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Subject Section

MaNGA: a novel multi-objective multi-niche genetic algorithm for QSAR modelling

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Abstract

Background: Quantitative structure-activity relationship (QSAR) modelling is currently used in multiple fields to relate structural properties of compounds to their biological activities. This technique is also used for drug design purposes with the aim of predicting parameters that determine drug behaviour. To this end, a sophisticated process, involving various analytical steps concatenated in series, is employed to identify and fine-tune the optimal set of predictors from a large dataset of molecular descriptors. The search of the optimal model requires to optimize multiple objectives at the same time, as the aim is to obtain the minimal set of features that maximizes the goodness of fit and the applicability domain. Hence, a multi-objective optimization strategy, improving multiple parameters in parallel, can be applied.

Results: Here we propose a new multi-niche/multi-objective genetic algorithm (MaNGA) that simultaneously enables stable feature selection as well as obtaining robust and validated regression models with maximized applicability domain. We benchmarked our method on two simulated datasets. Moreover, we analyzed an aquatic acute toxicity dataset and compared the performances of single- and multi-objective fitness functions on different regression models.

Conclusion: Our results show that our multi-objective algorithm is a valid alternative to classical QSAR modelling strategy, for continuous response values, since it automatically finds the model with the best compromise between statistical robustness, predictive performance, widest AD, and the smallest number of molecular descriptors.

Availability: The python implementation of MaNGA is available at https://github.com/Greco-Lab/MaNGA. **Contact:** dario.greco@tuni.fi

Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

QSAR methods have been widely applied in different fields such as toxicity prediction and drug design (Cherkasov *et al.*, 2014). QSAR models link the numerical description of molecular structures, the molecular descriptors (MDs), to their known physical-chemical properties and biological activities (Cherkasov *et al.*, 2014; Todeschini and Consonni, 2009).

Despite continuous methodological developments in the QSAR field, obtaining accurate, reliable, and stable models can still be challenging. A large number of easily computable descriptors, such as topological indices, two-dimensional (2D) and three-dimensional (3D) fingerprints are of utmost use to describe chemical structures (Todeschini and Consonni, 2009; Tropsha, 2010). However, selecting an optimal combination of predictive MDs in QSAR modelling is analytically difficult (Goodarzi et al., 2012). A comprehensive search of the best subset of features is

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computationally expensive, since as many as 2^N possible subsets of features for a dataset with N descriptors can be considered. Different methods for feature selection have been used in OSAR studies, such as multivariate adaptive regression splines and lasso (Eklund et al., 2012), random forest importance selection (Svetnik et al., 2003), univariate methods (Liu, 2004), backward elimination and forward selection (Yasri and Hartsough, 2001). Particular attention has been given to optimization search strategies such genetic algorithms (Leardi, 2001). QSAR models should ideally contain the smallest set of MDs with best goodness of fit. Indeed, considering the Topliss and Costello rule (Topliss, 1972), the number of MDs should be smaller than 1/5 of the training set compounds. Multi-objective optimization has, therefore, been applied to identify the smallest set of MDs with the highest predictive ability. For example, Nicolotti et al. (Nicolotti et al., 2002) proposed two multiobjective methods that use genetic programming to provide an adequate compromise between the number of model descriptors and accuracy. Soto et al. (Soto et al., 2009) developed a two-step feature selection using a multi-objective genetic algorithm that first performs a preliminary screening of the possible optimal solutions and, in a second step, enables model refinements. Recently Barycki et al. suggested a multi-objective genetic algorithm as a feature selection strategy for the development of ionic liquids' quantitative toxicity-toxicity relationship models, where the same set of features was evaluated against multiple endpoints (Barycki et al., 2018). Another important challenge related to the modelling of high-dimensional data is the stability of the selected features, i.e., how stable the selected features are with respect to variations in the training set (Kalousis et al., 2007). Model stability may become of particular concern when the number of features is much higher than the number of compounds (Fortino et al., 2014), as it is in QSAR datasets (Goodarzi et al., 2012). Stability is usually neglected in traditional OSAR applications, where a single endpoint is predicted. However, considering the role of QSAR in the context of safe-by-design, stability becomes important for obtaining more robust and reproducible models. Thus, stability should be reported along with the other validation metrics. Moreover, the criteria established by the Organization for Economic Co-operation and Development (OECD) (Oecd, 2014) must be met to ensure the validity of QSAR models. Along with the criteria addressing the transparency as well as internal and external statistical validity (Golbraikh and Tropsha, 2002; Shi et al., 2001a; Chirico and Gramatica, 2012a; Consonni et al., 2009a, 2010), a QSAR model is only reliable and valid with a defined applicability domain (AD) (Gramatica, 2007). The AD is the theoretical extent of the structural and response spaces in which the model is applicable to make reliable predictions for compounds with no experimental data. In brief, a good QSAR model should exhibit the best compromise between the number of MDs, predictive performance and the widest AD. To date, QSAR modeling has been traditionally carried out by following certain sequential steps: i) data preprocessing (preparation of the modelable dataset and training/test set splitting), ii) feature selection and modeling based on the optimization of individual parameters considered separately (e.g. R2), iii) internal and external validation of the models, and iv) AD definition (Cherkasov et al., 2014; Tropsha, 2010; Gramatica, 2007; Tropsha and Golbraikh, 2007; Gramatica et al., 2013a; Gramatica, 2013; Roy, 2007). In general, these steps are followed iteratively until the best model is identified. Also, each step is usually considered separately, not allowing a holistic evaluation.

