CI         1.516         65         CCN2C=C(C(O)=O)         1.694           28         NC1=CC=C(CI)=CCI         1.458         66         CCC1C(C)=C(N3CCNCC3)C=C12)=O         1.694           29         NC1=CC(CI)=C(CI)=C(I)         1.458         66         CCC1C(C)(O)C(O)C(C)C(C)C(C)C(C)         0.00           30         NC1=CC(CI)=C(CI)C=C1CI         1.160         C(OC2CC(C)(OCC)(O)C(C)C(C)C(C)C(C)         1.838           31         NC1=C(CI)C=C(CI)C=C1CI         0.959         0.959         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968         0.968	20	ClC1=CC(Cl)=C(Cl)C(Cl)=C1Cl	0.368	61	C1(C2=CC=CC=C2)=CC=CC=C1	0.887
Cicl=c(cl)c([N+]([O-])=O)=CC=C1	21	ClC1=CC=CC=C1[N+]([O-])=O	1.775	62	C1=CC=CS1	1.981
24 NC1=CC=CC=C1 2.418  25 NC1=CC=CC=C1C1 1.938 64 CC1=CC(C)=NC(NS(=0)(C2=CC=C(N) C=CC))=NC1=CC=CC(C1)=C1 1.948 C=CC=CC(C1)=C1 1.516 65 CCN2C=CC=C(C1)=C1 1.559 C(C1=CC(F)=C(N3CCNCC3)C=C12)=O 1.694  27 NC1=CC=C(C1)C(C1)=C1 1.559 C(C1=CC(F)=C(N3CCNCC3)C=C12)=O 1.694  29 NC1=CC(C1)=CC(C1)=C1 1.458 66 CCC1C(C)(O)C(O)C(O)C(C)CC(C)C(C1)=C1 1.194 (O)C(O23C(O)C(N(C)C)CC(C)O3)C(C1)=C1C1 1.194 (C0C2CC(C)(OC1)C(C1)CC(C1)=C1C1 1.160 C(C1)=C(C1)C=C(C1)C=C1C1 1.160 C(C1)=C(C1)C=C(C1)=C1C1 1.160 C(C1)=C(C1)C=C1C1 1.160 C(C1)=C(C1)C(C1)=C(C1)C=C1C1 1.160 C(C1)=C(C1)C(C1)=C(C1)C=C1C1 1.160 C(C1)=C(C1)C(C1)=C(C1)C=C1C1 1.160 C1=CC=CC([N+]([0-])=O)=C1 2.310  30 NC1=CC(C1)=C(C1)C(C1)=C1C1 0.882 67 OCC(NC(C(C1)C1)=O)C(O) C1=CC=CC=C([N+]([0-])=O)C=C1 2.310  31 NC1=C(C1)C(C1)=CCC=C2 1.266 68 O=C(O)CC1=CC=CC=C1NC2=C(C1) 2.943  32 CC1=CC=CC=C1 1.352 69 O=C(C)NC1=CC=CC=C1C1 2.943  33 CC1=CC=CC=C1 1.434 70 OC1=CC=CC=C1(O)=O 2.230  34 CCCCCCC 0.738 71 OC(CC2=CC1=CC=C2) 2.344	22	C1C1=CC=CC([N+]([O-])=O)=C1	1.693	63		
24 NC1=CC=CC=C1 2.418  25 NC1=CC=CC=C1C1 1.938 64 CC1=CC(C)=NC(NS(=O)(C2=CC=C(N) C=CC=C(N) C=CC=CC(C)) C=CC=C(C) C=CC=C C=CC=CC=CC=CC=CC=CC=CC=CC=CC=CC=	23	ClC1=C(Cl)C([N+]([O-])=O)=CC=C1	1.386			1.158
26 NC1=CC=CC(Cl)=C1 1.948	24	NC1=CC=CC=C1	2.418		C4=C3C=CC=C4O)=O)=O	
