SciMatics SciQSAR model for potency of mutagens in the Bacterial Reverse Mutation Test (Ames test) in *S. typhimurium in vitro*

1. QSAR identifier

1.1 QSAR identifier (title)

SciMatics SciQSAR model for potency of mutagens in the Bacterial Reverse Mutation Test (Ames test) in *S. typhimurium in vitro*, Danish QSAR Group at DTU Food.

1.2 Other related models

MultiCASE CASE Ultra model for potency of mutagens in the Bacterial Reverse Mutation Test (Ames test) in *S. typhimurium in vitro*, Danish QSAR Group at DTU Food.

Leadscope Enterprise model for potency of mutagens in the Bacterial Reverse Mutation Test (Ames test) in *S. typhimurium in vitro*, Danish QSAR Group at DTU Food.

1.3. Software coding the model

SciQSAR version 3.1.00.

2. General information
2.1 Date of QMRF
January 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
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2.6 Date of model development and/or publication

January 2014.

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

Contrera, J.F., Matthews, E.J., Kruhlak, N.L., and Benz, R.D. (2004) Estimating the safe starting dose in phase I clinical trials and no observed effect level based on QSAR modelling of the human maximum recommended daily dose. *Regulatory Toxicology and Pharmacology*, 40, 185 – 206.

SciQSAR (2009) Reference guide: *Statistical Analysis and Molecular Descriptors*. Included within the SciMatics SciQSAR software.

2.8 Availability of information about the model

The model is based on publicly available data collected and kindly donated by Proctor and Gamble (personal communication). The model algorithm is proprietary from commercial software.

2.9 Availability of another QMRF for exactly the same model

- 3. Defining the endpoint
- 3.1 Species

Salmonella typhimurium (multiple strains).

3.2 Endpoint

QMRF 4.10. Mutagenicity

OECD 471 Bacterial Reverse Mutation Test

3.3 Comment on endpoint

The bacterial reversed mutation *in vitro* assay using *Salmonella typhimurium* is also referred to as the Ames Test. The test is used to evaluate compounds mutagenic properties as it detects point mutations, which involve substitution, addition or deletion of one or a few DNA base pairs. The test uses amino acid-dependent strains of *S. typhimurium*. These strains contain a mutation that makes them unable to synthesize the amino acid histidine. Therefore, in the absence of an external histidine source, the cells cannot grow and form colonies. Colony growth is resumed if a reversion of the mutation occurs, allowing the production of histidine to be resumed. Spontaneous reversions occur within each of the strains. If a compound cause an increase in the number of revertant colonies relative to the background level it is said to be positive in the Ames test and therefore assigned mutagenic. Different strains of *S. typhimurium* exist and these have several features that make them more sensitive for the detection of mutations, including responsive DNA sequences at the reversion sites, increased cell permeability to large molecules and elimination of DNA repair systems or enhancement of error-prone DNA repair processes. The specificity of the tester strains can provide some useful information on the types of point mutations that are induced such as frameshift mutations or base-pair mutations.

If a chemical is predicted positive by the overall Ames test QSAR model or if a positive test result for the Ames test exists, this model can refine the information for the endpoint and predict if it is a very clear positive in the test or not. A very clear positive in this context means that a chemical cause at least 10 times more bacterial colonies than the control substance (spontaneous reversions).

3.4 Endpoint units

No units, 1 for positives and 0 for negatives.

3.5 Dependent variable

Potency in the Bacterial Reverse Mutation Test (Ames test) in *Salmonelle typhimurium in vitro*, positive or negative.

3.6 Experimental protocol

The experimental protocol is described in OECD guideline 471 (1997). Briefly, suspensions of bacterial cells are exposed to the test substance in the presence and in the absence of an exogenous metabolic activation system. The most commonly used system is a cofactor supplemented post-mitochondrial fraction (S9) prepared from the livers of rodents. In the plate incorporation method, these suspensions are mixed with an overlay agar and plated immediately onto minimal medium. In the preincubation method, the treatment mixture is incubated and then mixed with an overlay agar before plating onto minimal medium. For both techniques, after two or three days of incubation, revertant colonies are counted and compared to the number of spontaneous revertant colonies on solvent control plates (OECD guideline 471, 1997).

3.7 Endpoint data quality and variability

The Ames test is in general accepted to have a high performance and relatively low interlaboratory variability.

"The reproducibility of Ames tests is limited by the purity of the tested chemical, inconsistencies in the interpretation of dose-response curves, interference of further toxic side effects (such as cytotoxicity), variations in the methodology employed, and variations in the materials used (bacterial strains and metabolic activation mixtures). Nevertheless, the average interlaboratory reproducibility of a series of Ames test data from the National Toxicology Program (NTP) was determined to be 85%." (Kazius *et al.* 2005)

4. Defining the algorithm

4.1 Type of model

This is a categorical (Q)SAR model based on calculated molecular descriptors, and if available the modeller's own or third-party descriptors or measured endpoints can be imported and used as descriptors.

4.2 Explicit algorithm

This is a categorical (Q)SAR model made by use of parametric discriminant analysis to create a linear discriminant function (see 4.5). The specific implementation is proprietary within the SciQSAR software.