In a multi-objective optimization problem, multiple objective functions are involved. More formally, a multi-objective optimization problem can be formulated as follows: $\min (f_1(x), f_2(x), \ldots, f_k(x))$, s.t. $x \in X$, where $k \geq 2$ is the number of objective functions, while X is the set of feasible decision vectors, that is defined by the constraint functions (Konak et al., 2006). In many cases, multiple objectives under consideration conflict with each other, thus, the optimization with respect to a single objective can lead to unacceptable results with respect to the others.

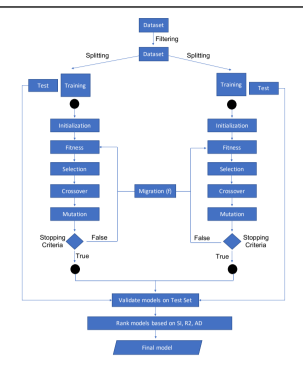


Fig. 1. Multi-niches multi-objective genetic algorithm methodology: A multi-objective multi-niches genetic algorithm that is able to identify smaller and stable sets of molecular descriptors that better predict activity with an optimal applicability domain.

A good compromise is to identify a set of solutions that satisfy the objective at different level and are not dominated by other solutions, meaning that they cannot be improved with respect to any objective without worsening at least one other objective (Konak *et al.*, 2006). The set of all feasible non dominated solutions is called Pareto optimal set, and their corresponding objective values are called Pareto front. Different multi-objective optimization approaches have been proposed with particular enphasys on evolutionary algorithms such as genetic algorithms (Konak *et al.*, 2006). Here, we propose a new multi-objective strategy based on genetic algorithms that simultaneously enables stable feature selection as well as robust and validated regression models with an optimal applicability domain.

2 Methods

2.1 Multi-niches multi objective genetic algorithm

Single- and multi-objective genetic algorithms were applied for feature selection in high-dimensional QSAR modelling. A multi-niche multiobjective genetic algorithm (MaNGA), (Figure 1), was implemented to compare the behaviour of 15 different objective functions (Table 1) and to select the most stable solutions in terms of feature robustness. The multi-objective non-dominated sorting genetic algorithm II (NSGA-II) (Deb et al., 2002), was implemented in python with the DEAP computational framework (Fortin et al., 2012). The pseudocode of the MaNGA algorithm is available as supplementary file 1. The NSGA-II algorithm was selected since it guarantees less computational complexity compared to other evolutionary algorithms and uses elitism to prevents the loss of good solutions among different iterations. In the proposed method, the individual solutions are binary chromosomes of length equal to the number of MDs in the dataset. A one (or zero) in the i-th position indicates that the i-th feature is (or is not) selected to be in the solution. Niching methods segment the genetic algorithm population

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Table 1. Objective functions and internal and external evaluation metrics. The 15 single- and multi- objective functions implemented in this work are reported (marked with Roman number) as combination of six base functions: R^2 , RMSE, AD, Q^2 , Q_{F3}^2 and NFeat. The x symbol is used to specify which base functions compose the objective functions. Furthermore, the list of internal and external validation metrics is reported. The threshold for the metrics are the following: $AD_{te} = 100$, $CCC_{te} > 0.85$, $Q^2 > 0.6$, $Q_{F1}^2 > 0.6$, $Q_{F2}^2 > 0.6$, $Q_{F3}^2 > 0.6$, NFeat < 1/5 of the nr. of training set samples. Arrows show if the objective functions have to be minimize or maximize.

Original Dataset																		
Training Set														Test Dataset				
	Objective functions combination for feature selection Internal Validation														External Validation			
Min/Max	Base Function	I	II	III	IV	V	VI	VII	VIII	IX	X	XI	XII	XIII	XIV	XV	R_{tr}^2	R_{te}^2
↑	R^2				Х	х	X	X				х	x	X		х	$RMSE_{tr}$	$RMSE_{te}$
+	RMSE	X				x			X	X		X	x		X	х	AD_{tr}	AD_{te}
↑	AD		X				X		X		X		X	X	X	х	Q^2	CCC_{te}
↑	Q^2				Х	x	X	X				X	x	X		х	Q_{F1}^2	
↑	Q_{F3}^2				X	X	X	X				X	X	X		X	Q_{F2}^2	
+	NFeat			х				X		х	x	х		X	X	х	Q_{F3}^2	

Table 2. Evaluation metrics formulas and their accepted thresholds.

