## **RSC Advances**



**PAPER** 

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Cite this: RSC Adv., 2014, 4, 50713

# Enalos InSilicoNano platform: an online decision support tool for the design and virtual screening of nanoparticles

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Engineered nanoparticles (ENPs) are being extensively used in a great variety of applications with a pace that is increasingly growing. The evaluation of the biological effects of ENPs is of utmost importance and for that experimental and most recently computational methods have been suggested. In an effort to computationally explore available datasets that will lead to ready-to-use applications we have developed and validated a QNAR model for the prediction of the cellular uptake of nanoparticles in pancreatic cancer cells. Our insilico workflow was made available online through the Enalos InSilicoNano platform (http://enalos.insilicotox.com/QNAR\_PaCa2/), a web service based solely on open source and freely available software that was developed with the purpose of making our model available to the interested user wishing to generate evidence on potential biological effects in the decision making framework. This web service will facilitate the computer aided nanoparticle design as it can serve as a source of activity prediction for novel nano-structures. To demonstrate the usefulness of the web service we have exploited the whole PubChem database within a virtual screening framework and then used the Enalos InSilicoNano platform to identify novel potent nanoparticles from a prioritized list of compounds.

Received 29th July 2014 Accepted 23rd September 2014

DOI: 10.1039/c4ra07756c

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#### 1. Introduction

Nanotechnology has already contributed a wide range of significant products in several areas of application such as medicine, environment, electronics, cosmetics, defense *etc.*<sup>1-4</sup> Due to the unique physical and chemical properties that particles possess in the nano scale, the research in the field is swiftly progressing and many more promising applications are rapidly being developed. As a result, nanoparticles (NPs) are increasingly used in our everyday life followed by concerns that have now been raised for their safety that is still to be explored.<sup>5-10</sup> The bioactivity profile and risk assessment of NPs, including exposure and hazard assessment, is now gaining greater concern by academia, government and industry and many initiatives worldwide are working on defining the strategies and setting the priorities towards this goal (*i.e.* NNI National Nanotechnology Initiative, NanoSafety Cluster).

The evaluation of NPs biological activity and toxicity by *in vitro* and *in vivo* studies is costly and time consuming and therefore alternative novel techniques that are fast, inexpensive and reduce the animal testing are required. To date a great number of Quantitative Structure Activity (QSAR) models have been proposed in literature. These models usually cover the biological profile of small organic molecules and have been proven accurate in predicting the biological effect for a wide

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range of molecular scaffolds. This is not the case for NPs that have recently emerged as important chemical structures with a wide range of significant properties that find applications in different areas of interest. Although 'classic' QSAR models own a great proportion of their success in the presence of organized databases, no such databases are available for NPs. Experimental data are scarce and produced by different groups of scientists following different protocols and it is often difficult to select and combine the available information from different sources. On top of that, the structural characteristics of NPs cannot be encoded by the "conventional" widely used 2D and 3D molecular descriptors. NPs include organic as well as inorganic elements with sometimes unknown composition and highly complex structures that demand new approaches for developing molecular descriptors. These hurdles have already been recognized and now international efforts are being organized towards the development of large datasets for NPs and the computational exploration of these results.

The potential of computational methods for advancing risk assessment of NPs is commonly accepted and a few computational attempts to predict the toxicity of NPs are reported in the literature the last few years. 18-24 As mentioned, although "classic" Quantitative Structure Activity Relationship (QSAR) models have been for long proposed in the literature to assess different properties of compounds, Quantitative Nanostructure Activity (QNAR) models have not yet been extensively studied and limited examples have been published. 1,5,7,9 Many factors have contributed in this including major hurdles such as lack of

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organized datasets and inadequate descriptors for NPs. On top of that attempts on the computational exploration of the activity of NPs and the produced QNAR models in principal are not made directly available to the community to be further used as useful tools for the risk assessment of novel NPs. Thus their utility is quite limited, whereas an online version of the model could spread the knowledge gained and generate more advancement in the field.

One of the few organized datasets on NPs that has been presented in literature includes the cellular uptake of 109 NPs in pancreatic cancer cells (PaCa2). Each NP within this dataset includes the same metal core (iron oxide/NH2 cores) but different surface modifiers which are organic small molecules conjugated to the NP surface.25 Different computational approaches have been proposed in literature for the exploitation of this dataset with interesting results in model development. Recent models presented in the literature are briefly discussed below.

In 2010 Fourches et al.26 presented a QNAR model based on MOE descriptors calculated for the organic molecules conjugated to the NP surface and k-nearest neighbors (kNN) methodology. The proposed model was proven robust and accurate as indicated by external predictions, cross validation and Y randomization. Winkler and coworkers<sup>27,28</sup> also studied this dataset and generated quantitative, predictive and informative models of cellular uptake using a pool of molecular descriptors. In a recent publication, Y. T. Chau and C. W. Yap<sup>29</sup> used four different modeling methods, namely Naive Bayes, logistic regression, k nearest neighbor and support vector machine, to develop candidate models. A consensus model was developed using the top 5 candidate models and validated by repeating the entire model development process five times using different combinations of training and validation sets. The final consensus model had a sensitivity of 86.7 to 98.2% and a specificity of 67.3 to 76.6%. In a different publication Toropov et al.30 used CORAL software to build a QSAR model for the prediction of cellular uptake of this dataset. The software gave satisfactory and stable predictions of the cellular uptake of NPs in PaCa2 cancer cells for five random splits. Another attempt was made by Ghorbanzadeh et al.31 who presented an artificial neural network that was built based on descriptors calculated with Hyperchem program and Dragon. The results revealed the accuracy and reliability of the proposed model and moreover a sensitivity analysis indicated that the number of hydrogen-bond donor sites in the organic coating of a NP is the predominant factor responsible for cellular uptake. Moreover, Liu et al.32 proposed a robust Relevance Vector Machine (RVM) model built with nine descriptors, which demonstrated prediction accuracy as quantified by a 5-fold crossvalidated squared correlation coefficient. Ensemble learning based QNAR models for predicting the biological effects of this dataset were also constructed by Singh et al.33 based on simple structural descriptors and various statistical parameters suggested robustness of the model. Finally a recent attempt for the modeling of this dataset was reported by Kar et al.34 in their publication were a statistically significant regression - based QNAR model was developed using a PLS method and a small number of interpretable descriptors.

In this work we present a fully validated and predictive QNAR model that was developed based on Mold2 descriptors and the kNN algorithm. Our model was made publicly available through Enalos InSilicoNano platform (http://enalos.insilicotox.com/ QNAR PaCa2/), a web service developed with the aim to facilitate NPs design and evaluation. The user can draw a new structure, enter a SMILES notation or upload many structures in an sdf file. By the click of a button a prediction is made available together with a value that indicates if the structure can be tolerated by the model in terms of its domain of applicability. We have used our web service in a virtual screening framework mining PubChem database. We have successfully retrieved several potent inhibitors with the aim to prioritize compounds for screening. This online tool could be a useful aid for the decision making of both research groups and regulatory bodies interested in NPs' design and screening.