26         NC1=CC=CC(Cl)=C1         1.948         C=C2)=O)=N1           27         NC1=CC=C(Cl)C=C1Cl         1.516         65         CCN2C=C(C(O)=O)         1.694           28         NC1=CC=C(Cl)C(Cl)=C1         1.559         C(C1=CC(F)=C(N3CCNCC3)C=C12)=O         1.694           29         NC1=CC(Cl)=CC(Cl)=C1         1.458         66         CCC1C(C)(O)C(O)C(C)C(C(C)CC(C)         0           30         NC1=CC(Cl)=C(Cl)C=C1Cl         1.194         (O)C(OC3C(O)C(N)C)C)C(C)O3C(C)         0         0           31         NC1=C(Cl)C=C(Cl)C=C1Cl         1.160         C(OC2CC(C)(OC)C(O)C(C)OC(C)O2C(C)         1.838           32         NC1=CC(Cl)=C(Cl)C(Cl)=C1Cl         0.959         0 </td <td>25</td> <td>NC1=CC=CC=C1Cl</td> <td>1.938</td> <td>64</td> <td></td> <td>2.106</td>	25	NC1=CC=CC=C1Cl	1.938	64		2.106
28         NC1=CC=C(Cl)C(Cl)=C1         1.559         C(C1=CC(F)=C(N3CCNCC3)C=C12)=0         1.694           29         NC1=CC(Cl)=CC(Cl)=C1         1.458         66         CCC1C(C)(O)C(O)C(C)C(C)CC(C)         0           30         NC1=CC(Cl)=C(Cl)C=C1Cl         1.194         (O)C(OC3C(O)C(N(C)C)CC(C)O3)C(C)         0           31         NC1=C(Cl)C=C(Cl)C=C1Cl         1.160         C(OC2CC(C)(OC)C(O)C(O)C(C)O2)C(C)         1.838           32         NC1=CC(Cl)=C(Cl)C(Cl)=C1Cl         0.959         0 <t< td=""><td>26</td><td>NC1=CC=CC(Cl)=C1</td><td>1.948</td><td></td><td>C=C2)=O)=N1</td><td>2.190</td></t<>	26	NC1=CC=CC(Cl)=C1	1.948		C=C2)=O)=N1	2.190
28  NC1=CC=C(Cl)C(Cl)=C1	27	NC1=CC=C(Cl)C=C1Cl	1.516	65		1.604
30 NC1=CC(Cl)=C(Cl)C=C1Cl 1.194 (O)C(OC3C(O)C(N(C)C)CC(C)O3)C(C) 31 NC1=C(Cl)C=C(Cl)C=C1Cl 1.160 C(OC2CC(C)(OC)C(O)C(C)O2)C(C) 1.838 32 NC1=CC(Cl)=C(Cl)C(Cl)=C1Cl 0.959 33 NC1=C(Cl)C(Cl)=C1Cl 0.882 67 OCC(NC(C(Cl)Cl)=O)C(O) 34 CC1=CC=CC=C1 1.540 C1=CC=C([N+]([O-])=O)C=C1 2.310 35 C12=CC=CC=C1 1.266 68 O=C(O)CC1=CC=CC=C1NC2=C(Cl) 36 C23=C1C(CC3)=CC=CC1=CC=C2 0.968 C=CC=C2Cl 2.943 37 CC1=CC=CC=C1 1.352 69 O=C(C)NC1=CC=CC(O)C=C1 3.450 38 C=CC1=CC=CC1 1.434 70 OC1=CC=CC=C(O)C=C1 3.450 39 CCCCCCC 0.738 71 OC(COC2=C1=CC=C2) 37 OCCCCCCC 2.344	28	NC1=CC=C(Cl)C(Cl)=C1	1.559		C(C1=CC(F)=C(N3CCNCC3)C=C12)=O	1.094
31 NC1=C(CI)C=C(CI)C=C1CI 1.160 C(OC2CC(C)(OC)C(O)C(C)O2)C(C) 1.838  32 NC1=CC(CI)=C(CI)C(CI)=C1CI 0.959  33 NC1=C(CI)C(CI)=C1CI 0.882 67 OCC(NC(C(CI)CI)=O)C(O) 34 CC1=CC=CC=C1 1.540 C1=CC=C([N+]([O-])=O)C=C1  35 C12=CC=CC=C1 1.266 68 O=C(O)CC1=CC=CC=C1NC2=C(CI) 36 C23=C1C(CC3)=CC=CC1=CC=C2 0.968 C=CC=C2CI  37 CC1=CC=CC=C1 1.352 69 O=C(C)NC1=CC=CC(O)C=C1 3.450  38 C=CC1=CC=CC=C1 1.434 70 OC1=CC=CC=C(O)C=C1 3.450  39 CCCCCCC 0.738 71 OC(COC2=C1=CC=C2)  2344	29	NC1=CC(Cl)=CC(Cl)=C1	1.458	66		