Metric	Min/Max	Threshold	Ref
$RMSE = \sqrt{\sum_{i=1}^{N} (y_i - \hat{y}_i)^2}$	Min		(Aptula et al., 2005)
$R^{2} = 1 - \frac{\sum_{i=1}^{n} (y_{i} - \hat{y}_{i})^{2}}{\sum_{i=1}^{n} (y_{i} - \bar{y}_{i})^{2}}$	Max	$R^2 > 0.6$	(Golbraikh and Tropsha, 2002)
$Q^{2} = 1 - \frac{\sum_{i=1}^{n} (y_{i} - \hat{y}_{i})^{2}}{\sum_{i=1}^{n} (y_{i} - \bar{y}_{i})^{2}}$	Max	$Q^2 > 0.5$	(Golbraikh and Tropsha, 2002)
$Q_{F_1}^2 = 1 - \frac{\sum_{i=1}^{n_{test}} (y_i - \hat{y_i})^2}{\sum_{i=1}^{n_{test}} (y_i - y_{trx})^2}$	Max	$Q_{F_1}^2 > 0.6$	(Shi et al., 2001b; Chirico and Gramatica, 2012b)
$Q_{F_2}^2 = 1 - \frac{\sum_{i=1}^{n_{test}} (y_i - \hat{y_i})^2}{\sum_{i=1}^{n_{test}} (y_i - y_{test}^-)^2}$	Max	$Q_{F_2}^2 > 0.6$	(Schüürmann <i>et al.</i> , 2008; Chirico and Gramatica, 2012b)
$Q_{F_3}^2 = 1 - \frac{\left[\sum_{i=1}^{n_{test}} (y_i - \hat{y}_i)^2\right]/n_{test}}{\left[\sum_{i=1}^{n_{tr}} (y_i - y\bar{t}_r)^2\right]/n_{tr}}$	Max	$Q_{F_3}^2 > 0.6$	(Consonni et al., 2009b, 2010; Chirico and Gramatica, 2012b)
$CCC = \frac{2\sum_{i=1}^{n_{test}} (y_i - \bar{y})(\hat{y}_i - \bar{\hat{y}})}{\sum_{i=1}^{n_{test}} (y_i - \bar{y})^2 + \sum_{i=1}^{n_{test}} (\hat{y}_i - \bar{\hat{y}})^2 + n_{test}(\bar{y} - \bar{\hat{y}})^2}$	Max	CCC > 0.85	(Chirico and Gramatica, 2012b)
** Q^2 , $Q^2_{F_1}$, $Q^2_{F_2}$ and $Q^2_{F_3}$ are computed with cross-validation s	trategies		

 $i\in P$, called niches, such as $A_i\cap A_j=\emptyset$ whenever $i\neq j$. These methods lead to a better cover of the searching region and of the local optima. MaNGA is implemented with a multi-niche schema with 20 different niches independently evolving their own populations by means of crossover and mutation. The niches interact between them by a genetic operator called migration that swap the top best 25% element of their populations, selected with the Pareto strategy. The interaction between the niches was implemented by using a Queue structure. Every niche contains a population of 500 individuals, with a mutation rate of 5% evolving for 500

generations. The best parameters setting was determined prior to running

 $P = \{1, \dots, n\}$ containing n individuals, into k disjoint sets N_i with

2.2 Objective functions and evaluation criteria

the genetic algorithm by preliminary analyses.

MaNGA was applied to explore the feature space that optimizes the fitness functions (Table 1). In particular, the first class of fitness function called the R^2 , maximizes the R^2 , Q^2 , and $Q^2_{F_3}$. The second class of

fitness function, defined as MSE minimizes the mean squared error in cross-validation on the training set. The third class of objective function, called NFeat is meant to minimize the number of MDs selected by the GA. The AD objective function aims at maximizing the applicability domain of the models both on the training and test set samples. These objective functions were combined to create 15 single- and multi- objective functions which performances in the MaNGA method were investigated. For every niche, 20% of the original dataset was set aside as the test set, and not used in the model selection phase, but only to externally validate the trained model. The remaining 80% of the dataset was used to perform feature selection, train and internally validate the model, by using a 5-fold cross validation repeated three times (Table 1). In order to check the stability of the feature selection algorithm, for every niche, a different training/test sets split was generated. Different regression models were compared. In particular, linear regression model (Freedman, 2009), support vector regression (Basak et al., 2007) and k-nearest neighbours for regression (Zhou and Li, 2005) were implemented. Once the multiobjective optimization and feature selection step was performed, a pool

Table 3. Simulated datasets. The number of compounds and MDs and the generative models for the two synthetic datasets.