#### 2. Results and discussion

#### 2.1 **Building a KNIME QNAR workflow**

In this work we have tried to address the need of robust and predictive QNAR models for the assessment of the biological profile of ENPs and on top of that the proposed model has been made available online through Enalos InSilicoNano platform. The platform was used in a virtual screening framework to identify promising compounds within PubChem. For our study we have worked with 109 ENPs with the same metal core and different organic coating.25 The model was built based on a KNIME workflow that was developed for this purpose. KNIME is a freely available tool and has an extended community of users and developers and is increasingly gaining more attention for solving cheminformatics problems.35 Our efforts to address the lack of ready-to-use applications based on QSAR models were facilitated by the use of this open source platform.

Our overall strategy is targeting the development of a validated QNAR model and the release of this model to the wider community through a web service. For the model development a KNIME35 workflow was developed that executes the following procedures: (i) data preprocessing, (ii) descriptors calculation, (iii) variable selection and model development, (iv) model validation, (v) domain of applicability determination. In the proposed workflow all these computational steps were incorporated and this complete line of operations was made feasible with the invaluable help of our in house made Enalos KNIME nodes, namely Enalos Mold2 node, Enalos Model Acceptability Criteria node and Enalos Domain - Similarity node.36 These nodes have been developed by Novamechanics Ltd and are publicly available through the KNIME Community and the company's website.37

**2.1.1** Model development. To initiate our model development all data including organic structures and cellular uptake values were preprocessed and randomly partitioned into training and validation set. Among the 109 compounds originally included in the dataset 89 constituted the training set and 20 the test set.25 Only compounds included in the training set were used to develop the QNAR model whereas compounds included in the test set were not involved in the model

development. Since the organic coating differentiated the NPs we have encoded the organic structure using Mold2 descriptors. Enalos Mold2 KNIME node was used to calculate a number of 777 descriptors for each compound that account for their topological, geometric and structural characteristics. From this original pool of descriptors a number was removed as some of the descriptors do not have any discrimination power (no variation) and for this a node called 'Low Variance Filter' was applied. After removal of these descriptors, 382 descriptors remained and were used as possible inputs for the QNAR model development.

The CfsSubset variable selection with BestFirst evaluator method was then applied on the training data to select the most significant descriptors. Among the available descriptors, nine have emerged as the most critical in capturing the significant structural characteristics that affect the biological profile of the studied NPs as proposed by the variable selection algorithm. These descriptors include:

Geary topological structure autocorrelation length-7 weighted by atomic van der Waals volumes (D461), Geary topological structure autocorrelation length-5 weighted by atomic Sanderson electronegativities (D467), number of total quaternary C-sp3 (D599), number of group secondary amines (aliphatic) (D649), number of group donor atoms for H-bonds (with N and O) (D712), number of group CH3R and CH4 (D714), number of group phenol or enol or carboxyl OH (D753), number of group Al2–NH (D758) and hydrophilic factor index (D775). Their physical meaning is briefly described below.

Descriptors D461 and D467 encode information as described by Geary topological structure autocorrelation length-7 weighted by atomic van der Waals volumes and length-8 weighted by atomic Sanderson electronegativities. Geary index is a general index of spatial autocorrelation and is a distance-type function varying from zero to infinite. In each descriptor the index is either weighted by atomic van der Waals volumes or atomic Sanderson electronegativities. The hydrophilic factor index (D775) accounts for the hydrophilicity of each of the structures described. All other descriptors included are counting for the number of different important features present in the structure such as total quaternary C-sp3 (D599), secondary amines (D649), donor atoms for H-bonds (D712), the presence of CH3R and CH4 (D714), phenol or enol or carboxyl OH (D753) and Al2–NH (D758).

The proposed KNIME workflow gave us the opportunity to test the performance of various algorithms included in the WEKA suite of programs and select the combination that best describes our data. The kNN algorithm was selected to describe the significant correlation among the selected descriptors and the cellular uptake in PaCa2. This algorithm outperformed various different algorithms that were also tested. The kNN methodology was applied on our training data with an optimized value of k equal to 2.<sup>43</sup> Euclidean distance was used with all nine descriptors and contributions of neighbors weighted by the inverse of distance.

**2.1.2 Model validation – domain of applicability.** The proposed model was validated using the techniques mentioned in the Materials and methods section.<sup>44–46</sup> Our model was

Criterion	Assessment	Result
R^2 > 0.6	PASS	$R^2 = 0.848$
Rcvext^2 > 0.5	PASS	Rcvext^2 = 0.82
(R^2-R0^2)/R^2 < 0.1	PASS	(R^2-R0^2)/R^2 = 0.038
(R^2-R'0^2)/R^2 < 0.1	PASS	$(R^2-R^0^2)/R^2 = 0.0$
abs(R0^2-R'0^2) < 0.1	PASS	abs(R0^2-R'0^2) = 0.032
0.85 < k < 1.15	PASS	k = 1.019
0.85 < k' < 1.15	PASS	k' = 0.979
N	lodel Predictiv	re

Scheme 1 Model evaluation summary results.

internally and externally validated using various validation algorithms to assess its robustness and predictivity. Lessons learned from the long standing 'classic' QSAR modeling have to be taken into account from the very beginning when building a QNAR model. It is very important that principles recommended by OECD including robust validation of results are addressed when the modeling of NPs' biological profile is requested. The Enalos Model Acceptability Criteria KNIME node has been used for this purpose. The model successfully passed Tropsha's recommended tests for predictive ability as shown from the results below. Scheme 1 is a screenshot of the results as they are produced by Enalos Model Acceptability Criteria KNIME node.

 $R^2$  is the coefficient of determination between experimental values and model prediction on the test set ( $R^2_{\rm pred}$ ). Mathematical calculations of  $R^2_{\rm o}$ ,  $R'^2_{\rm o}$ , k, and k' are based on regression of the observed activities against the predicted activities and *vice versa* using the equations described in Materials and methods section.

The model was also quite stable to the inclusion–exclusion of compounds measured by the ten-fold cross validation procedure. The  $R_{\rm L10O}^2$  was calculated equal to 0.74. In addition the Y-randomization test was used as a method for testing the robustness and statistical significance of the model. Since low values of the correlation coefficient were measured we can eliminate the possibility of chance correlation.

The values of all the above statistical tests illustrate the accuracy, significance and robustness of the proposed model.

It is important that the limitations of the model are also described *via* the domain of applicability. This gives an important indication as the user can freely and creatively design novel molecules but will be warned for the reliability of the prediction when the structural characteristics cannot be tolerated by the model. After model validation, the domain of applicability of our model was also defined to ascertain that a given prediction can be considered reliable.<sup>47–50</sup> The applicability domain limit value was defined equal to 2.153 based on the equation provided in Materials and methods section. All compounds in the test set had values in the range of 0.019–1.06 except for one which slightly falls outside with a value of 2.29. The predictions for all compounds that fell inside the domain of applicability of the model can be considered reliable.