4.3 Descriptors in the model

Molecular connectivity indices

Molecular shape indices

Topological indices

Electrotopological (Atom E and HE-States) indices

Electrotopological bond types indices

SciQSAR software provides over 400 built-in molecular descriptors. Additionally, SciQSAR makes it possible to import the modeller's own or third-party descriptors or use measured endpoints as custom descriptors.

4.4 Descriptor selection

The initial descriptor set is manually chosen by the model developer from the total set of built-in descriptors. Furthermore, the set of descriptors applied in the modelling by the program is on top of this selection determined by thresholds for descriptor variance and number of nonzero values likewise defined by the model developer.

47 descriptors were selected from the initial pool of descriptors by the system and used to build the model.

4.5 Algorithm and descriptor generation

For a binary classification problem SciQSAR uses discriminant analysis (DA) to make a (Q)SAR model. SciQSAR implements a broad range of discriminant analysis (DA) methods including parametric and non-parametric approaches. The classic parametric method of DA is applicable in the case of approximately

normal within-class distributions. The method generates either a linear discriminant function (the within-class covariance matrices are assumed to be equal) or a quadratic discriminant function (the within-class covariance matrices are assumed to be unequal). When the distribution is assumed to not follow a particular law or is assumed to be other than the multivariate normal distribution, non-parametric DA methods can be used to derive classification criteria. The non-parametric DA methods available within SciQSAR include the kernel and *k*-nearest-neighbor (kNN) methods. The main types of kernels implemented in SciQSAR include uniform, normal, Epanechnikov, bi-weight, or tri-weight kernels, which are used to estimate the group specific density at each observation. Either Mahalanobis or Euclidean distances can be used to determine proximity between compound-vectors in multidimensional descriptor space. When the kNN method is used, the Mahalanobis distances are based on the pooled covariance matrix. When the kernel method is used, the Mahalanobis distances are based on either the individual within-group covariance matrices or the pooled covariance matrix. (Contrera *et al.* 2004)

If the data outcome is continuous, regression analysis is used to build the predictive model. Within SciQSAR several regression methods are available: ordinary multiple regression (OMR), stepwise regression (SWR), all possible subsets regression (PSR), regression on principal components (PCR) and partial least squares regression (PLS). The choice of regression method depends on the number of independent variables and whether correlation or multicollinearity among the independent variables exists: OMR is acceptable with a small number of independent variables, which are not strongly correlated. SWR is used under the same circumstances as OMR but with greater number of variables. PSR is used for problems with a great number of independent variables. PCR and PLS are useful when a high correlation or multicollinearity exist among the independent variables. (SciQSAR 2009)

To test how stable the developed models are, SciQSAR have built-in cross-validation procedures (see 6.).

For this model, the linear method was used.

4.6 Software name and version for descriptor generation

SciQSAR version 3.1.00.

4.7 Descriptors/chemicals ratio

In this model 47 descriptors were used. The training set consists of 187 compounds. The descriptor/chemical ratio is 1:4 (47:187).

5. Defining Applicability Domain

5.1 Description of the applicability domain of the model

The definition of the applicability domain consists of two components; the definition in SciQSAR and the inhouse further refinement algorithm on the output from SciQSAR to reach the final applicability domain call.

1. SciQSAR

The first criterion for a prediction to be within the models applicability domain is that all of the descriptor values for the test compound can be calculated by SciQSAR. If SciQSAR cannot calculate each descriptor value for the test chemical no prediction value is given by SciQSAR and it is considered outside the model's applicability domain.

2. The Danish QSAR group

The Danish QSAR group has applied a stricter definition of applicability domain for its SciQSAR models. In addition to the applicability domain definition made by SciQSAR a second criterion has been applied for predictions generated from (Q)SAR models with a binary endpoint. For each prediction SciQSAR calculates the probability (p) for the test compound's membership in one of the two outcome classes (positive or negative). The probability of membership in a class is a measure of how well training set knowledge is able to discriminate a positive prediction from a negative prediction within the nearest space of the subject compound-vector. The probability of membership value is also a measure of the degree of confidence of a prediction. The Danish QSAR group uses this probability for a prediction to further define the model's applicability domain. Only positive predictions with a probability equal to or greater than 0.7 and negative predictions with a probability equal to or less than 0.3 are accepted. Positive predictions with a probability between 0.5 and 0.7 as well as negative predictions with a probability between 0.3 and 0.5 are considered outside the model's applicability domain. When these predictions are wed out the accuracy of the model in general increases at the expense of reduced model coverage. Furthermore, as SciQSAR does not define a structural domain, only predictions which were within either Leadscope structural domain (defined as at least one training set chemical within a Tanimoto distance of 0.7) or CASE Ultra structural domain (no unknown fragments for negatives and maximum 1 unknown fragment for positives) were defined as being inside the SciQSAR applicability domain.

5.2 Method used to assess the applicability domain

The system does not generate predictions if it cannot calculate each descriptor value for the test compound.