Dataset Nr.Comp MDs
Sim_{D_1} 77 55
$Y_{D_1} = 0.3 + 7.33 \text{ MD8} + 11.65 \text{ MD11} + 12.47 \text{ MD29} + 0.05 \text{ MD30} + 11.51 \text{ MD31} + 12.65 \text{ MD33} + 9.6 \text{ MD35} + 9.12 \text{ MD36} + 12.58 \text{ MD44} + 8.055 \text{ MD55} + \epsilon + 12.65 \text{ MD35} + 12.65 \text{ MD36} + 12.65 \text{ MD37} + 12.65 \text$
Sim_{D_2} 518 115
$Y_{D_2} = 0.3 + 10.09~\text{MD}10 - 9.50~\text{MD}15 + 8.72~\text{MD}17 + 9.79~\text{MD}22 + 12.70~\text{MD}25 + 7.75~\text{MD}26 + 14.01~\text{MD}27 + 8.32~\text{MD}55 + 12.95~\text{MD}64 + 8.62~\text{MD}65 + \epsilon + 12.00~\text{MD}10 $

of pareto-front optimal solutions were identified as the first output of the MaNGA algorithm. Thus, predictive performances for these models were evaluated in terms of the up-to-date criteria for QSAR models. Independently from the objective function used in the optimization process, $RMSE_{tr}$ (Aptula $et\ al.$, 2005), the coefficient of determination R_{tr}^2 , the leave-more-out correlation coefficient Q_{LMO}^2 , the $Q_{F_1}^2$, $Q_{F_2}^2$, $Q_{F_3}^2$, and the AD_{tr} were computed, on the training set as internal validation metrics, for every model. Furthermore, the $RMSE_{te}$ (Aptula $et\ al.$, 2005), R_{te}^2 , AD_{te} and the concordance correlation coefficient CCC_{te} , were computed as external validation metrics on the test set. The detailed description of the evaluation metrics is reported in Table 2.

2.3 Number of features

An objective function minimizing the number of features to be included in each solution was introduced (Table 1). This objective function is optimized under the constraint that the MDs in the solutions do not exceed $\frac{1}{5}$ of the number of compounds in the training set. Solutions that do not satisfy this requirement were heavily penalized during the optimization process by assigning them a high fitness value. However, the maximum number of MDs in the solutions is a parameter of the algorithm that the user can change arbitrary. Moreover, in order to easily converge to solutions satisfying the requirement, the initial binary population in every niche was generated with a probability of having value "0" equal to 0.99 and value "1" equal to 0.01.

2.4 Applicability Domain

The applicability domain was defined by means of the Williams Plot based on standardized residuals and leverage values. The Williams Plot helps identifying the response outliers as the ones following outside the 3σ range of the normally distributed standardized residuals, that covers 99% of the samples. The leverage value (h) measures the distance from the centroid of the modelled space. A warning leverage, referred to as the critical hat value (h^*) , is set at 3(p+1)/n, where p is the number of variables appearing in the model and n is the number of samples in the training set. Thus, a compound was considered influential and identified as a highleverage compound if $h > h^*$. In a Williams plot (Gramatica, 2007), the leverage values were mapped against the standardized residuals to define the structural and the response spaces visually. Finally, the AD was reported as the percentile coverage for the training (AD_{tr}) and the test (AD_{te}) set, respectively. Solutions with AD_{te} less than 100% were heavily penalized during the optimization process by assigning them a negative fitness value.

2.5 Selection of the final model

Multi-objective optimization methods, give as a result a set of optimal solutions distributed over the Pareto front represent different compromises between the best values of the multiple objectives to be optimized. Indeed, at the end of the iterations, a pool of 200 solutions was obtained by selecting the first 10 ranked solutions from every niche. However, as per traditional QSAR, a subsequent step for the models prioritization was carried out. Hence, every solution was further validated on the test set. Among these solutions, the unique sets of selected features were identified and ranked

based on their occurrence frequency over the pool of 200. The solutions were then filtered based on thresholds already established in the literature: $R_{tr}^2>0.6$ (Golbraikh and Tropsha, 2002), $R_{te}^2>0.6$ (Golbraikh and Tropsha, 2002), Q_{F1}^2 (Chirico and Gramatica, 2012a; Shi *et al.*, 2001a), Q_{F2}^2 and $Q_{F3}^2>0.6$ (Chirico and Gramatica, 2012a; Consonni *et al.*, 2009a, 2010; Schüärmann *et al.*, 2008), $CCC_{te}>0.85$ (Chirico and Gramatica, 2012a), $AD_{te}=100$. Only the solutions that satisfy these requirements were considered eligible. Finally, the most frequent solution surviving the filtering was selected as the best one.

2.6 Time-Complexity Analysis

The computational complexity of our method is bounded by the complexity of the NSGA-II algorithm, that is $O(kS^2)$ (Deb et al., 2002), where k is the number of objectives to be optimized (from 1 to 5 for our approach) and S is the population size (500 for our approach). The timecomplexity of the fitness functions are reduced to the time-complexity of the regression methods for the evaluation of the subset of features. Let n be the number of compound in the training sample, and p the number of MDs, the time complexity for the multiple linear regression method is $O(p^2n + p^3)$, the time complexity for the kernel SVR is $O(n^2p + n^3)$ and the time complexity for the k-nearest neighbors is O(np). Moreover, it has to be considered that the fitness function is computed in a repeated cross validation strategy, with 5 folds and 3 repetitions. The computational complexity of each iteration of our MaNGA algorithm is given by $O(kRS^2)$, where R is the time complexity of the selected regression model. Finally, this has to be multiplied by the number of iterations (500 in our approach). Even though, the method is computationally intensive compared with other FS methods (Guyon and Elisseeff, 2003), CPU-time is not a crucial issue provided that the algorithm may be executed in a reasonable polynomial time and feature selection is not aimed to be applied in real time.