31 NC1=C(CI)C=C(CI)C=C(CI)	30	NC1=CC(Cl)=C(Cl)C=C1Cl	1.194			1 020
32       NC1=CC(Cl)=C(Cl)C(Cl)=C1Cl       0.959         33       NC1=C(Cl)C(Cl)=C1Cl       0.882       67       OCC(NC(C(Cl)Cl)=O)C(O)         34       CC1=CC=CC=C1       1.540       C1=CC=C([N+]([O-])=O)C=C1       2.310         35       C12=CC=CC=C1C=CC=C2       1.266       68       O=C(O)CC1=CC=CC=C1NC2=C(Cl)       2.943         36       C23=C1C(CC3)=CC=CC1=CC=C2       0.968       C=CC=C2Cl       2.943         37       CC1=CC=CC=C1C       1.352       69       O=C(C)NC1=CC=C(O)C=C1       3.450         38       C=CC1=CC=CC=C1       1.434       70       OC1=CC=CC=C1C(O)=O       2.230         39       CCCCCCC       0.738       71       OC(COC2=C1C=CC=CC)       2.344	31	NC1=C(Cl)C=C(Cl)C=C1Cl	1.160			1.838
34     CC1=CC=CC=C1     1.540     C1=CC=C([N+]([O-])=O)C=C1     2.310       35     C12=CC=CC=C1C=CC=C2     1.266     68     O=C(O)CC1=CC=CC=C1NC2=C(C1)     2.943       36     C23=C1C(CC3)=CC=CC1=CC=C2     0.968     C=CC=C2C1     2.943       37     CC1=CC=CC=C1C     1.352     69     O=C(C)NC1=CC=C(O)C=C1     3.450       38     C=CC1=CC=CC=C1     1.434     70     OC1=CC=CC=C1C(O)=O     2.230       39     CCCCCCC     0.738     71     OC(COC2=C1C=CC=CC1=CC=C2)     2.344	32	NC1=CC(Cl)=C(Cl)C(Cl)=C1Cl	0.959			
34	33	NC1=C(Cl)C(Cl)=CC(Cl)=C1Cl	0.882	67		2.210
36 C23=C1C(CC3)=CC=CC1=CC=C2 0.968 C=CC=C2Cl 2.943  37 CC1=CC=CC=C1C 1.352 69 O=C(C)NC1=CC=C(O)C=C1 3.450  38 C=CC1=CC=CC=C1 1.434 70 OC1=CC=CC=C1C(O)=O 2.230  39 CCCCCCC 0.738 71 OC(COC2=C1C=CC=C2)  2.943	34	CC1=CC=CC=C1	1.540		C1=CC=C([N+]([O-])=O)C=C1	2.310
36	35	C12=CC=CC=C1C=CC=C2	1.266	68		2 042
38	36	C23=C1C(CC3)=CC=CC1=CC=C2	0.968		C=CC=C2Cl	2.943
39 CCCCCCC 0.738 71 OC(COC2=C1C=CC=CC1=CC=C2)	37	CC1=CC=CC=C1C	1.352	69	O=C(C)NC1=CC=C(O)C=C1	3.450
$\sim$ 2 344	38	C=CC1=CC=CC=C1	1.434	70	OC1=CC=CC=C1C(O)=O	2.230
40 CC(Cl)(Cl)Cl 1.655 CNC(C)C 2.344	39	CCCCCC	0.738	71	,	2 2 4 4
	40	CC(Cl)(Cl)Cl	1.655		CNC(C)C	2.344



