

	QMRF identifier (JRC Inventory): Q15-33-0012
	QMRF Title: QSARINS model for prediction of (Benzo-)Triazoles toxicity in <i>Oncorhynchus mykiss</i>
	Printing Date: Dec 11, 2019

1. QSAR identifier

1.1. QSAR identifier (title):

QSARINS model for prediction of (Benzo-)Triazoles toxicity in *Oncorhynchus mykiss*

1.2. Other related models:

1.3. Software coding the model:

PaDEL-Descriptor

A software to calculate molecular descriptors and fingerprints, version 2.18 [ref 2; sect 9.2]

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<http://padel.nus.edu.sg/software/padeldescriptor/index.html>

QSARINS

Software for the development, analysis and validation of QSAR MLR models, version 1.2 (also verified with version 2.2, 2015)

Prof. Paola Gramatica, paola.gramatica@uninsubria.it

<http://www.qsar.it/>

2. General information

2.1. Date of QMRF:

04/02/2015

2.2. QMRF author(s) and contact details:

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2.3. Date of QMRF update(s):

2.4. QMRF update(s):

2.5. Model developer(s) and contact details:

[1]Stefano Cassani Insubria University, Department of Theoretical and Applied Sciences (DiSTA), via J.H. Dunant 3, 21100 Varese (Italy) +390332421439 stefano.cassani@uninsubria.it
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[2]Paola Gramatica Insubria University, Department of Theoretical and Applied Sciences (DiSTA), via J.H. Dunant 3, 21100 Varese (Italy) +390332421573 paola.gramatica@uninsubria.it
<http://www.qsar.it/>

2.6. Date of model development and/or publication:

Developed in 2013, Published in 2014 [ref 4; sect 9.2]

2.7. Reference(s) to main scientific papers and/or software package:

[1]Cassani S et al (2013). Daphnia and fish toxicity of (benzo)triazoles: Validated QSAR models, and interspecies quantitative activity–activity modelling. Journal of Hazardous Material 258-259, 50-60.

DOI: 10.1016/j.jhazmat.2013.04.025

[2]Yap CW , (2011). PaDEL-descriptor: an open source software to calculate molecular descriptors and fingerprints. Journal of Computational Chemistry, 32 (7), 1466-1474. DOI: 10.1002/jcc.21707

[3]Gramatica P et al (2013). QSARINS: A new software for the development, analysis and validation of QSAR MLR models. Journal of Computational Chemistry (Software News and Updates). 34 (24), 2121-2132. DOI: 10.1002/jcc.23361

[4]Gramatica P et al (2014). QSARINS-chem: Insubria datasets and new QSAR/QSPR models for environmental pollutants in QSARINS., Journal of Computational Chemistry (Software News and Updates) 35 (13), 1036-1044. DOI: 10.1002/jcc.23576

2.8. Availability of information about the model:

Non-proprietary. Defined algorithm, available in QSARINS [ref 3,4; sect 9.2]. Training and prediction sets are available in the Supporting Information of the related paper [ref 1; sect 9.2], in the attached sdf files of this QMRF (section 9) and in the QSARINS-Chem database [ref 4; sect 9.2].

2.9. Availability of another QMRF for exactly the same model:

No other information available

3. Defining the endpoint - OECD Principle 1

3.1. Species:

Oncorhynchus mykiss

3.2. Endpoint:

3. Ecotoxic effects 3.3. Acute toxicity to fish (lethality)

3.3. Comment on endpoint:

A selected set of experimental LC 50(96h) data was taken from FOOTPRINT PPDB (Pesticide Properties DataBase) online database [ref 5; sect 9.2].

3.4. Endpoint units:

The median lethal concentrations are reported as the logarithm of the inverse molar concentration: $\log(1/LC50)$

3.5. Dependent variable:

$\log(1/LC50)$ or pLC50

3.6. Experimental protocol:

OECD 203 test protocol

3.7. Endpoint data quality and variability:

The data classified as "verified data", "verified data used for regulatory purposes" and "unverified data from known source" were included in model development. The classification of "verified data", "verified data used for regulatory purposes" and "unverified data from known source" was given directly by the FOOTPRINT PPDB database [ref 5; sect 9.2].