2.7 Simulated dataset

In order to test the effectiveness of the proposed method, two simulated datasets with different number of compounds and MDs were generated, where the association between the MDs and the response is known. First, the original matrix of descriptors from a subset of the drugs in the connectivity map database (CMap) (Lamb et al., 2006) was generated. For each drug, the correspective 3D SDF file was downloaded from PubChem (Kim et al., 2016) and given in input to the DRAGON v. 7 software (Mauri et al., 2006) to compute 5.325 MDs. The constant (>80%) and highly interrelated MDs (pairwise correlation among all pairs of descriptors (>95%)) were filtered, as suggested (Gramatica et al., 2013b). Next, a population of true coefficients with only 10 relevant extracted from a normal distribution with mean 10 and standard deviation 2 were chosen, while the other non-relevant were set to zero. The relevant values were selected in order to obtain a full applicability domain with all the drugs inside the AD range. Furthermore, the error term ϵ was defined as an independent random normal vector, with mean 0 and standard deviation of 0.1. The intercept value was set as b0 = 0.3, and the response variable y as $y = b0 + X + \epsilon$ was computed. Finally, the MDs correlated with the ten relevant ones or correlated with the response y were removed. The

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Table 4. Selected features, internal and external validation metrics and frequencies for the solutions of MaNGA run on the Sim_{D_1} dataset. MDs column show the MDs selected by the MaNGA algorithm. The best solutions are those with: i) $RMSE_{tr}$ and $RMSE_{te}$ values as close as possible to zero; ii) AD_{tr} and AD_{te} as close as possible to 100; iii) $R_{tr}^2, Q^2, Q_{F1}^2, Q_{F2}^2, Q_{F3}^2$ and CCC_{te} as close as possible to 1.

ObjFun	Reg	NF	$RMSE_{tr}$	$RMSE_{te}$	AD_{tr}	AD_{te}	R_{tr}^2	R_{te}^2	Q^2	$Q_{F_1}^2$	$Q_{F_2}^2$	$Q_{F_3}^2$	CCC_{te}	Freq
R^2	LR	4/10	0.09	0.12	98.55	100	0.96	0.94	0.94	0.97	0.97	0.97	0.96	10.20
RMSE	LR	4/10	0.07	0.54	95.65	87.5	0.97	92.32	0.95	9.95	0.95	0.95	0.06	30.00
AD	LR	1/4	0.44	0.44	98.55	100	0.06	-0.04	-0.07	0.77	0.77	74	0.03	26.19
NFeat	LR	0/3	0.44	0.52	97.10	100	0.09	0.10	-13.36	-12.63	-12.67	-11.93	0.12	34.00
R^2 -RMSE-AD-NFeat	LR	2/3	0.14	0.18	100	100	0.90	0.84	0.88	0.90	0.90	0.90	0.92	26.19
R^2 -RMSE-AD-NFeat	SVM	3/3	0.06	0.11	98.55	100	0.98	0.98	0.95	0.97	0.97	0.96	0.96	24.00
R^2 -RMSE-AD-NFeat	kNN	3/3	0.06	0.10	98.55	100	0.99	0.99	0.97	0.97	0.97	0.97	0.97	31.03

Table 5. Selected features, internal and external validation metrics and frequencies for the solutions of MaNGA run on the SimD2 dataset. MDs column show the MDs selected by the MaNGA algorithm. The best solutions are those with: i) $RMSE_{tr}$ and $RMSE_{te}$ values as close as possible to zero; ii) AD_{tr} and AD_{te} as close as possible to 100; iii) R_{tr}^2 , Q^2 , Q_{F1}^2 , Q_{F2}^2 , Q_{F3}^2 and CCC_{te} as close as possible to 1.

ObjFun	Reg	NF	$RMSE_{tr}$	$RMSE_{te}$	AD_{tr}	AD_{te}	R_{tr}^2	R_{te}^2	Q^2	$Q_{F_1}^2$	$Q_{F_2}^2$	$Q_{F_3}^2$	CCC_{te}	Freq
R^2	LR	2/5	0.07	0.06	97.85	94.23	0.90	0.92	0.89	0.91	0.91	0.91	0.94	11.63
RMSE	LR	6/14	0.03	0.03	97.42	98.08	0.98	0.98	0.98	0.99	0.99	0.99	0.99	40.00
AD	LR	1/13	0.17	0.17	100	100	0.37	0.35	0.33	0.91	0.91	0.92	0.52	59.38
NFeat	LR	0/3	0.21	0.22	97.00	100	0.02	0.05	0.01	0.87	0.87	0.86	0.04	32.00
R^2 -RMSE-AD-NFeat	LR	2/7	0.06	0.07	99.14	100	0.91	0.89	0.91	0.97	0.97	0.97	0.95	100
R^2 -RMSE-AD-NFeat	SVM	2/3	0.05	0.05	99.79	100	0.95	0.95	0.94	0.88	0.88	0.87	0.97	50.00
R^2 -RMSE-AD-NFeat	kNN	2/3	0.03	0.04	98.50	100	0.98	0.96	0.95	0.95	0.95	0.96	0.97	33.33