No	Smile	Log CL <sub>50</sub>	No	Smile	Log CL <sub>50</sub>
41	ClC1(Cl)C(Cl)(Cl)CCCC(Cl)1Cl	1.117	72	CCCC(NC1=CC(C(C)=O)=C(OCC(O)	3.676
42	C1/C(C1)=C(C1)/C1	1.218		CNC(C)C)C=C1)=O	3.070
43	ClC1=C(Cl)C(Cl)=C(Cl)C(Cl)1Cl	0.431	73	CC(C)(NCC(O)	2.752
44	CC(O)CO	3.292		COC1=NSN=C1N2CCOCC2)C	3.753

<sup>&</sup>lt;sup>a</sup> the values of Log CL<sub>50</sub> are expressed in ug/ml.

#### Validation of model

An important aspect of the QSAR model is validation. Consequently, the obtained model must be carefully checked and thoroughly validated. The QSARINS offers several tools to confirm that the model meets the standards, set by the OECD for the development, validation, acceptance and use of QSAR models in order to increase the confidence in the reliability of data predicted by it. Based on

these principles, the QSAR-MLR model must meet the following criteria: 1) a defined end point; 2) an unambiguous algorithm; 3) a defined domain of applicability; 4) appropriate measures of goodness-of-fit, robustness and predictivity; and 5) a mechanistic interpretation, if possible [26]. In a previous paper we give a complete explanation of all these aspects; for details see reference [27].

In this work we use some cross-validation techniques for internal validation. First, *leave-one-out* (LOO) exercises were done because, with the disturbance of a single compound in a small database, one gets criteria of its robustness. Later, a *leave-many-out* (LMO) technique was developed, which allows studying the behavior of the model when a greater number of compounds are excluded. In order to demonstrate that the model is not the result of a casual correlation, the randomization (*Y-Scrambling*) procedure was also applied. In this process, the answers are randomly located, so that there is no correlation with the descriptors and, as a result, the model performance should decay dramatically. In order to prove the model predictivity, an external validation was performed. For this, we use the prediction set, which has never been used in the model calculation.

#### Acute Toxicity Test with *L. sativa* Seeds

The acute toxicity test was carried out in the ecotoxicology laboratory of the Center of Toxicology and Biomedicine (TOXIMED) in Santiago de Cuba. We followed the procedure previously described by Sobrero and Ronco in 2004 [28], to develop acute toxicity test (120-h exposure) with *L. sativa* seeds; it was used for assessing the phytotoxic effects of the chemical compounds on the seed germination process; the toxicity was assessed as inhibition of the elongation of the radicle and hypocotyl. This test has been recommended and implemented by different agencies for environmental protection for the ecotoxicological assessment of environmental samples and pure compounds and also for evaluating the phytotoxic effect of pesticides on non-target species for the registration of these compounds [29, 30].



## RESULTS AND DISCUSSION

# Develop of the QSAR-MLR Model

Several models were obtained while exploring the best combinations of molecular descriptors which show a high correlation with the response variable (Log CL<sub>50</sub>). Therefore, taking into account the principle of parsimony (describe the maximum information with the least number of descriptors), an analysis of the model's parameters was done. Based on it, a QSAR-MLR model was developed to evaluate the toxic effects with seven variables. In the development of the model compounds the number 64 was detected as statistical outlier and removed; once rejected the model was obtained. The equation and the statistical parameters of the best model are the following:

$R^2 = 0.917$	$R_{adj}^2 = 0.904$	$R^2 - R^2_{adj} = 0.013$	LOF = 0.101
$K_{xx} = 0.315$	Delta K= 0.052	$RMSE_{tr} = 0.236$	$MAE_{tr} = 0.190$
$RSS_{tr} = 3.003$	$CCC_{tr} = 0.957$	s = 0.256	F = 72.231