4. Defining the algorithm - OECD Principle 2

4.1. Type of model:

QSAR - Multiple linear regression model (OLS - Ordinary Least Square)

4.2. Explicit algorithm:

Log 1/LC50 O.mykiss SOM Split model

OLS-MLR method. Model developed on a training set of 52 compounds

Log 1/LC50 O.mykiss Ordered response split model

OLS-MLR method. Model developed on a training set of 52 compounds

Log 1/LC50 O.mykiss FULL model

OLS-MLR method. Model developed on a training set of 75 compounds

SOM Split model equation: $pLC50 = 3.49 + 0.32 VP-1 - 0.18 nHBAcc - 1.23 minHBd$

Ordered Response Split model equation: $pLC50 = 3.48 + 0.32 VP-1 - 0.18 nHBAcc - 1.11 minHBd$

Full model equation: $pLC50 = 3.62 + 0.30 VP-1 - 0.17 nHBAcc - 1.20 minHBd$

The modeling descriptors, calculated in PaDEL-Descriptor 2.18, are:

VP-1, nHBAcc, minHBd. See section 4.3 for a more detailed description of the three descriptors.

4.3. Descriptors in the model:

[1]VP-1 dimensionless Valence path, order 1. It is related to molecular dimension and branching, and increasing values of this variable are related to an increase in the observed toxicity values. Additionally, being slightly correlated with LogP in the studied dataset, VP-1 also encodes for hydrophobic properties of molecules

[2]nHBAcc dimensionless Number of hydrogen bond acceptors (using CDK HBondAcceptorCountDescriptor algorithm)

[3]minHBd dimensionless Minimum E-States for (strong) Hydrogen Bond donors

4.4. Descriptor selection:

A total of 729 molecular descriptors of differing types (0D, 1D, 2D) were calculated in PaDEL-Descriptor 2.18 [ref 2; sect 9.2]. Constant and semi-constant values and descriptors found to be correlated pairwise were excluded in a pre-reduction step (one of any two descriptors with a correlation greater than 0.98 was removed to reduce redundant information), and a final set of 227 molecular descriptors were used as input variables for variable subset selection. The models were initially developed by the all-subset-procedure, and then GA was applied to obtain the final population of models (three variables). The optimized parameter used was Q 2LOO (leave-one-out). The GA-VSS, by Ordinary Least Squares regression (OLS), included in QSARINS [ref 3,4; sect 9.2], was applied to select only the best combination of descriptors from input pool: 3 modeling descriptors selected from 227.

4.5. Algorithm and descriptor generation:

Multiple linear regression (Ordinary Least Square method) was applied to generate the model.

Molecular descriptors were generated with the PaDEL-Descriptor software 2.18 [ref 2; sect 9.2].

The input files for descriptor calculation contain information on atom and bond types, connectivity, partial charges and atomic spatial coordinates, relative to the minimum energy conformation of the molecule, and were firstly obtained by the semi empirical AM1 method using the package HYPERCHEM 7.03. Then, these files were converted by OpenBabel 2.3.2 into MDL-MOL format and used as input for the calculation of descriptors in PaDEL-Descriptor. Any user can re-derive the model calculating the molecular descriptors with the PaDEL-Descriptor software 2.18 (recently included in QSARINS 2.2) and applying the given equation (automatically done by QSARINS 2.2).

4.6. Software name and version for descriptor generation:

PaDEL-Descriptor

A software to calculate molecular descriptors and fingerprints, version 2.18
Yap Chun Wei, Department of Pharmacy, National University of Singapore.
<http://padel.nus.edu.sg/software/padeldescriptor/index.html>

HYPERCHEM

Software for molecular drawing and conformational energy optimization, ver. 7.03, 2002.
Phone: (352)371-7744
<http://www.hyper.com/>

OpenBabel

Open Babel: The Open Source Chemistry Toolbox. Used for conversion between HYPERCHEM files (hin) and MDL-MOL files, version 2.3.2, 2012.