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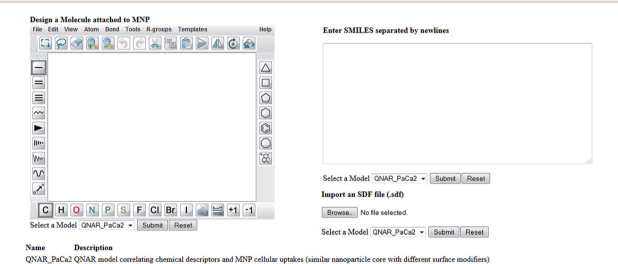
Our proposed model requires only structural information from the small organic molecules involved and was proven accurate and reliable for given applicability limits. Thus our model could be used as a useful aid to the costly and time consuming experiments for determining cellular uptake of NPs and could further be used to screen existing databases or virtual chemical structures to identify NPs with desired properties. In this effort, the applicability domain will play an important role as it will filter out chemical structures that could not be tolerated by the model.

#### **Building Enalos InSilicoNano platform**

An important aspect that, in the vast majority of examples presented in literature, remains forgotten is the dissemination

of the results to the wider community. It is very crucial that the developed model does not remain within the developers' group but is widely disseminated to the community so that it could immediately serve as an important source of information as it was initially designed to be. Moreover as recently highlighted by many initiatives it is crucial that this is done with open source tools that could be easily expanded and adjusted to the special needs of each project.

To enable its role and make the model predictions available to the interested users, our proposed model was made publicly available online through Enalos InSilicoNano platform.<sup>51</sup> Enalos InSilicoNano platform is a webservice that can host several validated and predictive models that can be utilized in the NPs design process. Our validated model was made publicly



Scheme 2 Screen shot of Enalos InSilicoNano platform input page.

## Prediction of MNPs Uptake in PaCa2 Cancer Cells

Knime report powered by Birt

"PaCa2 cellular uptake (log10 [nanoparticles]/cell p	M)" "Domain of Applicability Prediction"
4.429	reliable
4.293	reliable
3.494	reliable
4.197	reliable
3.895	reliable
3.895	reliable
4.197	reliable
4.197	reliable
3.895	reliable
4.197	reliable
3.523	reliable
3.495	reliable
4.209	reliable
3.895	reliable
4.283	reliable
Date: Jun 5, 2014 6:07 AM Auth	or: NovaMechanics Ltd 1 of 1

Scheme 3 Screen shot of Enalos InSilicoNano platform results.

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Table 1 Compounds included in our dataset with experimental and predicted values