The number of compounds used as the training set was 56 compounds:  $R^2$  is the coefficient of determination,  $R^2_{adj}$  is adjusted  $R^2$ , s is standard error of estimate, F is variance ratio, LOF is Friedman lack of fit [31, 32] and  $K_{xx}$  is the correlation among descriptors [33]. Delta K is the difference of the correlation between the descriptors ( $K_x$ ) and the descriptors plus the responses ( $K_{xy}$ ), RMSE<sub>tr</sub> is Root-Mean-Square Error in fitting (for the training set), MAE<sub>tr</sub> is Mean Absolute Error in fitting (calculated on the training set), RSS<sub>tr</sub> is the Residual Sum of Squares in the fitting (also for the training set) and CCC<sub>tr</sub> is the concordance correlation coefficient calculated over the training set [34-36].

The model developed in this study showed a  $R^2$  value of 0.917, indicating an appropriate fit to model the toxicity in *L. sativa*; this means that the model explains almost the 92 % of the experimental variance. The LOF parameter showed a low value (0.101), indicating that there was no overfitting risk. The correlation between the descriptors of the model is low because the  $K_{xx}$  is small (0.315), so we can say that there is little redundant information in the selected descriptors. In addition, the correlation between the descriptors and the modeled response is appropriate, in accordance with the Delta K parameter (0.052), with a small error on the calculations of training and parameters estimation (RMSE<sub>tr</sub> = 0.236; MAE<sub>tr</sub> = 0.190; s = 0.256). The Figure 1A shows the scatter plot of the predicted response against experimental, in which it is evident that the compounds are located next to the diagonal line.



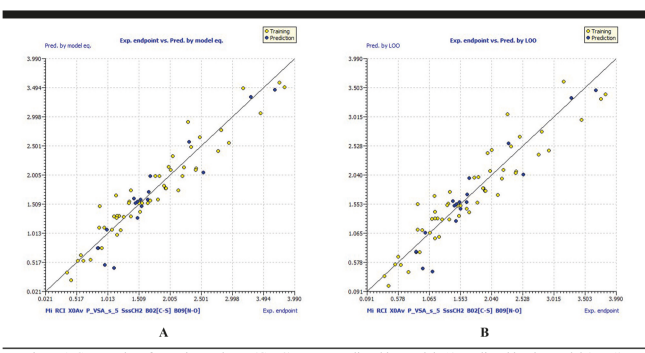


Figure 1. Scatter plot of experimental Log (CL50) versus predicted by model. **A)** predicted by the model (Eq. 1) and **B)** predicted in by LOO croos validation..

# Validation of the developed Model

The principal importance of the validation is to prove the predictability and the robustness of the model. In this work we performed both: internal (LOO, LMO and *Y*-scrambling) and external (prediction set) validation experiments only for the final model. According to the results of the *LOO cross-validation*, it can be stated that internal predictions are good and  $Q^2_{LOO} = 0.874$  has a value very close to  $R^2$  value, so the model is considered internally stable and robust, with a small error in the predictions (RMSE<sub>cv</sub> = 0.289 and MAE<sub>cv</sub> = 0.233). The Figure 1B shows the values predicted by LOO *vs.* experimental Log (CL<sub>50</sub>) values for the training and test sets.

The model performance of internal validation by LMO ( $Q^2_{LMO} = 0.875$ ) is very similar to the obtained with LOO, although it should be noticed that this technique is more useful with larger databases. The Figure 2 shows  $Q^2_{LMO}$  vs.  $K_{XY}$  (correlation between the descriptors and toxicity). Notice that  $Q^2_{LMO}$  values are similar to each other and comparable to  $K_{XY}$  values, corroborating the good fit and stability of the model.