Table 6. Multi-objective results for different training/test splits for each niche. The table reports the results for the dataset log-mmol-acqua, for each objective function and for each regression model. Optimal solutions have R^2 close to 1, FReq close to 100, RMSE close to 0, NF as small as possible and AD close to 100. Validation criteria fulfilled are in bold. The best solutions are those with: i) $RMSE_{tr}$ and $RMSE_{te}$ values as close as possible to zero; ii) AD_{tr} and AD_{te} as close as possible to 100; iii) $R_{tr}^2, Q^2, Q_{F1}^2, Q_{F2}^2, Q_{F3}^2$ and CCC_{te} as close as possible to 1.

ObjFun	Reg	NF	$RMSE_{tr}$	$RMSE_{te}$	AD_{tr}	AD_{te}	R_{tr}^2	R_{te}^2	Q^2	$Q_{F_1}^2$	$Q_{F_2}^2$	$Q_{F_3}^2$	CCC_{te}	Freq
R^2	LR	13	0.59	0.53	94.98	100	0.78	0.85	0.75	0.68	0.68	0.7	0.85	13.64
RMSE	LR	15	0.53	0.63	94.21	100	0.82	0.74	0.79	0.61	0.61	0.6	0.88	33.33
AD	LR	6	1.00	0.91	100	100	0.42	0.48	0.38	0.74	0.73	0.76	0.56	33.33
NFeat	LR	3	0.93	0.83	97.30	100	0.49	0.59	0.46	0.74	0.74	0.73	0.63	28.38
R^2 -RMSE-AD-NFeat	LR	5	0.62	0.63	95.75	100	0.78	0.83	0.76	0.88	0.88	0.86	0.86	12.82
R^2 -RMSE-AD-NFeat	SVM	14	0.36	0.45	97.30	100	0.92	0.91	0.83	0.91	0.91	0.92	0.89	20.93
R^2 -RMSE-AD-NFeat	kNN	10	0.39	0.44	96.91	100	0.91	0.89	0.81	0.86	0.86	0.88	0.88	22.22

thresholds used for the correlation filtering were -0.2 and 0.2. The first simulated dataset (called Sim_{D_1}) contains 77 drugs and 55 MDs. The second simulated dataset (called Sim_{D_2}) consists of 518 drugs and 115 MDs. Details on the simulated dataset are shown in (Table 3). The two simulated datasets are available as supplementary files 2 and 3.

2.8 Fathead minnow acute toxicity dataset

A dataset was retrieved from the literature (He and Jurs, 2005) containing measured acute toxicity as 96-h pLC50 (in mmol/L unit) of 288 compounds to Pimephales promelas (fathead minnow). The 3D .SDF files were downloaded from PubChem (Kim *et al.*, 2016) and used as input in the software DRAGON v.7 (Mauri *et al.*, 2006) to obtain 5,325 molecular descriptors. Unsupervised feature reduction was applied to filter the constant (>80%) and highly intercorrelated descriptors (pairwise correlation among all pairs of descriptors >95%) prior to training/test set splitting, and variable selection (Gramatica *et al.*, 2013b). After the preprocessing, 954 molecular descriptors were considered for further analysis. The list of compounds used in this study is available in supplementary file 4.

3 Results and discussion

Here, we compared the performances of single- and multi-objective functions (Table 1), using linear and nonlinear regressions, for QSAR modelling. We investigated the performance of our novel algorithm, MaNGA, on two simulated datasets. Furthermore, we present a case study on a fathead minnow acute toxicity dataset (hereafter referred to as log-mmol-acqua) (He and Jurs, 2005).

3.1 MaNGA algorithm selects relevant features in simulated datasets

In order to show the effectiveness of our proposed methodology, we tested the performances of the MaNGA algorithm on two simulated datasets with known optimal set of features. We compared different objective functions (comprising R^2 , RMSE, NFeat, AD and R^2 -RMSE-NFeat-AD) in combination with three regression models (linear, SVR and kNN). MaNGA was run with the following parameters: 500 individuals, 50 number of iterations, 5 niches. We selected the first 10 best solutions for every niche for a total of 50 solutions and investigated the set of features identified by the algorithm along with the internal and external validation metrics and the applicability domain value. Our results proved that the

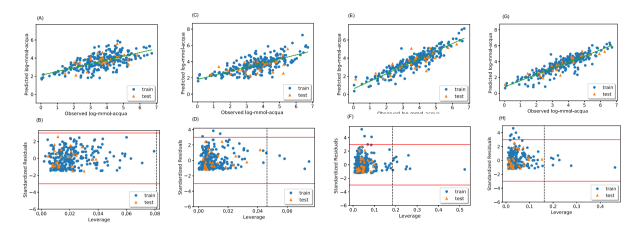


Fig. 2. Comparison of the results obtained from the linear regression model with single objective functions. The observed versus predicted log-mmol-acqua values (A) and the Williams plot representing the applicability domain (B) of the linear model maximizing the AD. The observed versus predicted log-mmol-acqua values (C) and the Williams plot representing the applicability domain (D) of the linear model minimizing the NFeat. The observed versus predicted log-mmol-acqua values (E) and the Williams plot representing the applicability domain (F) of the linear model minimizing the MSE. The observed versus predicted log-mmol-acqua values (G) and the Williams plot representing the applicability domain (H) of the linear model maximizing the \mathbb{R}^2 .

