# Annex I –(Q)SAR model reporting format (QMRF) v.2.1

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QMRF v.2.1 is a minor update of the QMRF template, as it only concerns the description of the QMRF fields. The only exception is Section 10, which has been entirely removed. This section referred to the JRC QSAR Model Database, which is not updated anymore.

The update is based on the version 2.0[[1]](#footnote-1).

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|  | **Element** | **Explanation** |
| **1.** | **QSAR identifier** |  |
| 1.1. | QSAR identifier (title) | 1. Stacked PLS q-RASPR model for zeta potential of engineered nanomaterials
2. Stacked MLP q-RASPR model for zeta potential of engineered nanomaterials
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| 1.2 | Other related models | MLR q-RASPR models (M1-M4) for zeta potential of engineered nanomaterials |
| 1.3. | Software coding the model | PLS\_SingleY\_1.0 (Roy’s group), Jupyter Notebook using Scikit-learn |
| **2.** | **General information** |  |
| 2.0 | Abstract | Four MLR q-RASPR models for zeta potential of engineered nanomaterials were developed after suitable feature selection. The predictions for these models were taken as descriptors to develop Stacked PLS and MLP q-RASPR models for zeta potential of engineered nanomaterials |
| 2.1. | Date of QMRF | 29th February 2024 |
| 2.2. | QMRF author(s) and contact details | Arkaprava Banerjee, Joyita Roy, Kunal Roy\*kunal.roy@jadavpuruniversity.in  |
| 2.3. | Date of QMRF update(s) | -NA- |
| 2.4. | QMRF update(s) | -NA- |
| 2.5. | Model developer(s) and contact details | Arkaprava Banerjee, Joyita Roykunal.roy@jadavpuruniversity.in  |
| 2.6. | Date of model development and/or publication | 2024 |
| 2.7. | Reference(s) to main scientific papers and/or software package | 1. Banerjee A, Roy K. Mol Divers 2022, 26, 2847-2862
2. Banerjee A, Gajewicz-Skretna A, Roy K. Mol Inform 2023, 42, 2200261
3. Banerjee A, Kar S, Pore S, Roy K. Nanotoxicology 2023, 17, 78-93
4. Wold S, Sjostrom M, Eriksson L. Chemom Intell Lab Syst 2001, 58, 109-130
5. Hackeling G. Mastering Machine Learning with scikit-learn. Packt Publishing Ltd; 2017
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| 2.8. | Availability of information about the model  | The model is non-proprietary: A full description of the model algorithm is available, and training and test sets are available as the Supplementary material of the original research article |
| 2.9. | Availability of another QMRF for exactly the same model | No |
| **3** | **Defining the endpoint - OECD Principle 1: “A DEFINED ENDPOINT"** | **PRINCIPLE 1: “A DEFINED ENDPOINT". ENDPOINT refers to any physicochemical, biological, or environmental property/activity/effect that can be measured and therefore modelled. The intent of PRINCIPLE 1 (a (Q)SAR should be associated with a defined endpoint) is to ensure clarity in the endpoint being predicted by a given model, since a given endpoint could be determined by different experimental protocols and under different experimental conditions. It is therefore important to identify the experimental system and test conditions that is being modelled by the Q)SAR.** |
| 3.1. | Species | -NA- |
| 3.2. | Endpoint | Zeta potential of engineered nanomaterials |
| 3.3 | Comment on endpoint | -NA-  |
| 3.4. | Endpoint units | mV |
| 3.5. | Dependent variable | Zeta potential of engineered nanomaterials was the dependent variable that was modelled. No logarithmic transformation was done. |
| 3.6. | Experimental protocol | -NA-. |
| 3.7. | Endpoint data quality and variability | The physicochemical data were obtained under the EU-FP7 NanoMILE project. The Zeta Potential was measured in water (pH=6.5-8.5). |
| **4** | **Defining the algorithm - OECD Principle 2 : “AN UNAMBIGUOUS ALGORITHM”** | **PRINCIPLE 2: “AN UNAMBIGUOUS ALGORITHM”. The (Q)SAR estimate of an endpoint is the result of applying an ALGORITHM to a set of structural parameters which describe the chemical structure. The intent of PRINCIPLE 2 (a (Q)SAR should be associated with an unambiguous algorithm) is to ensure transparency in the model algorithm that generates predictions of an endpoint from information on chemical structure and/or physicochemical properties. In this context, algorithm refers to any mathematical equation, decision rule or output approach.** |
| 4.1. | Type of model | Equation-based (PLS q-RASPR), Machine Learning model (MLP q-RASPR) |
| 4.2. | Explicit algorithm | After suitable feature selection (QSPR descriptors) from the training set, these features were used to compute the RASPR descriptors. Data fusion was performed, by clubbing the initially selected QSPR descriptors with the computed RASPR descriptors, and a further feature selection algorithm was applied. Four different MLR q-RASPR models were developed using different combinations of selected descriptors. The predictions from these models were used as descriptors to generate PLS and MLP q-RASPR models that were used as the final stacking regressors after optimization of the associated hyperparameters. The detailed algorithm of q-RASAR/q-RASPR can be explored from the following literature sources: 1. Banerjee A, Roy K. Mol Divers 2022, 26, 2847-2862