Contact not available

http://openbabel.org/wiki/Main_Page

4.7. Chemicals/Descriptors ratio:

Split by SOM model: 52 chemicals / 3 descriptors = 17.33
Split by Ordered response model: 52 chemicals / 3 descriptors = 17.33

Full model: 75 chemicals / 3 descriptors = 25

5. Defining the applicability domain - OECD Principle 3

5.1. Description of the applicability domain of the model:

The applicability domain of the model was verified by the leverage approach and fixed thresholds has been used to define both structural and response outliers (see section 5.4). The plot of leverages (hat diagonals) versus standardised residuals, i.e. the Williams plot, verified the presence of response outliers (i.e. compounds with)

cross-validated standardized residuals greater than 2.5 standard deviation units) and chemicals very structurally influential in determining model parameters (i.e. compounds with a leverage value (h) greater than $3p'/n$ (h^*), where p' is the number of model variables plus one, and n is the number of the objects used to calculate the model). For new compounds without experimental data, leverage can be used as a quantitative measure for evaluating the degree of extrapolation: for compounds with a high leverage value ($h > h^*$), that are structural outliers, predictions should be considered less reliable. In QSARINS the Insubria graph allows to identify for which chemicals the predictions are inter- or extrapolated by the model.

Response and descriptor space: Range of experimental pLC50 *O.mykiss* values: 1.92 / 6.7
Range of descriptor values: VP-1: 1.51 / 10; nHBAcc: 1 / 13;
minHBd: 0 / 0.80.

5.2. Method used to assess the applicability domain:

As stated in section 5.1, the structural applicability domain of the model was assessed by the leverage approach, providing a cut-off hat value ($h^*=0.160$). HAT values are calculated as the diagonal elements of the HAT matrix:

$$H = X(X^T X)^{-1} X^T$$

The response applicability domain can be verified by the standardized residuals in cross-validation greater than 2.5 standard deviation units

5.3. Software name and version for applicability domain assessment:

QSARINS

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5.4. Limits of applicability:

SOM Split model domain: outliers for structure, $hat > 0.277$ (h^*): no. Outliers for response, standardised residuals > 2.5 standard deviation units: amitrole (61-82-5), prothioconazole (178928-70-6). Ordered Response Split model domain: outliers for structure, $hat > 0.277$ (h^*): tricyclazole (41814-78-2), Flumetsulam (98967-40-9), N-(2,6-difluorophenyl)-8-fluoro-5-hydroxy[1,2,4]triazolo[1,5-c]pyrimidine-2-sulfonamide (NO CAS). Outliers for response, standardised residuals > 2.5 standard deviation units: prothioconazole (178928-70-6), 4(1H)-Pyrimidinone (2814-20-2). FULL model domain: outliers for structure, $hat > 0.186$ (h^*): no. Outliers for response, standardised residuals > 2.5 standard deviation units: amitrole (61-

82-5), prothioconazole (178928-70-6),
4(1H)-Pyrimidinone (2814-20-2).

6.Internal validation - OECD Principle 4

6.1.Availability of the training set:

Yes

6.2.Available information for the training set:

CAS RN: Yes

Chemical Name: Yes

Smiles: Yes

Formula: Yes

INChI: No

MOL file: No

6.3.Data for each descriptor variable for the training set:

All

6.4.Data for the dependent variable for the training set:

All

6.5.Other information about the training set:

To verify the predictive capability of the models, the whole dataset (n=75) was split, before model development, into training sets used for model development and prediction sets used later for external validation. Two different splitting techniques were applied: by structural similarity (Self Organizing Maps, SOM, n training= 52) and by ordered response (n training=52). In the SOM splitting, the training and prediction set are structurally balanced, since the splitting was based on the structural similarity analysis (performed with Kohonen artificial neural network, K-ANN or SOM method included in KOALA software [ref 11; sect 9.2]). In the ordered response splitting chemicals were ordered according to their increasing toxicity and one out of every three chemicals was assigned to the prediction set (always including the most and the least persistent compound in the training set, i.e. the lowest and the highest pLC50). This splitting guarantees that the training set covers the entire range of the modeled response.