The last experiment of the internal validation was the *Y-scrambling* procedure, as we pointed out before; it was done to demonstrate that the model is not the result of a casual correlation. Here, the answers were placed at random, so that there is no correlation with the descriptors. As a consequence of this, the model performance decays dramatically. The values of  $R^2$  and  $Q^2$  of every iteration, and their averages ( $R^2_{Y-scr}$  and  $Q^2_{Y-scr}$ ) offer the criteria that the model is good, since these parameters are ever lower with regard to the values of the model ( $R^2_{Y-scr} = 0.130$  and  $Q^2_{Y-scr} = -0.265$ ). The  $R^2_{Y-scr}$  and



 $Q^2_{Y-scr}$  values against  $R^2$  and  $Q^2$  of the model are shown in Figure 3. Notice that the values of  $R^2$  and  $Q^2$  of the model are far from the values obtained for those parameters in the *Y-scrambling* experiment, which indicates that the model is not obtained as a result of a random correlation.

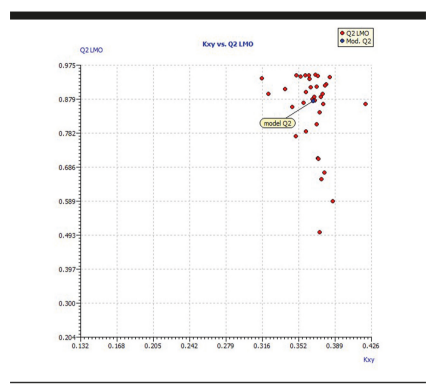


Figure 2. Plot of LMO validations compared with the original model.



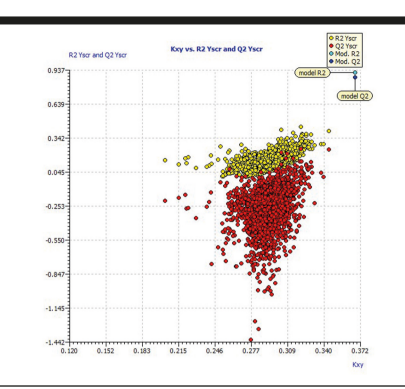


Figure 3. Plot of *Y-scrambled* model compared with the original model.

Table 2. Experimental and predicted values (expressed as Log  $CL_{50}$ ) of the toxicity in L. Sativa

No.	Experimental <sup>a</sup>	Prediction <sup>b</sup>	Residual <sup>c</sup>	No.	Experimentala	Prediction <sup>b</sup>	Residual <sup>c</sup>
1	1.818	1.588	-0.231	38*	1.434	1.608	0.174
2*	1.650	1.588	-0.063	39	0.738	0.563	-0.175
3	1.381	1.303	-0.078	40	1.655	1.533	-0.122
4	1.112	1.303	0.191	41*	1.117	0.427	-0.690
5*	1.002	1.079	0.077	42	1.218	1.067	-0.151
6	1.079	1.079	0.000	43	0.431	0.221	-0.210
7	1.516	1.558	0.042	44*	3.292	3.340	0.048
8	1.362	1.558	0.196	45	2.826	2.780	-0.046
9	2.428	2.125	-0.303	46	2.485	2.656	0.171
10*	1.554	1.482	-0.072	47*	1.664	1.719	0.055
11	1.914	1.826	-0.088	48	0.584	0.647	0.063
12	1.909	1.826	-0.083	49	1.530	1.379	-0.151
13*	1.487	1.279	-0.208	50	1.146	1.660	0.514
14	1.160	0.991	-0.169	51	3.829	3.504	-0.325
15	0.863	0.763	-0.100	52	2.048	2.334	0.286
16	0.920	0.763	-0.157	53*	1.492	1.556	0.064
17*	0.858	0.763	-0.095	54	3.172	3.487	0.315