2 FC(F)( 3 FC(F)( 4 CC1(C) 5 O=C1 6 CC1= 7 CC1= 8 CCCC 9 CC1C(C) 10 O=C1 11 O=C1 11 O=C1 12 Bre1cc(I) 13 O=C1 14 Fc1c(II 15 O=C1 16 Oc1cc 17 O=C1 18 Clc1cc 19 O=C1 20 ClC1= 21 CC(= 22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 28 O=C1 31 Cc1cc 32 O=C1 31 Cc1cc 32 O=C1 33 O=C1 34 O=C1 35 O=C1 37 CN1C 38 O=C1 38 O=C1 38 O=C1 40 CC(= 41 Bre1c( 42 O=C1 42 O=C1 44 CC(= 44 CC(C) 45 CC(C) 46 ClC1= 47 CCCC 48 Nc1cc 49 CCCC 48 Nc1cc 49 CCCC 48 Nc1cc 49 CCCC 50 O=C1 51 O=C1 52 O=C1 51 O=C1 52 O=C1		uptake (log 10 [NP]/cell)	Predicted PaCa2 cellular uptake (log 10 [NP]/cell)
2 FC(F)(3) 3 FC(F)(4) 4 CC1(C) 5 O=C1 6 CC1= 7 CC1= 8 CCCC 9 CC1C(C) 10 O=C1 11 O=C1 1	F)(F)C(=O)OC(=O)C(F)(F)F	4.17	4.17
3 FC(F)( 4 CC1(C) 5 O=C1 6 CC1= 7 CC1= 8 CCC0 9 CC1C0 10 O=C1 11 O=C1 12 Bretec 13 O=C1 14 Fc1c(I 15 O=C1 16 Oc1ec 17 O=C1 18 Clc1cc 17 O=C1 18 Clc1cc 17 O=C1 18 Clc1cc 17 O=C1 18 Clc1cc 19 O=C1 20 ClC1= 21 CCC= 22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 28 O=C1 29 CCC0 30 OC(= 31 Cc1cc 32 O=C1 33 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 CN1C 38 O=C1 38 O=C1 38 O=C1 40 CC(= 41 Bretec 42 O=C1 44 ICC(= 44 ICC(= 45 ClC1 45 ClC1 46 ClC1= 47 CCCC 48 Nc1cc 48 Nc1cc 49 CCCC 48 Nc1cc 50 O=C1 51 O=C1 51 O=C1 51 O=C1 52 O=C1	F)(Cl)C(=O)OC(=O)C(F)(F)Cl	3.95	3.95
4 CC1(C) 5 O=C1 6a CC1= 7 CC1= 8 CCC0 9 CC1C0 10 O=C1 11 O=C1 12 Bretec 13 O=C1 14 Fete(II 15 O=C1 16 Octec 17 O=C1 18 Clctec 17 O=C1 18 Clctec 17 O=C1 18 Clctec 19 O=C1 20 ClCt= 22 Clctec 23 O=C1 24 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 28 O=C1 29 CCC0 30a OC(= 31 Cctcc 32 O=C1 33 O=C1 33 O=C1 33 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 CN1C0 38 O=C1 38 O=C1 40 CC(= 41a Bretec 44a O=C1 44b CC(= 44c O=C1 44c O=C1 44c CCC 44c CCC 45 CCCC 46 CCCC 47 O=C1 48 Netec 49a CCCC 48 Netec 48a Netec 49a CCCC 48a Netec 49a CCCC 48a Netec 49a CCCC 48a Netec 49a CCCCC 48a Netec 49a CCCC 48a Netec 48a CCCCC	F)(F)C(F)(F)C(=O)OC(=O)C(F)(F)C(F)(F)F	4.08	4.08
5	(C)CC(=O)OC1=O	4.11	3.80
6 <sup>a</sup> CC1= 7 CC1= 8 CCC0 9 CC1C0 10 O=C1 11 O=C1 12 O=C1 13 O=C1 14 Fc1c[I 15 O=C1 16 Oc1cc 17 O=C1 18 Clc1cc 19 O=C1 20 Clc1cc 21 Clc1cc 22 Clc1cc 22 Clc1cc 22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 29 CCC0 30 <sup>a</sup> OC(= 31 Cc1cc 32 O=C1 33 O=C1 33 O=C1 33 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 CN1C 38 O=C1 38 O=C1 38 O=C1 40 CC(= 41 <sup>a</sup> Brc1c( 44 <sup>a</sup> O=C1 44 <sup>a</sup> O=C1 45 ClCC(= 44 <sup>a</sup> CCCC 48 Nc1cc 48 Nc1cc 49 <sup>a</sup> CCCC 48 Nc1cc 50 <sup>a</sup> O=C1 51 O=C1 52 O=C1	C1OC(=0)C=C1	3.98	4.11
7	=CC(=0)OC1=O	3.58	3.65
8	=C(C)C(=O)OC1=O	3.48	3.80
9	CCCC(=O)OC(=O)CCCCC	3.65	3.65
10	CC(=0)OC1=0	3.64	3.65
11	C1OC(=O)c2cc(ccc12)C(=O)c1ccc2C(=O)OC(=O)c2c1	3.51	3.53
12 <sup>a</sup> Bre1cc 13 O=C1 14 Fc1c[I 15 O=C1 16 Oc1cc 17 O=C1 18 Clc1cc 19 O=C1 20 ClC1= 21 <sup>a</sup> CC(= 22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 29 CCC 30 <sup>a</sup> OC(= 31 Cc1cc 32 O=C1 33 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 CN1C 38 O=C1 38 O=C1 40 CC(= 40 CC(= 41 <sup>a</sup> Bre1c(= 42 <sup>a</sup> O=C1 43 O=C1 44 ICC(= 44 ICC(= 45 ClCC(= 46 ClC1= 47 <sup>a</sup> CCCC 48 Nc1cc 48 Nc1cc 49 <sup>a</sup> CCCC 48 Nc1cc 49 <sup>a</sup> CCCC 48 Nc1cc 50 <sup>a</sup> O=C1 51 O=C1 52 O=C1	C1OC(=O)c2cc(ccc12)N(=O)=O	3.27	3.29
13	lccc2C(=O)OC(=O)c3cccc1c23	3.63	3.52
14 Fc1c(I 15 O=C1 16 Oc1cc 17 O=C1 18 Clc1cc 19 O=C1 20 ClC1= 21 CC(= 22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 29 CCC 30 OC(= 31 Cc1cc 31 O=C1 32 O=C1 33 O=C1 33 O=C1 34 O=C1 33 O=C1 34 O=C1 35 O=C1 40 CC(= 40 CC(= 41	C1OC(=O)c2ccc3C(=O)OC(=O)c4ccc1c2c34	3.67	3.68
15 O=C1 16 Octoc 17 O=C1 18 Clctcc 19 O=C1 20 ClCt= 21 CC(= 22 Clctcc 23 O=C1 24 O=C1 25 O=C1 27 O=C1 28 O=C1 29 CCC 30 OC(= 31 Cctcc 31 O=C1 33 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 O=C1 38 O=C1 40 CC(= 40 CC(= 41	c(F)c(F)c2C(=O)OC(=O)c2c1F	3.83	3.84
16 Octoc 17 O=C1 18 Clc1cc 19 O=C1 20 ClC1= 21 CC(= 22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 29 CCC 30 OC(= 31 Cc1cc 32 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 CN1C 38 CC1Cc 38 O=C1 40 CC(= 41 Brc1c( 42 O=C1 43 O=C1 44 ICC(= 45 ClCC( 46 ClC1= 47 CCCC 48 Nc1cc 49 CCCC 48 Nc1cc 49 CCCC 50 O=C1 51 O=C1 51 O=C1	C1OC(=O)c2cc(cc3cccc1c23)N(=O)=O	4.11	4.09
17	cccc2C(=O)OC(=O)c12	3.97	3.97
18	C1OC(=O)C2C3CCC(C=C3)C12	3.9	3.87
19	.ccc2NC(=0)OC(=0)c2c1	4.18	4.17
20	C1OS(=O)(=O)c2cccc12	3.88	3.93
21a         CC(=           22         Clc1cc           23         O=C1           24         O=C1           25         O=C1           26         O=C1           27         O=C1           28         O=C1           30a         OC(=           31         Cc1cc           32         O=C1           33         O=C1           34         O=C1           35a         O=C1           37         CN1C           38a         CC1C0           39         O=C1           39         O=C1           40         CC(=           41a         Brc1c(           42a         O=C1           43         O=C1           44         ICC(=           45         ClCC(           46         ClC1=           47a         CCCC           48         Nc1cc           49a         CCCC           50a         O=C1           51         O=C1           52         O=C1			
22 Clc1cc 23 O=C1 24 O=C1 25 O=C1 25 O=C1 26 O=C1 27 O=C1 28 O=C1 30 O=C1 31 Cc1cc 32 O=C1 33 O=C1 34 O=C1 35 O=C1 36 O=C1 37 CN1C 38 CC1CC 40 CC(= 41 Brc1c( 44 ICC(= 44 ICC(= 44 ICC(= 45 ClCC( 48 Nc1cc 49 CCCC 48 Nc1cc 49 CCCC 48 Nc1cc 49 CCCC 48 Nc1cc 49 CCCC 48 Nc1cc 50 O=C1 51 O=C1 52 O=C1	1=C(Cl)C(=0)OC1=0 =0)SC1CC(=0)OC1=0	3.84	3.87