6.6.Pre-processing of data before modelling:

Transformation of LC50 (mg/L) into Log1/LC50 (mol/L)

6.7.Statistics for goodness-of-fit:

SOM Split model:

$R^2 = 0.76$; CCCtr [ref 6,7; sect 9.2]=0.86; RMSE= 0.54

Ordered response split model:

$R^2 = 0.76$; CCCtr=0.87; RMSE= 0.52

6.8.Robustness - Statistics obtained by leave-one-out cross-validation:

SOM Split model:

$Q^2_{LOO} = 0.71$; CCCcv=0.84; RMSEcv= 0.59

Ordered response Split model:

$Q^2_{LOO} = 0.72$; $CCC_{cv} = 0.84$; $RMSE_{cv} = 0.56$

6.9. Robustness - Statistics obtained by leave-many-out cross-validation:

SOM Split model:

$Q^2_{LMO} 30\% = 0.73$. Ordered response split model:

$Q^2_{LMO} 30\% = 0.73$.

Values of Q^2_{LMO} (average value for 2000 iterations, with 30% of chemicals put out at every iteration) that are high and close to the original Q^2_{LOO} , mean that the model is robust and stable.

6.10. Robustness - Statistics obtained by Y-scrambling:

SOM Split model:

$R^2_{y-sc} = 0.06$

Ordered response split model:

$R^2_{y-sc} = 0.06$.

Low value of scrambled R^2 (average value for 2000 iterations, in where the Y-responses are randomly scrambled), means that the model is not given by chance-correlation.

6.11. Robustness - Statistics obtained by bootstrap:

No information available (since we have calculated Q^2_{LMO})

6.12. Robustness - Statistics obtained by other methods:

No information available

7. External validation - OECD Principle 4

7.1. Availability of the external validation set:

Yes

7.2. Available information for the external validation set:

CAS RN: Yes

Chemical Name: Yes

Smiles: Yes

Formula: Yes

INChI: No

MOL file: No

7.3. Data for each descriptor variable for the external validation set:

All

7.4. Data for the dependent variable for the external validation set:

All

7.5. Other information about the external validation set:

To verify the predictive capability of the models, the dataset ($n=75$) was split, before model development, into a training set used for model development and a prediction set used later for external validation. Two different splitting techniques were applied: by structural similarity (SOM) and by sorted response (n external validation set = 23 in both cases); the range of $pLC50$ are: 2.85 / 5.99 for SOM prediction set, 2.85 / 6.26 for Ordered Response prediction set.

7.6. Experimental design of test set:

In the case of split by sorted response model, chemicals were ordered according to their increasing activity, and one out of every three chemicals was put in the prediction set (always including the most and the least active compounds in the training set). The splitting based on structural similarity (SOM) takes advantage of the clustering capabilities of Kohonen Artificial Neural Network, allowing the selection of a structurally meaningful training set and an equally representative prediction set. Through its clustering capabilities, SOM ensures that both sets are homogeneously distributed within the entire area of the descriptor space; in this case the chemicals in both sets, selected to maximize the coverage of the descriptor space (i.e. representativity), represent the structural variety of the studied data set in a balanced way. The selected training chemicals are those with the minimal distance from the centroid of each cell in the top map. In this case, the representative points of the prediction set are close (in the same cell of the top map) to representative points of the training set in the multidimensional structural descriptor.

7.7. Predictivity - Statistics obtained by external validation:

SOM Split model:

Q^2_{extF1} [ref 8; sect 9.2]= 0.76; Q^2_{extF2}

[ref 9; sect 9.2]= 0.75; Q^2_{extF3}

[ref 10; sect 9.2]= 0.83; CCC_{ex}=0.88; RMSE= 0.46

Ordered response split model:

Q^2_{extF1} = 0.74; Q^2_{extF2} = 0.74; Q^2_{extF3} =

0.77; CCC_{ex}=0.86; RMSE= 0.51 The high values of external Q^2 and concordance

correlation coefficient-CCC (threshold for accepting the external $Q^2_{\text{F1-F2-F3}}$

is 0.70, threshold for CCC is 0.85, [ref 7; sect 9.2]), show that the

model is predictive, when applied to chemicals not used during the model

development (prediction sets).