No.	Experimentala	Prediction <sup>b</sup>	Residual <sup>c</sup>	No.	Experimental <sup>a</sup>	Prediction <sup>b</sup>	Residual <sup>c</sup>
18	0.623	0.545	-0.078	55	2.293	2.915	0.622
19	0.536	0.545	0.009	56*	2.538	2.050	-0.488
20	0.368	0.350	-0.019	57	2.778	2.421	-0.357
21	1.775	1.994	0.219	58	2.140	1.745	-0.395
22*	1.693	1.994	0.301	59	2.020	2.096	0.076
23	1.386	1.752	0.366	60	2.207	1.995	-0.212
24	2.418	2.092	-0.326	61	0.887	1.483	0.596
25*	1.938	1.783	-0.155	62	1.981	2.148	0.167
26	1.948	1.783	-0.165	63	1.158	1.276	0.118
27	1.516	1.530	0.014	64	2.196	-outlier-	-outlier-
28	1.559	1.530	-0.029	65	1.694	1.576	-0.118
29*	1.458	1.530	0.072	66	1.838	1.989	0.151
30	1.194	1.310	0.116	67*	2.310	2.576	0.266
31	1.160	1.310	0.150	68	2.943	2.560	-0.384
32	0.959	1.112	0.153	69	3.450	3.065	-0.385
33	0.882	1.112	0.230	70	2.230	2.137	-0.094
34*	1.540	1.588	0.048	71	2.344	2.490	0.146
35	1.266	1.296	0.030	72*	3.676	3.467	-0.209
36*	0.968	0.477	-0.491	73	3.753	3.586	-0.167
37	1.352	1.531	0.179				

<sup>&</sup>lt;sup>a</sup> Experimental values (See Table 1). <sup>b</sup> Predicted values using Eq. 1. <sup>c</sup> Residual values: log(Obsd)–log(Pred). \* Compounds used as prediction set.

However, it has been recognized that the only way to establish the real predictivity power of any model is through an external validation [37]. The use of an external prediction set is the best way of validating a QSAR model. The result of our model for the prediction showed excellent results (R<sup>2</sup><sub>ext</sub> = 0.896, RMSE<sub>ext</sub> = 0.269, MAE<sub>ext</sub> = 0.199, PRESS<sub>ext</sub> = 1.300, Q<sup>2</sup>-F1 = 0.869, Q<sup>2</sup>-F2 = 0.868, Q<sup>2</sup>-F3 = 0.892, CCC<sub>ext</sub> = 0.939, r<sup>2</sup>m\_aver = 0.797, r<sup>2</sup>m\_delta = 0.097). Where, R<sup>2</sup><sub>ext</sub> is the external determination coefficient [37]; RMSE<sub>ext</sub> is the Root-Mean-Square Error in external prediction; MAE<sub>ext</sub> is the Mean Absolute Error in external prediction; PRESS<sub>ext</sub> is the Predictive Residual Sum of Squares (external validation); Q<sup>2</sup>-F1 [38], Q<sup>2</sup>-F2 [39], Q<sup>2</sup>-F3 [40] are the explained variances in external prediction; CCC<sub>ext</sub> is the Concordance Correlation Coefficient [34-36] and r<sup>2</sup><sub>m\_aver</sub> and r<sup>2</sup><sub>m\_aver</sub> delta are the Roy criteria: average and delta [41]. The predictions of compounds in the external set can be seen in Figure 1 (blue circles) and Table 2.

#### **Aplicability Domain**

A critical aspect in chemometrics and QSAR studies, for either classification or regression models, is the definition of the applicability domain (AD), because only predictions for compounds that fall



within the domain of applicability may be considered reliable. In this work we use the *leverage* (h) and standardized residual approach, described in the literature [42]. The Figure 4 shows the graph of William for the training and prediction sets; as can be seen in this figure, most compounds are within the AD of the model. Only rather few chemicals of the training set showed values of leverage reater than the *critical leverage* value ( $h^*$ =0.444), although they showed values of standard deviation within the limits, which means that they should be considered as influential compounds instead of *outliers*. Therefore, we can say that these models, for their applicability domain, can be used with high accuracy.

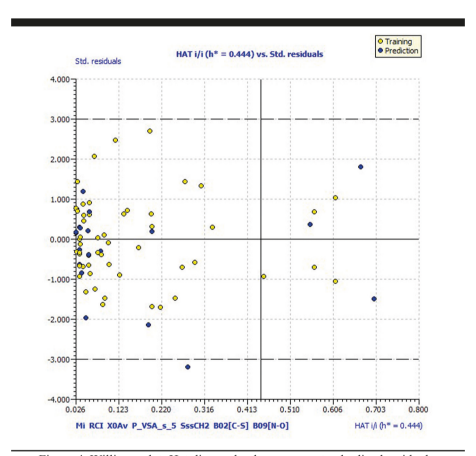


Figure 4. Williams plot. Hat diagonal values versus standardized residuals.