23		3.59	3.85
24	.cc2C(=0)OC(=0)c2cc1Cl	4.12	4.07
25 O=Ci 26 O=Ci 27 O=Ci 27 O=Ci 28 O=Ci 29 CCCC 30° OC(= 31 Cc1cc 32 O=Ci 33 O=Ci 34 O=Ci 35° O=Ci 37 CN1Ci 38° CC1Cc 39 O=Ci 40 CC(= 41° Brc1c( 42° O=Ci 44 ICC(= 44° CCCC 48 Nc1cc 49° CCCC 48 Nc1cc 50° O=Ci 51 O=Ci 52 O=Ci 51 O=Ci 52 O=Ci 51 O=Ci 52 O=Ci 54 O=Ci 55 O=Ci 56 O=Ci 57 O=C	C1OC(=0)C2C3OC(C=C3)C12	3.82	3.80
26	C1OC(=0)C2C3C=CC(C12)C1C3C(=0)OC1=O	3.63	3.65
27	C1OC(=0)C2CC=CCC12	3.89	3.86
28 $O = C1$ 29 $CCCC$ $30^a$ $OC(=$ 31 $Cc1cc$ 32 $O = C1$ 33 $O = C1$ 34 $O = C1$ 35 $^a$ $O = C1$ 37 $CN1C$ 39 $O = C1$ 40 $CC(=$ 41 $^a$ $Bre1c($ 42 $^a$ $O = C1$ 43 $O = C1$ 44 $ICC(=$ 45 $CICC(=$ 46 $CIC1=$ 47 $^a$ $CCCC$ 48 $Nc1cc$ 49 $^a$ $CCCC$ 50 $^a$ $O = C1$ 51 $O = C1$ 52 $O = C1$	C1OC(=0)c2cccc2-c2cccc12	3.77	3.77
29       CCCC $30^a$ OC(=         31       Cc1cc         32       O=C1         33       O=C1         34       O=C1         35 $^a$ O=C1         36       O=C1         38 $^a$ CC1C0         39       O=C1         40       CC(=         41 $^a$ Bre1c(         42 $^a$ O=C1         43       O=C1         44       ICC(=         45       CICC(         46       CIC1=         47 $^a$ CCCC         48       Nc1cc         49 $^a$ CCCC         50 $^a$ O=C1         51       O=C1         52       O=C1	C1OC(=O)c2ccc(c3cccc1c23)N(=O)=O	3.93	3.92
30 <sup>a</sup> OC(= 31 Cc1cc 32 O=C1 33 O=C1 33 O=C1 34 O=C1 35 <sup>a</sup> O=C1 36 O=C1 37 CN1C 38 <sup>a</sup> CC1Cc 39 O=C1 40 CC(= 41 <sup>a</sup> Bre1c( 42 <sup>a</sup> O=C1 43 O=C1 44 ICC(= 45 ClCC( 46 ClC1= 47 <sup>a</sup> CCCC 48 Nc1cc 49 <sup>a</sup> CCCC 50 <sup>a</sup> O=C1 51 O=C1 52 O=C1	C1OC(=0)C2C1C1C2C(=0)OC1=O	3.77	3.86
31 Celece 32 O=Cl 33 O=Cl 33 O=Cl 34 O=Cl 35a O=Cl 36 O=Cl 37 CN1Cl 38a CC1Cl 39 O=Cl 40 CC(= 41a Brele( 42a O=Cl 43 O=Cl 44 ICC(= 45 ClCC( 46 ClCl= 47a CCCC 48 Nelece 49a CCCC 50a O=Cl 51 O=Cl 52 O=Cl	CCCCCCCC(=0)OC(=0)CCCCCCCCC	3.82	3.82
32 $O = C1$ 33 $O = C1$ 34 $O = C1$ 35a $O = C1$ 36 $O = C1$ 37 $CN1C$ 39 $O = C1$ 40 $CC(=$ 41a $Bre1c(=$ 42a $O = C1$ 43 $O = C1$ 44 $ICC(=$ 45 $CICC(=$ 46 $CIC1=$ 47a $CCCC$ 48 $Nc1cc$ 49a $CCCC$ 50a $O = C1$ 51 $O = C1$ 52 $O = C1$	=O)c1ccc2C(=O)OC(=O)c2c1	3.55	3.62
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	ccc2C(=O)OC(=O)c2c1	3.98	3.97
$34$ O=C1 $35^a$ O=C1 $36$ O=C1 $36$ O=C1 $37$ CN1C1 $38^a$ CC1C0 $39$ O=C1 $40$ CC(= $41^a$ Bre1c( $42^a$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^a$ CCCC $48$ Ne1ce $49^a$ CCCC $50^a$ O=C1 $51$ O=C1 $52$ O=C1	C1OC(=O)c2c1cccc2N(=O)=O	3.5	3.54
$35^{a}$ O=C1 $36$ O=C2 $36$ O=C1 $37$ CN1C1 $38^{a}$ CC1C0 $39$ O=C1 $40$ CC(= $41^{a}$ Bre1c( $42^{a}$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^{a}$ CCCC $48$ Ne1ce $49^{a}$ CCCC $50^{a}$ O=C1 $51$ O=C1 $52$ O=C1	C1Cc2cccc2C(=O)O1	3.78	3.81
$36$ O=C1 $37$ CN1C1 $38^a$ CC1C0 $39$ O=C1 $40$ CC(= $41^a$ Brc1c( $42^a$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^a$ CCCC $48$ Nc1cc $49^a$ CCCC $50^a$ O=C1 $51$ O=C1 $52$ O=C1	C1CCCC(=0)O1	4.07	4.06
$37$ CN1C $38^a$ CC1C( $39$ O=C1 $40$ CC(= $41^a$ Brc1c( $42^a$ O=C1 $43$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^a$ CCCC $48$ Nc1cc $49^a$ CCCC $50^a$ O=C1 $51$ O=C1 $52$ O=C1	C1CN(CCN2CC(=0)OC(=0)C2)CC(=0)O1	3.93	3.76
$38^{a}$ CC1C0 $39$ O=C1 $40$ CC(= $41^{a}$ Brc1c( $42^{a}$ O=C1 $43$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^{a}$ CCCC $48$ Nc1cc $49^{a}$ CCCC $50^{a}$ O=C1 $51$ O=C1 $52$ O=C1	C1Nc2cccc2C(=O)O1	4.44	4.43
$39$ O=C1 $40$ CC(= $41^a$ Bre1c( $42^a$ O=C1 $43$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^a$ CCCC $48$ Ne1ce $49^a$ CCCC $50^a$ O=C1 $51$ O=C1 $52$ O=C1	C(=O)OC(=O)c2ccccc12	3.36	3.38
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	CC(=O)OC(=O)C1	3.91	3.68
$41^{a}$ Bre1c( $42^{a}$ O=C1 $43$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^{a}$ CCCC $48$ Ne1ce $49^{a}$ CCCC $50^{a}$ O=C1 $51$ O=C1 $52$ O=C1	C1OC(=O)C2=C1CCCC2	3.73	3.74
$42^{a}$ O=C1 $43$ O=C1 $44$ ICC(= $45$ ClCC( $46$ ClC1= $47^{a}$ CCCC $48$ Ne1ce $49^{a}$ CCCC $50^{a}$ O=C1 $51$ O=C1 $52$ O=C1	=O)OC1C(OC(C) $=$ O)C( $=$ O)OC1 $=$ O	3.91	3.91
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	c(Br)c(Br)c2C(=O)OC(=O)c2c1Br	3.8	3.66
44 ICC(= 45 ClCC(= 46 ClC1= 47 <sup>a</sup> CCCC 48 Nc1cc 49 <sup>a</sup> CCCC 50 <sup>a</sup> O=C1 51 O=C1 52 O=C1	C1OC(=O)C2CCCCC12	3.93	3.88
$45$ ClCC( $\frac{1}{46}$ ClC1= $\frac{1}{47^a}$ CCCC $\frac{1}{48}$ Nc1cc $\frac{1}{49^a}$ CCCC $\frac{1}{50^a}$ O=C1 $\frac{1}{52}$ O=C1	C1OC(=O)C2=C1CCC2	3.69	3.71
$ 46 $ $ ClC1 = 47^{a} $ $ CCCC $ $ 48 $ $ Nc1cc $ $ 49^{a} $ $ CCCC $ $ 50^{a} $ $ 0 = C1 $ $ 51 $ $ 0 = C2 $ $ 52 $ $ 0 = C1 $	(=O)OC(=O)CI	3.42	3.42
$47^{a}$ CCCC $48$ Netce $49^{a}$ CCCC $50^{a}$ O=C1 $51$ O=C1 $52$ O=C1	C(=O)OC(=O)CCI	3.63	3.62
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	1 = C(Cl)C2(Cl)C3C(C(=O)OC3=O)C1(Cl)C2(Cl)Cl	3.47	3.49
$49^{a}$ CCCC $50^{a}$ O=C1 $0$ =C1 $0$	CCCCCCCCCCCC(=0)OC(=0)CCCCCCCCCCCCCC	3.55	3.78
50 <sup>a</sup> O=C1 51 O=C1 52 O=C1	ccc2C(=O)OC(=O)c3cccc1c23	3.64	3.63
50 <sup>a</sup> O=C1 51 O=C1 52 O=C1	CCCCCCC(=O)OC(=O)CCCCCCCC	4.03	3.78
51 O=C1 52 O=C1	C1CC2(CCCC2)CC(=0)O1	4.06	3.88
52 O=C1	C1OC(=O)C2C3CCC(C3)C12	3.94	3.91
	C1OC(=O)c2cccc3cccc1c23	3.96	3.95
53 O=C1	C1CCC(C(=0)O1)c1ccccc1	4.02	4.00
	c(Cl)c(Cl)c2C(=O)OC(=O)c2c1Cl	3.83	3.66
,	.ccc(Cl)c2C(=O)OC(=O)c12	3.9	3.88
	(C)CCC(=0)OC1=O	3.94	3.80
57 CCCC		3.78	3.78
	C)CC(C)N	3.85	3.85
	.C(O)CC(CO)C(O)C1O	3.36	3.36
	CCCCN	3.75	3.77
	C)(C)N	3.86	3.87
62 CC(C)		3.72	3.74