2. Banerjee A, Gajewicz-Skretna A, Roy K. Mol Inform 2023, 42, 2200261
3. Banerjee A, Roy K. Chem Res Toxicol 2023, 36, 446-464.
 |
| 4.3. | Descriptors in the model | 4 descriptors (Ypred(M1), Ypred(M2), Ypred(M3), Ypred(M4)) |
| 4.4. | Descriptor selection | A total of 72 structural, physicochemical, and periodic table QSPR descriptors were initially screened by employing the Stepwise Selection and the Genetic Algorithm approaches. From these 72 descriptors, the Best Subset Selection was applied to select a pool of 10 QSPR descriptors. Additionally we have taken the log-transformed hydrodynamic diameter since we found that it possessed significant correlation with the training set response values. The RASPR descriptors were computed on these (10+1) QSPR descriptors. The generated 18 different RASPR descriptors were clubbed with the selected 11 QSPR descriptors to obtain a pool of 29 descriptors. Feature selection in the form of Best Subset Selection was applied and four different MLR q-RASPR models were generated. The predictions for these four models were then used as descriptors and the final stacked PLS q-RASPR and MLP q-RASPR models were generated.  |
| 4.5. | Algorithm and descriptor generation | Structural, physicochemical, and periodic table descriptors (QSPR)Similarity and error-based descriptors (q-RASPR) |
| 4.6. | Software name and version for descriptor generation | Elemental Descriptor Calculator, RASAR-Desc-Calc-v3.0.2 |
| 4.7. | Chemicals/Descriptors ratio | 53/4 |
| **5** | **Defining the applicability domain - OECD Principle 3: “A DEFINED DOMAIN OF APPLICABILITY”** | **PRINCIPLE 3: “A DEFINED DOMAIN OF APPLICABILITY”. APPLICABILITY DOMAIN refers to the response and chemical structure space in which the model makes predictions with a given reliability. Ideally the applicability domain should express the structural, physicochemical and response space of the model. The CHEMICAL STRUCTURE (x variable) space can be expressed by information on physicochemical properties and/or structural fragments. The RESPONSE (y variable) can be any physicochemical, biological or environmental effect that is being predicted. According to PRINCIPLE 3 a (Q)SAR should be associated with a defined domain of applicability. Section 5 can be repeated (e.g., 5.a, 5.b, 5.c, etc) as many times as necessary if more than one method has been used to assess the applicability domain.** |
| 5.1. | Description of the applicability domain of the model | The applicability domain was defined using the Leverage approach that utilizes a HAT matrix and identifies structural outliers. These structural outliers are compounds that are structurally different from the other compounds and thus, do not fall under the chemical space defined by the model. The traditional approach to identifying structural outliers with the leverage approach uses the descriptor values as the source of information, however, since our final regressors are stacked models, we have treated the predicted values of the individual MLR q-RASPR models as descriptors, From the training set, one compound (#28) had a higher leverage value and thus, was considered as an outlier. In the test set, no compounds were outside the AD.  |
| 5.2. | Method used to assess the applicability domain | Leverage approach |
| 5.3. | Software name and version for applicability domain assessment | Hi\_Calculator-v2.0 |
| 5.4. | Limits of applicability | Compounds having leverage values lower than the threshold (h\*=0.283) are considered as inside the applicability domain (AD). From the training set, one compound (#28) had a higher leverage value and thus, was considered as an outlier. In the test set, no compounds were outside the AD.  |
| **6** | **Defining goodness-of-fit and robustness (internal validation) – OECD Principle 4: “APPROPRIATE MEASURES OF GOODNESS-OF-FIT, ROBUSTENESS AND PREDICTIVITY”** | **PRINCIPLE 4: “APPROPRIATE MEASURES OF GOODNESS-OF-FIT, ROBUSTENESS AND PREDICTIVITY”. PRINCIPLE 4 expresses the need to perform validation to establish the performance of the model. GOODNESS-OF-FIT and ROBUSTNESS refer to the internal model performance.** |
| 6.1. | Availability of the training set | It is available in the Supplementary Information. |
| 6.2. | Available information for the training set | Available information for the training set: Chemical names are available and the data points are for nanomaterials. |
| 6.3. | Data for each descriptor variable for the training set | Available and attached as the Supporting information |
| 6.4. | Data for the dependent variable for the training set | Available and attached as the Supporting information |
| 6.5. | Other information about the training set | The number of training set data points: 53 engineered nanomaterials. The training set was obtained by randomly assigning 75% of the data points from the whole dataset.  |