In addition, QSARINS software provides a new approach to determine the application domain: the Insubria graph. This method is based on the *leverage* and model predictions. The Insubria graph is useful to evaluate the position of the molecules, lacking experimental response compared to the structural application domain, and to compare their predictions to those of the compounds having experimental values. We detect that the same compounds that have values over the  $h^*$  in the previous graph showed the same behavior with this approach. The Figure 5 shows the Insubria graph for the applicability domain.



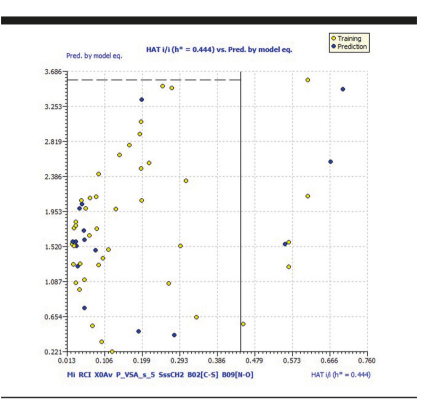


Figure 5. Insubria graph. Hat diagonal values versus predicted data.

# **Experimental Evaluation**

Finally, some chemical compounds, used routinely in the ecotoxicology assays of the laboratory of the TOXIMED, were experimentally tested against *L. sativa*. We carried out the assay as explained above and the results are presented in the following table, as well as the prediction obtained by the model (Eq. 1) for each compound.

Table 3. Experimental results of the evaluated compounds

Compound	Experimentala	Prediction <sup>b</sup>
Methanol	4.670	3.244
Ethanol	4.170	2.967
Chromotropic acid	2.673	3.147
Phenol	1.974	1.900
p-nitroaniline	1.419	2.587

<sup>&</sup>lt;sup>a</sup> Experimental values obtained in the assay in our lab.



<sup>&</sup>lt;sup>b</sup> Predicted values using Eq. 1.

As can be seen, the compounds were sorted in decreasing order of toxicity (according to the experimental results). We can classify as more toxic those compounds with value of Log  $CL_{50}$ <3.0 and as less toxic those with values of Log  $CL_{50}$ >3.0; taking this into account, the computational model predicts the more toxic compounds better. Notice that the prediction of two compounds, phenol and chromotropic acid are good and showed low residual values; it also predicts the p-nitroaniline acceptably well (with a residual about 1.1). However, the prediction of the least toxic compounds (ethanol and methanol) has not been so good (residuals being greater than 1.5). For both compounds the model overestimates the toxicity when compared to the experimental values; the increment in the difference could be caused because in the development of the experiment perhaps there was a loss by evaporation (notice that both compounds are volatile) and the value of the assay could underestimate the real toxicity of the compound. Thus, the experiment should be carried out in the future taking into account this inconvenience, as well as one should evaluate other chemical of interest for the above mentioned laboratory. In general, we can say that this model is a good predictor tool to predict the toxicity over *L. sativa* of chemical compounds and that, for future works, it could become an important tool for the work in the ecotoxicology laboratory of TOXIMED.

#### **CONCLUSION**

In the present work, a QSAR-MLR model was developed by using molecular descriptors calculated using the Dragon software, which adequately predicts the acute toxicity against seeds of *L. sativa*, so as to assess the phytotoxic effects of the chemical compounds on the seed germination. The model was preformed with QSARINS software and extensively validates following the principles established by the OECD; the robustness of the model was tested through internal validation techniques (*LOO*, *LMO*, and *Y-scrambling*), and its predictability was checked through an external prediction set (external validation). The obtained model showed good statistical parameters for training and a prediction set concluding that the proposed computational tools are efficient to predict the acute toxicity against *L. sativa*.

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