Table 1 (Contd.)

ID	Smiles	Observed PaCa2 cellular uptake (log 10 [NP]/cell)	Predicted PaCa2 cellular uptake (log 10 [NP]/cell)
63	CC(C)(C)CN	3.75	3.91
64	CC(C)CCN	3.83	3.82
65	CCC(N)CC	3.81	3.82
66	CCC(C)(C)N	4.07	3.91
67	NCCN	3.46	3.46
68	CCCCCCCCCCCCN	4.06	4.03
69	NCCCN	3.49	3.49
70	NCCCCN	3.48	3.48
71	NCCCCCCN	3.62	3.62
72	CCCCC(CC)CN	3.95	3.94
73	CCCCCCCCCCCCN	3.97	4.00
74	CCCCC(C)N	3.63	3.64
75 <sup>a</sup>	CCCCCCCCCCCN	4.27	4.02
76 76	NCCNCCN	3.77	3.77
77	NCC12CC3CC(CC(C3)C1)C2	2.84	2.87
78	NCCc1ccc(O)c(O)c1	2.53	2.53
79	NCCc1ccc(O)cc1	2.77	2.77
$80^a$	NCCCNCCCNCCN	2.41	2.37
81	NCCNCCCNCCN	2.23	2.24
82	NCCNCCNCCNCCN	2.54	2.54
83	NC12CC3CC(CC1C3)C2	3.12	3.14
84	NC1C2CC3CC(C2)CC1C3	3.18	3.15
85	NCC(0)=0	2.57	2.58
86	COC(=O)C(N)Cc1ccccc1	3.39	3.39
87	NC(CO)C(O)=O	3.36	3.35
		3.21	
88	CC(O)C(N)C(O)=O $NC(CO1c[n]U[O]=O(O)=O$		3.21
89	NC(Cc1c[nH]c2cccc12)C(O)=O	3.19	3.19
90 91 <sup>a</sup>	NC(Cc1ccc(O)cc1)C(O)=O	3.07	3.07
	CC(C)C(N)C(O)=O	3.27	3.04
92	NCCCCC(N)C(O) = O	3.25	3.25
93	NC(C(O)=O)c1ccc(Cl)cc1	3.06	3.07
94 95 <sup>a</sup>	CC(N)C(O)=O	2.9	2.90
	NC(CCCNC(N)=N)C(O)=O	3.15	3.28
96 <sup>a</sup>	NC(CC(0)=0)C(0)=0	3.29	3.35
97	NC(CCC(N)=O)C(O)=O	3.32	3.32
98	NC(CCC(0)=0)C(0)=0	3.4	3.40
99	NC(Cc1c[nH]cn1)C(O)=O	3.38	3.38
100	CSCCC(N)C(O) = O	3.23	3.23
101	NC(Cc1cccc1)C(O)=O	3.29	3.29
102	O=C1CCC(=O)O1	4.24	4.11
103 <sup>a</sup>	CC(=0)OC(C)=0	4.05	3.80
104	C=C1CC(=0)OC1=O	4.04	4.06
105	O=C1COCC(=0)01	3.99	3.96
106	O=C1OC(=O)c2cccc12	3.9	3.92
107	OC(=0)CC1CC(=0)OC1=0	4.03	4.03
108	Fc1ccc(F)c2C(=O)OC(=O)c12	3.91	3.87
109	OC(=O)CN(CCN1CC(=O)OC(=O)C1)CCN1CC(=O)OC(=O)C1	4.1	4.09

available through this platform and can thus be of help to the wider community of end users interested in NP's design. The web service needs no special computational skills and can be easily used by different groups of scientists like chemists, biologists *etc.* or even non experts involved or interested in the NPs biological evaluation.

Enalos InSilicoNano platform has a user friendly interface with minimum steps required and no authentication and authorization procedure. To initiate a prediction the user must first select the model of interest from the drop down menu provided. When the model "QNAR\_PaCa2" is selected the prediction can be initiated when a structure or a batch of structures is uploaded. For that the web service provides three different options described as follows: (i) the user draws a chemical structure of interest using the drawing tool. The user can easily select from the different panels the atoms, bonds or substructures of interest and construct the molecule. What is important is that the user can also open, save and convert files

Table 2 Virtual screening results for the most promising compounds in PubChem database

ID	Compound	Predicted value PaCa2 cellular uptake (log 10 [NP]/cell)	Domain of applicability (limit: 2.153)
679	HN	4.41	0.03
604	F H N N	4.41	0.01
958		4.41	0.06
676	HN I	4.40	0.01
678	HN	4.40	0.05
677	HN N O	4.39	0.02
107	N H	4.39	0.02
368	NH NH	4.38	0.10
293	Br	4.37	0.09
493	HN HN	4.37	0.10
494	HN N	4.37	0.10
550	HN	4.36	0.11
196		4.35	0.06

Table 2 (Contd.)

ID	Compound	Predicted value PaCa2 cellular uptake (log 10 [NP]/cell)	Domain of applicability (limit: 2.153)
200		4.35	0.06
626	T N N	4.35	0.06
602	F N N	4.35	0.06
981	o H	4.34	0.10
925		4.34	0.10
192	Br O	4.34	0.06
65	O N	4.34	0.05

with a variety of chemical formats (i.e. SMILES, IUPAC chemical Identifier, MDL MOL file) using the drop down menu of the online sketcher; (ii) the user enters the SMILES notation of a structure or several structures separated by newlines. Even if the SMILES notation is not initially known it is important that the chemical sketcher included gives the users the opportunity to design the chemical structure and then copy the structure as SMILES from the Edit drop down menu. This is very significant as it facilitates the generation of several structures since the user can make several modifications using the sketcher and copy all structures as SMILES so that a prediction for the whole set of produced structures is generated. The user can thus visualize the modifications and make multiple predictions at once; (iii) the user can select and import an SDF file (.sdf) with several structures.

When structures are uploaded in either way a prediction can be generated by clicking the submit button. The output is then presented in a different html page. The results include the predicted value for each structure entered and an indication of whether the prediction could be considered reliable based on the domain of applicability of the model. A screen shot of the web service and the results page are presented in the following schemes.