Only positive predictions with probability equal to or greater than 0.7 and negative predictions with probability equal to or less than 0.3 were accepted.

5.3 Software name and version for applicability domain assessment

SciQSAR version 3.1.00.

5.4 Limits of applicability

The Danish QSAR group applies an overall definition of structures acceptable for QSAR processing which is applicable for all the in-house QSAR software, i.e. not only SciQSAR. According to this definition accepted structures are organic substances with an unambiguous structure, i.e. so-called discrete organics defined as: organic compounds with a defined two dimensional (2D) structure containing at least two carbon atoms, only certain atoms (H, Li, B, C, N, O, F, Na, Mg, Si, P, S, Cl, K, Ca, Br, and I), and not mixtures with two or more 'big components' when analyzed for ionic bonds (for a number of small known organic ions assumed not to affect toxicity the 'parent molecule' is accepted). Structures with less than two carbon atoms or containing atoms not in the list above (e.g. heavy metals) are rendered out as not acceptable for further QSAR processing. Calculation 2D structures (SMILES and/or SDF) are generated by stripping off accepted organic and inorganic ions. Thus, all the training set and prediction set chemicals are used in their non-ionized form. See 5.1 for further applicability domain definition.

6. Internal validation
6.1 Availability of the training set
Yes
6.2 Available information for the training set
CAS
SMILES
6.3 Data for each descriptor variable for the training set
No
6.4 Data for the dependent variable for the training set
All

6.6 Pre-processing of data before modelling

6.5 Other information about the training set

187 compounds are in the training set: 104 positives and 83 negatives.

Only structures acceptable for SciQSAR were used in the final training set. That is, only discrete organic chemicals as described in 5.4 were used. In case of replicate structures, one of the replicates was kept if all the compounds had the same activity and all were removed if they had different activity. No further structures accepted by the software were eliminated (i.e. outliers).

6.7 Statistics for goodness-of-fit

SciQSARs own internal performance test of the model gave the following Cooper's statistics for predictions within the applicability domain as defined by SciQSAR (i.e. the first criterion described in 5.1):

- Sensitivity (true positives / (true positives + false negatives)): 82.7%
- Specificity (true negatives / (true negatives + false positives)): 84.3%
- Concordance ((true positives + true negatives) / (true positives + true negatives + false positives + false negatives)): 83.5%

6.8 Robustness – Statistics obtained by leave-one-out cross-validation
Not performed.
6.9 Robustness – Statistics obtained by leave-many-out cross-validation
SciQSAR's own internal 10-fold cross-validation (10*10% out) procedure was used for predictions within the applicability domain as defined by SciQSAR (i.e. the first criterion described in 5.1). As the probability domain was not applied (i.e. the second criterion described in 5.2) the accuracy of the predictions when applying this domain can be expected to be higher than reflected in these cross-validation results. This gave the following Cooper's statistics:
 Sensitivity (true positives / (true positives + false negatives)): 75.0% Specificity (true negatives / (true negatives + false positives)): 74.7% Concordance ((true positives + true negatives) / (true positives + true negatives + false positives + false negatives)): 74.9%
6.10 Robustness - Statistics obtained by Y-scrambling
Not performed.
6.11 Robustness - Statistics obtained by bootstrap Not performed.
6.12 Robustness - Statistics obtained by other methods
Not performed.

- 7. External validation
- 7.1 Availability of the external validation set
- 7.2 Available information for the external validation set
- 7.3 Data for each descriptor variable for the external validation set
- 7.4 Data for the dependent variable for the external validation set
- 7.5 Other information about the training set
- 7.6 Experimental design of test set
- 7.7 Predictivity Statistics obtained by external validation
- 7.8 Predictivity Assessment of the external validation set
- 7.9 Comments on the external validation of the model

External validation not performed.

8. Mechanistic interpretation

8.1 Mechanistic basis of the model

The SciQSAR software provides over 400 calculated physico—chemical, electrotopological E-state, connectivity and other molecular descriptors. The descriptors selected for the model may indicate modes of action that are obvious for persons with expert knowledge about the endpoint.

8.2 A priori or posteriori mechanistic interpretation

A posteriori mechanistic interpretation. The descriptors selected for the model may provide a basis for mechanistic interpretation.

8.3 Other information about the mechanistic interpretation

9. Miscellaneous information

9.1 Comments

This model can only be used if the compound has been predicted positive in an overall Ames model or if a positive Ames test result exists. If so, the model can be used to predict if the chemical will be clearly positive and cause at least 10 times more bacterial colonies than the control substance.

9.2 Bibliography

Kazius, J., McGuire, R., and Burs, R. (2005) Derivation and Validation of Toxicophores for Mutagenicity Prediction. *J. Med. Chem.*, 48, 312-320.

OECD guideline 471 (1997) OECD Guidelines for the Testing of Chemicals No. 471, Bacterial Reverse Mutation Test. Organisation for Economic Cooperation and Development; Paris, France. Available online at: http://www.oecd-ilibrary.org/environment/oecd-guidelines-for-the-testing-of-chemicals-section-4-health-effects 20745788.

9.3 Supporting information