7.8. Predictivity - Assessment of the external validation set:

The splitting methodology based on similarity analysis and by Ordered response allowed for the selection of meaningful training sets and representative prediction sets.

Training and prediction sets are balanced according to both structure and response. In particular, for response the range of pLC₅₀ values are [1.92 / 6.7][2.85 / 5.99] and [1.92 / 5.99][2.85 / 6.7] respectively for SOM and Ordered Response training and prediction sets.

As concerns structural representativity, the range of descriptors values is:

VP-1: SOM Split training set (1.51 / 10), prediction set (2.74 / 9.61); Ordered response split training set (1.51 / 10), prediction set (2.74 / 9.61)

nHB_{acc}: SOM Split training set (3 / 12), prediction set (1 / 13);

Ordered response split training set (2 / 13), prediction set (1 / 10)

minHB_d: SOM Split training set (0 / 0.78), prediction set (0 / 0.80);

Ordered response split training set (0 / 0.80), prediction set (0 / 0.78)

The applicability domain of the model on the prediction set was verified by the Williams plot: in SOM splitting, only one compound of the prediction set is recognised as outlier for structure (no outliers for response); in the ordered by response splitting, only 1 chemical of the prediction set is outlier for response (no outliers for structure). These results demonstrate the broad applicability domain of the model.

7.9. Comments on the external validation of the model:

The full (N Training set=75) model was then applied to a further Evaluation Set (for an additional external validation) composed of 18 (B)TAZs, became available in the literature. The following parameters are related to this further external validation:

RMSE_{ex}= 0.51; CCC_{ex}= 0.86; Q²_{extF1}=0.78; Q²_{extF2}=0.78; Q²_{extF3}=0.77.

8. Providing a mechanistic interpretation - OECD Principle 5

8.1. Mechanistic basis of the model:

The model was developed by a statistical approach. No mechanistic basis for this physico-chemical property was set a priori, but a mechanistic interpretation of molecular descriptors was provided a posteriori (see 8.2).

8.2. A priori or a posteriori mechanistic interpretation:

A posteriori mechanistic interpretation:

The PaDEL-descriptor full model equation, included in QSARINS [ref 3,4; sect 9.2] is: $pLC50 = 3.62 + 0.30 VP-1 - 0.17 nHBAcc - 1.20 minHBd$

VP-1 is Valence path order 1, which it is related to molecular dimension and branching, and increasing values of this variable are related to an increase in the observed toxicity values. Additionally, being slightly correlated with LogP in the studied dataset, VP-1 also encodes for hydrophobic properties of molecules; generally high values of this descriptor are connected to high/medium toxicity (correlation with pLC50 0.64). nHBAcc is Number of hydrogen bond acceptors (using CDK HBondAcceptorCountDescriptor algorithm), a constitutional descriptor that describes capability of chemicals in participating in H-bonding, and minHBd is Minimum E-States for (strong) Hydrogen Bond donors; the last two descriptors are negatively correlated with the toxicity in fish.

8.3. Other information about the mechanistic interpretation:

No other information available.

9. Miscellaneous information

9.1. Comments:

Given the results of the external validation, this model has a broad applicability domain and therefore unsuccessful applications are probably very reduced. Anyhow, the check of outliers by the Williams

plot and the Insubria graph for chemicals without experimental data (see section 5.1)

will allow to verify the model applicability.

To predict pLC50 for new (benzo-)triazoles without experimental data, it is suggested to apply the equation of the **Full Model**, developed on all the available chemicals (N Training=75).