Our developed KNIME workflow integrated with Enalos InSilicoNano web service made the online prediction of the biological effects of NPs feasible. In the web service presented in Scheme 2 and 3, the user can design or enter a chemical structure and get the prediction. The workflow behind the interface calculates the descriptors and generates the output. It is important that the output will appear on screen within seconds. The user can experiment with different scaffolds and substituents and study the structural characteristics that are responsible to induce a certain effect. The user can take advantage of the proposed QNAR model and immediately scan the structures of interest for a preliminary in silico testing. In this way we overcome a main point of controversy for QSAR models in general, that their results are not available for sharing and implementation. As recently highlighted<sup>52</sup> the advantages of making models available for use as software tools will increase in the future and this will enable the re-use of knowledge and will boost further developments. Enalos InSilicoNano platform uses a pipeline tool, KNIME, to address exactly this need of using and testing the models directly available on the web. With this platform we aim to address the need to reduce the amount of time spent by scientists in referencing disparate sources of data to aid decision making related to NPs design and bioactivity profile. Enalos InSilico-Nano is launched as an efficient port where models can be developed and published directly on the web using a user friendly interface.

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#### 2.3 Virtual screening

The presented model and web service can be used in a virtual screening framework for the prioritization of novel potent compounds.53,54 To demonstrate the usefulness of the Enalos InSilicoNano platform we have identified novel potent structures within PubChem database using similarity measurements based on Molecular Quantum Numbers (MQNs) as described in Materials and methods section.55 For this purpose we have selected among millions of compounds included in the Pub-Chem<sup>56</sup> database the most similar to the most active compound included in our initial dataset, that is compound 36 (isatoic anhydride). The above virtual screening procedure was used for the identification of the first 1000 neighbours of this most active compound included in our dataset in terms of chemical similarity. These first 1000 compounds within the PubChem that were identified as the most similar in the chemical space were evaluated using Enalos InSilicoNano platform for the assessment of their cellular uptake. For this purpose we have uploaded the sdf file containing all the proposed structures to Enalos InSilicoNano platform web service and asked for a prediction. Compounds were then sorted by their increasing potency and the most promising compounds were proposed for screening. The predicted values of the first 20 prioritized compounds are shown in Table 2.

Within this proposed strategy Enalos InSilicoNano platform emerges as a key component for evaluating novel nano-structures that have not been experimentally evaluated or even synthesized. It is also important to highlight that our proposed methodology and tools can also be expanded and applied to polymer–nanoparticle composites that are now gaining increasing attention.

We have succeeded to generate a novel computational activity assessment platform for nanoparticles by integrating two open science platforms: KNIME that combines a rich graphical workflow environment for integration of diverse analytics and Enalos InSilicoNano a platform for hosting and publishing models directly on the web allowing the researchers to do virtual screening and/or design of novel nanoparticles. Two milestones have been reached within this work, the first is the development of a validated QNAR model and the second is the development of a web service that will immediately give the opportunity of exploiting the model's results. To demonstrate the usefulness of the model we have also proposed a virtual screening framework that could be used to identify novel potent structures.

#### 3. Conclusions

In summary our goal in this work was dual, firstly to build a robust validated QNAR model for NPs and secondly to give immediate access to our model and results using an open access web service. Enalos InSilicoNano platform aspires to be a useful tool for design of novel NPs with desired properties. <sup>57,58</sup> To this end we have successfully built and validated a QNAR model that can reliably predict the cellular uptake of a dataset of 109 NPs. The model was made publicly available through

Enalos InSilicoNano platform and can be used for the predictions of cellular uptake of new structures that are designed or imported to the server. This online tool was successfully used for the virtual screening of a set of structures within PubChem database that were selected based on MQN descriptors to identify compounds similar to a known active structure. Besides the significance of the validated proposed model, to the best of our knowledge this is the first attempt to develop an online tool for the wider scientific community to use in the computer aided nanoparticles design with desired properties. Our model can now be directly used and easily applied to facilitate the special requirements of the user.

#### 4. Materials and methods

#### 4.1 KNIME workflow development

To address the needs for our project we have used the powerful KNIME (Konstanz Information Miner) which is an open source tool for data analysis that allows data integration, processing, analysis, and exploration and enables the user to visually create data flows selectively execute some or all analysis steps, and later investigate the results through interactive views on data and models.35 In this work we have used KNIME platform in order to integrate all the various components of our workflow as described below in details. Within our workflow we have also incorporated our newly developed Enalos KNIME Nodes that were used to facilitate our model development and validation.<sup>36</sup> Enalos KNIME nodes include among others: (1) Enalos Mold2 node for the calculation of Mold2 molecular descriptors, (2) Enalos Model Acceptability Criteria node that can be used to validate the Quality of Fit and Predictive Ability of a continuous QSAR Model and (3) Enalos Domain - Similarity node that can be used to define applicability domain (APD) based on the Euclidean distances. The Enalos KNIME Nodes are freely available via the KNIME Community and the company's website.37

**4.1.1 Data set.** The data set consists of 109 magneto-fluorescent NPs that have the same metal core decorated with different synthetic small molecules. Experimental values of cellular uptake in PaCa2 for each NP included are reported in literature and expressed as the decadic logarithm of the concentration (pM) of NP per cell with values ranging from 2.23 to 4.44.<sup>25</sup> SMILES notation of the organic surface modifier as well as the corresponding experimental values are given in Table 1.

**4.1.2 Molecular descriptors.** It is well known that for a successful QNAR development, descriptors that assess the structural characteristics of compounds involved are of outmost importance. As stated before this study involves NPs with the same metal core but different organic molecules as surface modifiers and thus we have chosen to encode the structural characteristics of these organic modifiers that change among the dataset. For this purpose we have included in our workflow Mold2 software developed by the National Center for Toxicological Research of FDA that has been previously used with great success in other applications.<sup>38</sup> Mold2 calculates a large and

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diverse set of molecular descriptors encoding two-dimensional chemical structure information.

Within our KNIME workflow we have included Enalos Mold2 KNIME node<sup>36</sup> that is able to calculate a number of 777 descriptors that account for the topological, geometric and structural characteristics of the small molecules. From this original pool of descriptors a number was removed as some of the descriptors do not have any discrimination power (no variation) and for this a node called 'Low Variance Filter' was applied.<sup>39</sup>

4.1.3 Model development. A variable selection method was first used to select the most important variables among the set of originally determined descriptors. Correlation - based feature subset selection (CfsSubset) variable selection combined with BestFirst evaluator were chosen to evaluate the most critical parameters. 40,41 CfsSubset algorithm evaluates the worth of a subset of attributes by considering the individual predictive ability of each feature along with the degree of redundancy between them. Subsets of features that were highly correlated with the class while having low inter-correlation were preferred. BestFirst evaluator searches the space of attribute subsets by greedy hillclimbing augmented with a backtracking facility. Setting the number of consecutive non-improving nodes allowed controls the level of backtracking done. Best first may start with the empty set of attributes and search forward, or start with the full set of attributes and search backward, or start at any point and search in both directions (by considering all possible single attribute additions and deletions at a given point). A forward search has been chosen for this work.