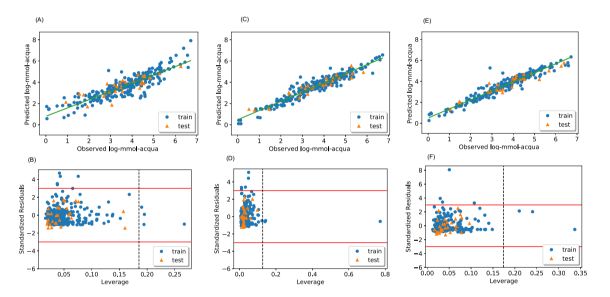


Fig. 3. Comparison of the results obtained with multi-objective functions that maximize the R2, and AD while minimizing the number of features and MSE. The observed versus predicted log-mmol-acqua values (A) and the Williams plot representing the applicability domain (B) of the linear model. The observed versus predicted log-mmol-acqua values (C) and the Williams plot representing the applicability domain (D) of kNN model. The observed versus predicted log-mmol-acqua values (E) and the Williams plot representing the applicability domain (H) of the SVR model.

MaNGA algorithm correctly preferred the selection of features belonging to the optimal solution, in both datasets (supplementary files 5 and 6).

When using the linear regression, the number of optimal MDs in the solutions is higher than the MDs in the solutions coming from SVM and kNN regressors (Tables 4 and 5). Moreover, the optimal solutions are not selected when solely minimizing the number of features. On the other hand, when performing the multi-objective optimization with both SVM and kNN regression, the best minimal set of features is included both in the solutions of the Sim_{D_1} (all the features) and Sim_{D_2} dataset (two out of three), respectively. With regards to the evaluation metrics, good predictive abilities were obtained by the models resulting from the optimization of the R^2 , MSE and multi-objective, while low predictive capacity is achieved by the models obtained by maximizing only the AD or minimizing the number of MDs.

3.2 Multi-objective functions provide the best compromise between number of MDs, widest AD and goodness of fit

We applied the MaNGA algorithm to a fathead minnow acute toxicity dataset (He and Jurs, 2005) by using different single- and multi-objective functions, and three regression models. The AD and NFeat objective functions did not produce models that met all the validation requirements, indeed R_{tr}^2 , R_{te}^2 and Q^2 were lower than 0.6 and CCC_{te} was lower than 0.85 (Table 6). On the other hand, the models obtained by optimizing the R^2 , RMSE and R^2 -RMSE-AD-NFeat functions fulfilled all the validation criteria. Moreover, the number of features selected by the multi-objective function was lower than the one selected by the single-objective ones (Table 6). Furthermore, the AD coverages of the multi-objective function were slightly higher than those obtained with the single-objective ones (Table 6). This was particularly clear when investigating the scatterplot of the observed versus predicted values and the Williams plot (Figure 2). Indeed,

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when MaNGA was run by only optimizing the AD values, none of the training or test set samples fell outside the applicability domain (Figure 2B), but the predictive performances of the model were quite low, with poor concordance between the predicted and experimental data (Figure 2A). Thus, the model did not fulfill the criteria regarding the goodness of fit and the internal and external validation requirements ($R_{tr}^2 < 0.45$ for LR and SVR models, R_{te}^2 and $Q^2 < 0.45$ and $CCC_{te} < 0.85$ for LR, SVR and kNN models). When the optimization was carried in terms of the number of selected MDs, the model contained only 3 features (Table 6), but there was no high concordance between the predicted and experimental data (Figure 2C), and AD was not satisfactory (Figure 2D). Moreover, this model did not fulfil the criteria regarding the goodness of fit and the internal and external validation requirements ($R_{tr}^2 < 0.6$ for LR and SVR models and Rte2 and $Q^2 < 0.6$ for LR, SVR and kNN models, Table 6). On the other hand, when the optimization was performed by minimizing the MSE or maximizing the R2, the prediction capability of the model were better. Indeed Figure 2E and Figure 2G, show a good concordance between the predicted and experimental data ($R_{tr}^2 > 0.6$), the solutions satisfied all the requested parameter of internal and external validation, but the AD values decreased to 94.21% on the training set when optimizing for the MSE (Figure 2F, Table 6) and to 94.96% on the training set when optimizing for the \mathbb{R}^2 (Figure 2H, Table 6). Moreover, the number of MDs in the models increased to 15 and 13 for MSE and R^2 , respectively (Table 6). The multi-objective optimization strategy allowed to meet as many requirements as possible with the same solution, particularly, when the optimization was performed by minimizing the number of features and the RMSE while maximizing the \mathbb{R}^2 and AD (Figure 3). Moreover, using a non-linear model such as kNN (Figure 3C and 3D) or SVR (Figure 3E and 3F) significantly improved the goodness of fit of the model compared to the linear regression (Figure 3A and 3B). The selected models are available in supplementary file 7. The predicted values for the multi-objective function R^2 -AD-RMSE-NFeat for the linear, SVR and kNN models are reported in supplementary files 8, 9 and 10.