| 6.6. | Pre-processing of data before modelling | Logarithmic transformation of the “hydrodynamic diameter measured by DLS” was performed. The raw data and processed data are not given as per this QMRF; however, the raw data may be available from other QMRFs of this round-robin exercise.  |
| 6.7. | Statistics for goodness-of-fit | Stacked PLS q-RASPR: R2=0.681, MAEtrain=13.255, RMSEC=18.417Stacked MLP q-RASPR: R2=0.695, MAEtrain=12.952, RMSEC=18.015 |
| 6.8. | Robustness - Statistics obtained by leave-one-out cross-validation | Stacked PLS q-RASPR: Q2(LOO)=0.657. MAE(LOO)=13.766Stacked MLP q-RASPR: Q2(LOO)=0.645, MAE(LOO)=13.957 |
| 6.9. | Robustness - Statistics obtained by leave-many-out cross-validation | -NA- |
| 6.10. | Robustness - Statistics obtained by Y-scrambling | -NA- |
| 6.11. | Robustness - Statistics obtained by bootstrap | -NA- |
| 6.12. | Robustness - Statistics obtained by other methods | -NA- |
| **7** | **Defining predictivity (external validation) – OECD Principle 4: “APPROPRIATE MEASURES OF GOODNESS-OF-FIT, ROBUSTENESS AND PREDICTIVITY”** | **PRINCIPLE 4: “APPROPRIATE MEASURES OF GOODNESS-OF-FIT, ROBUSTENESS AND PREDICTIVITY”. PRINCIPLE 4 expresses the need to perform validation to establish the performance of the model. PREDICTIVITY refers to the external model validation. Section 7 can be repeated (e.g., 7.a, 7.b, 7.c, etc) as many times as necessary if more validation studies need to be reported in the QMRF.** |
| 7.1. | Availability of the external validation set | Available |
| 7.2. | Available information for the external validation set | Available information for the test set: Chemical names are available and the data points are for nanomaterials. |
| 7.3. | Data for each descriptor variable for the external validation set | Available and attached as supporting information |
| 7.4. | Data for the dependent variable for the external validation set | Available and attached as supporting information |
| 7.5. | Other information about the external validation set | Number of test set data points: 18 engineered nanomaterials  |
| 7.6. | Experimental design of test set | Randomly setting aside chemicals before modeling |
| 7.7. | Predictivity - Statistics obtained by external validation | Stacked PLS q-RASPR: $r\_{test}^{2}=0.960, Q\_{ext}^{2}=0.951, MAE=4.402, RMSEP=6.320$Stacked MLP q-RASPR: $r\_{test}^{2}=0.961, Q\_{ext}^{2}=0.963, MAE=4.038, RMSEP=5.500$ |
| 7.8. | Predictivity - Assessment of the external validation set | The external validation set is sufficiently large (ntest=18) and representative of the applicability domain (No compounds outside AD (see section 5.1)). The range of descriptors for the training and test sets is as follows: Ypred(M1): Training set range=86.657, Test set range=95.196Ypred(M2): Training set range=102.768, Test set range=124.117Ypred(M3): Training set range=98.843, Test set range=111.952Ypred(M4): Training set range=99.156, Test set range=105.526The observed response range for the training and test sets is as follows:Yobs: Training set range=114.6 (Max=64.3, Min=-50.3), Test set=98.9 (Max=52.7, Min=-46.2) |
| 7.9. | Comments on the external validation of the model | Highly predictive |
| **8** | **Providing a mechanistic interpretation - OECD Principle 5: “A MECHANISTIC INTERPRETATION, IF POSSIBLE”** | **PRINCIPLE 5: “A MECHANISTIC INTERPRETATION, IF POSSIBLE”. According to PRINCIPLE 5, a (Q)SAR should be associated with a mechanistic interpretation, if possible.** |
| 8.1. | Mechanistic basis of the model | -NA- since the final models are stacked models that use the predicted values of individual MLR q-RASPR models as descriptors and not the structural, physicochemical, and periodic table descriptors.  |
| 8.2. | A priori or a posteriori mechanistic interpretation | -NA- |
| 8.3. | Other information about the mechanistic interpretation | -NA- |
| **9** | **Miscellaneous information** |  |
| 9.1. | Comments | None |
| 9.2. | Bibliography | 1. Srisongkram T. Chem Res Toxicol 2023, 36, 1961-1972.
2. Alexander DLJ, Tropsha A, Winkler DA. J Chem Inf Model 2015, 55, 1316-1322.
 |
| 9.3 | Supporting information | The training and test datasets used to develop stacked PLS and MLP q-RASPR models have been attached in the Supplementary Information. |

1. Triebe, J., Worth, A., Janusch Roi, A. and Coe, A., JRC QSAR Model Database: EURL ECVAM DataBase service on ALternative Methods to animal experimentation: To promote the development and uptake of alternative and advanced methods in toxicology and biomedical sciences: User Support & Tutorial, EUR 28713 EN, Publications Office of the European Union, Luxembourg, 2017, ISBN 978-92-79-71406-1, doi:10.2760/905519, JRC107491. [↑](#footnote-ref-1)