Subsequently a machine learning method that could best model the available dataset was applied. We have thus incorporated in our KNIME workflow k-nearest neighbors (kNN) methodology. ANN methodology belongs to instance-based (or lazy) learning that classifies objects based on the closest training examples in the feature space. An object is classified by a majority vote of its neighbors, with the object being assigned to the class most common amongst its k nearest neighbors (a positive integer, typically small). For our dataset we have used an optimal k value and Euclidean distance with all descriptors and contributions of neighbors weighted by the inverse of distance.

**4.1.4 Model validation.** Our developed model was fully validated in accordance to the principals of model validation for accepting QSAR models as described by the Organization for Economic Cooperation and Development (OECD).<sup>44</sup> Our model was both internally and externally validated as represented by goodness-of-fit, robustness and predictivity.

For external validation the partitioning KNIME node was applied and the dataset was separated into training and validation set leaving a number of 20 compounds for the external validation of the model. All compounds included in the test set were not involved by any means in the training procedure.

To evaluate the models performance the following statistical criteria were used: the coefficient of determination between experimental values and model predictions ( $R^2$ ), validation through an external test set, leave-many-out cross validation procedure and Quality of Fit and Predictive Ability of a

continuous QSAR Model according to Tropsha's tests.<sup>45,46</sup> The latter was made feasible by including Enalos Model Acceptability Criteria node in our workflow.

In particular the formulas for calculating Tropsha's tests<sup>45</sup> are given below:

$$R_{\text{cvext}}^2 = 1 - \frac{\sum_{i=1}^{\text{ntest}} (y_i - \tilde{y}_i)^2}{\sum_{i=1}^{\text{ntest}} (y_i - \overline{y}_{\text{tr}})^2}$$
(1)

$$k = \frac{\sum_{i=1}^{ntest} y_i \tilde{y}_i}{\sum_{i=1}^{ntest} \tilde{y}_i^2}$$
 (2)

$$R_{o}^{2} = 1 - \frac{\sum_{i=1}^{ntest} (\tilde{y}_{i} - \tilde{y}_{i}^{ro})^{2}}{\sum_{i=1}^{ntest} (\tilde{y}_{i} - \overline{\tilde{y}})^{2}}, \text{ where } \tilde{y}_{i}^{ro} = ky_{i}, \quad i = 1, ..., \text{ ntest}$$
(3)

In the above equation *ntest* is the number of compounds that constitute the validation data set,  $\bar{y}_{tr}$  is the averaged value for the dependent variable for the training set,  $y_i$ ,  $\tilde{y}$ , i=1,..., ntest are the measured values and the QSAR model predictions of the dependent variable over the available validation set and  $\bar{y}$  is the average over all  $\tilde{y}$ , i=1,..., ntest.

Tropsha *et al.*<sup>45</sup> considered a QSAR model predictive, if the following conditions are satisfied:

$$R_{\text{cvext}}^2 > 0.5 \tag{4}$$

$$R_{\rm pred}^2 > 0.6 \tag{5}$$

$$\frac{\left(R_{\text{pred}}^2 - R_{\text{o}}^2\right)}{R_{\text{pred}}^2} < 0.1 \tag{6}$$

$$0.85 \le k \le 1.15 \tag{7}$$

4.1.5 Domain of applicability. When proposing a validated model it is very important to simultaneously define its limits so that a well-defined applicability domain could indicate those predictions that can be considered reliable. When the model is used to screen new compounds it is important that structures that fall out the domain of applicability of the model are filtered out as the model cannot generate for these structures reliable predictions. Domain of applicability can be defined using similarity measurements based on the Euclidean distances among all training compounds and the test compounds. The distance of a test compound to its nearest neighbor in the training set is compared to a predefined threshold (APD) and the prediction is considered unreliable when the distance is higher than that. APD was calculated based on the following formula:

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$$APD = \langle d \rangle + Z\sigma \tag{8}$$

Calculation of  $\langle d \rangle$  and  $\sigma$  was performed as follows: first, the average of Euclidean distances between all pairs of training compounds was calculated. Next, the set of distances that were lower than the average was formulated.  $\langle d \rangle$  and  $\sigma$  were finally calculated as the average and standard deviation of all distances included in this set. Z was an empirical cutoff value and for this work, it was chosen equal to 0.5.47-50 Enalos Domain – Similarity node that executes the aforementioned procedure is included in our workflow and was used to assess domain of applicability of the proposed model.47-50

#### 4.2 Enalos InSilicoTox platform

Novamechanics Ltd has recently launched Enalos InSilico platform, a new toxicity and drug discovery platform freely available online.51 Enalos InSilico platform aims to address the need to reduce the amount of time spent by scientists in referencing disparate sources of data to aid decision making related to NPs design and bioactivity profile and it offers an efficient and cost-effective response to the EU REACH legislation and the desire to reduce animal testing. The available workflows are built based on diverse and reliable data sources and integrate advanced in silico tools to provide accurate predictions. The web service is solely based on open source and freely available software including the powerful KNIME (Konstanz Information Miner)35 which is a user friendly and comprehensive open-source platform for data analysis including also all analysis modules of the well-known Weka data mining.41 In this work we have used KNIME platform in order to simultaneously run and compare different modeling methodologies and explore which of the available methods (or combination) best suites our data. As previously mentioned, to address our needs for robust and accurate model development targeting structural optimization and design we have developed the Enalos family nodes that were made publicly available for all KNIME users.

Through Enalos InSilico platform, toxicity, biological activity and property predictions can be obtained for chemical structure provided by the user. Structures can be designed, entered as SMILES or imported in SDF format. The QNAR model described in this work can be selected from the pull down menu of the available workflows already developed and provided by the Enalos InSilicoNano platform.

#### 4.3 Virtual screening

In an effort to identify novel potent compounds, a virtual screening study was initiated for compounds included in Pub-Chem database. <sup>55</sup> PubChem is a publicly available database that archives the molecular structures and bioassay data within the National Institute of Health (NIH) Roadmap for Medical Research Initiative. PubChem is currently the largest publicly available molecular database with millions of entries.

PubChem database was used to retrieve potent compounds in the described virtual screening framework using the most active compound in our original database, compound 36, which has a PaCa2 cellular uptake value equal to 4.44 expressed as decadic logarithm of the concentration (pM) of NP per cell (log 10[NP]/cell). All compounds included in the PubChem database were compared to compound 36 in a similarity context on the basis of 42 integer value descriptors of molecular structure, called Molecular Quantum Numbers (MQNs). MQNs count elementary features that matter most for the properties of organic molecules: atoms, bonds, polar groups, and topological features. The MQN-space organises molecules by their global structural features, but also by their similarity in biological activity. Distances in MQN-space can be used to search for analogues of known drugs. The MQNs form a scalar fingerprint which can be used to measure the similarity between pairs of molecules and enable ligand-based virtual screening.

### Acknowledgements

This project has received funding from the European Union's Seventh Framework Programme for research, technological development and demonstration under grant agreement no 310451 (Project NanoMILE).

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