3.3 Analysis of selected descriptors

Next, we analyzed the MDs selected by the MaNGA algorithm on the fathead minnow acute toxicity dataset by their occurrence frequency across the generated models. The most frequently selected MD was BLTD48. BLTD48 is the Verhaar Daphnia (48-h) base-line toxicity from MLOGP (mmol/L), a molecular property negatively relating to the Moriguchi hydrophobicity ($BLTD48 = -0.95 \times MLOGP - 1.32$) (Verhaar et al., 1992; Moriguchi et al., 1992). BLTD48 showed a negative coefficient in the two linear regression models obtained with the MSE and R2 objective functions. The inverse relationship between the BLTD48 and MLOGP ultimately suggests that increasing hydrophobicity accounts for increasing toxicity. The second most frequently selected MD is another hydrophobicity term, the squared Ghose-Crippen octanol-water partition coefficient, ALOGP2, (Ghose et al., 1998). ALOGP2 showed a positive coefficient in the linear regression model when optimizing for the RMSE, thus, explaining increasing aquatic toxicity. Different octanolwater partition coefficients calculated based on different approaches, such as atomic contribution in the case of ALOGP and property based methods in the case of MLOGP, have been reported elsewhere (Martin et al., 2015). Other MDs selected by more than one model are the Mor12s, nN, P_VSA_logP_3, and Ui. Mor12s is the signal-12 3D-MoRSE descriptor weighted by the intrinsic state (I-state). The I-state of an atom is the possible partitioning of the influence of non- σ electrons throughout the σ bonds within a molecule starting from the atom in consideration (Todeschini and Consonni, 2009). Hence, the less partitioning of the electron influence can be attributed to that the valence electrons are more prone to intermolecular interactions, which possibly result in toxicity.

nN is a constitutional index counting the number of nitrogen atoms in a molecule also reported to be correlated to fathead minnow acute toxicity (Papa *et al.*, 2005); P_VSA_logP_3 (P_VSA-like on LogP, bin 3) is a molecular descriptor defined as the amount of van der Waals surface area (VSA) having a property in a certain range and is related to hydrophobicity (Labute, 2000). Finally, Ui is another molecular property descriptor representing the unsaturation index. The complete list of descriptors selected by MaNGA is available in supplementary file 11.

3.4 Comparison with previous models

We compared our models for fathead minnow acute toxicity, with previously independent models coming from literature (Supplementary file 12). Different regression models were applied such as linear regression, PLS, neural networks and k nearest neighbours. The models were not evaluated by an exhaustive set of metrics, thus we compared our results based only on those available. All the models, except one, were internally evaluated by using the R_{tr}^2 . The most used metric for the evaluation of the external predictive performance was the R_{te}^2 . Only the model described in (Cassotti et al., 2015) was identified by using the same MDs used in this study, and evaluated also in terms of Q^2 , RMSE and AD, thus we choose this model for comparison. The model was obtained with a combination of genetic algorithm and knn regression model. It contains six MDs, and depending on the different thresholds that were set to compute the AD, it reached a $R_{tr}^2 \in [0.62 - 0.73]$, $RMSE_{tr} \in [0.65 - 0.87]$, $Q^2 \in [0.61 - 0.79], R_{te}^2 \in [0.61 - 0.77] \text{ and } RMSE_{te} \in [0.68 - 0.88].$ Our models obtained by using the $R^2 - RMSE - AD - NFeat$ multi-objective function (Table 6), reached better performances on all the metrics, even thought, the models obtained by using the kNN and SVR regression models use a higher number of MDs, while the model obtained by using the linear regression method uses only five MDs. Furthermore, the applicability domain values or our models (95%-97%) are higher than those reported for the previous model (60%-80%) (Cassotti et al., 2015).

4 Conclusions

In this work, we presented MaNGA, a new multi-niche multi-objective genetic algorithm for feature selection in QSAR studies. We performed extensive analyses on two simulated datasets and showed that MaNGA correctly identifies sets of optimal features. Furthermore, we applied MaNGA to a real dataset to compare different single- and multi-objective functions as well as linear and non linear regression models. Our results suggest that multi-objective functions outperform the single-objective ones, since they allow to obtain the smallest possible set of features, with the widest possible applicability domain and the best possible goodness of fit with a single run of the MaNGA algorithm. We also provide facilities to monitor the stability of the selected features thus helping the evaluation of the proposed models. Finally, we want to highlight that the MaNGA methodology was developed for QSAR modelling with continuous response variables. However, a similar strategy can be applied to classification problems where the response variable is discrete.

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