The full model was then applied to a further Evaluation Set (for an additional external validation) composed of 18 (B)TAZs, became available in the literature. The equation (reported also in section 4.2) and the statistical parameters of the full model, also when applied to the EV,

are the following:

$$\mathbf{pLC50} = 3.62 + 0.30 \text{ VP-1} - 0.17 \text{ nHBAcc} - 1.20 \text{ minHBd}$$

N Training set= 75 (*full model applied to the Evaluation Set-EV,

N=18); $R^2 = 0.76$; $Q^2_{\text{LOO}} = 0.73$; Q^2_{LMO}

$30\% = 0.73$; $\text{CCC} = 0.86$; $\text{CCC}_{\text{cv}} = 0.85$; $\text{RMSE} = 0.51$; $\text{RMSE}_{\text{cv}} = 0.54$; $\text{RMSE}_{\text{ex}} = 0.51$; $\text{CCC}_{\text{ex}} =$

0.86 ; $Q^2_{\text{extF1}} = 0.78$; $Q^2_{\text{extF2}} = 0.78$;

$Q^2_{\text{extF3}} = 0.77$.

9.2. Bibliography:

[1] Cassani S et al (2013). Daphnia and fish toxicity of (benzo)triazoles: Validated QSAR models, and interspecies quantitative activity-activity modelling. *Journal of Hazardous Material* 258-259, 50-60.

DOI: 10.1016/j.jhazmat.2013.04.025

[2] Yap CW (2011). PaDEL-descriptor: an open source software to calculate molecular descriptors and fingerprints. *Journal of Computational Chemistry*, 32 (7), 1466-1474. DOI: 10.1002/jcc.21707

[3] Gramatica P et al (2013). QSARINS: A new software for the development, analysis and validation of QSAR MLR models. *Journal of Computational Chemistry (Software News and Updates)*. 34 (24), 2121-2132. DOI: 10.1002/jcc.23361

[4] Gramatica P et al (2014). QSARINS-chem: Insubria datasets and new QSAR/QSPR models for environmental pollutants in QSARINS, *Journal of Computational Chemistry (Software News and Updates)* 35 (13), 1036-1044. DOI: 10.1002/jcc.23576

[5] FOOTPRINT PPDB (Pesticide Properties DataBase), 2009 <http://sitem.herts.ac.uk/aeru/ppdb/en/> (accessed 30, Jan 2015)

[6] Chirico N and Gramatica P (2011), Real external predictivity of QSAR models: how to evaluate it? Comparison of different validation criteria and proposal of using the concordance correlation coefficient, *Journal of Chemical Information & Modeling*, 51, 2320-2335. DOI: 10.1021/ci200211n

[7] Chirico N & Gramatica P (2012). Real External Predictivity of QSAR Models. Part 2. New Intercomparable Thresholds for Different Validation Criteria and the Need for Scatter Plot Inspection, *Journal of Chemical Information & Modeling* 52 (8), 2044-2058. DOI: 10.1021/ci300084j

[8] Shi LM et al (2001). QSAR Models Using a Large Diverse Set of Estrogens. *Journal of Chemical Information & Computer Science*. 41 186-195. DOI: 10.1021/ci000066d

[9] Schuurman G et al (2008). External Validation and Prediction Employing the Predictive Squared Correlation Coefficient - Test Set Activity Mean vs Training Set Activity Mean. *Journal of Chemical Information & Modeling*. 48 (11), 2140-2145. Doi: 10.1021/ci800253u

[10]Consonni V et al (2009). Comments on the Definition of the Q2 Parameter for QSAR Validation, Journal of Chemical Information & Modeling. 49(7), 1669-1678. DOI: 10.1021/ci900115y

[11]KOALA Rel. 1.0 for Windows, 2001. Todeschini R, Consonni V, Mauri, A. Milan, Italy no url available

9.3.Supporting information:

Training set(s)Test set(s)Supporting information

10.Summary (JRC QSAR Model Database)

10.1.QMRF number:

Q15-33-0012

10.2.Publication date:

2015-06-11

10.3.Keywords:

PaDEL-Descriptor;Oncorhynchus mykiss;LC50;acute fish toxicity;benzotriazole;triazole;QSARINS;

10.4.Comments: