A method for regularization of evolutionary polynomial regressions

Francisco Coelho*

Departamento de Informática, Universidade de Évora, Rua Romão Ramalho 58, 7000-671 Évora, Portugal

João Pedro Neto*

Departamento de Informática, Faculdade de Ciências da Universidade de Lisboa, Campo Grande, 1749-016 Lisboa, Portugal

Abstract

While many applications require models that have no acceptable linear approximation, the simpler nonlinear models are defined by polynomials. The use of genetic algorithms to find polynomial models from data is known as Evolutionary Polynomial Regression. This paper introduces Evolutionary Polynomial Regression with Regularization, an algorithm that extends the EPR method and describes a set of experiences on common datasets that compare both flavors of EPR and other methods including Linear Regression, Regression Trees and Support Vector Regression.

The empiric conclusion of those experiments is that EPR with regularization is able to achieve better fitting than other non-ensemble methods and it has shorter computation time than plain EPR.

Keywords: evolutionary polynomial regression, regularization, feature extraction

— GAs are repeatedly mentioned as an approach to search for locally optimal parameters, this is simply incorrect and is a very surprising statement. GAs are

^{*}Corresponding author: Tel.: +351-919-006-379

Email addresses: fc@di.uevora.pt (Francisco Coelho), jpn@di.fc.ul.pt (João Pedro eto)

NOT a local optimizer in any shape or form.

- What is the reference for the GA encoding used?
- The penalty term introduces is the main contribution of this work, this should be clearly stated in the text. It is basically a complexity based penalty which is really not so novel, there are even several cases in literature of penalty based GA functions. Please dig up further in the state of the art to back up better your claim of novelty.
- In some box plots only ten simulations are performed, in others 25. These numbers are too low.

3 1. Introduction

With notable exceptions (e.g. neural networks) machine learning regression techniques produce linear models. The linearity assumption has many advantages including reduced computational complexity and strong theoretical framework. However nonlinearity is unavoidable in many application scenarios, specially those with phase transitions or feedback loops, so common in engineering, ecology, cybernetics and other areas. The kernel trick in Support Vector Machines (SVM) ([1, 2, 3]) alleviates this problem by allowing special nonlinear transformations of the feature-space. The condition such transformations must meet is known as the kernel trick, $k(x,x') = \langle \varphi(x), \varphi(x') \rangle$, where φ is the feature-space transformation and $\langle \cdot, \cdot \rangle$ denotes inner product. The "trick" 13 consists on computing the kernel k(x, x') while avoiding the computation of the 14 inner product and the transformations $\varphi(x)$, $\varphi(x')$. A special case of polyno-15 mial transformation, the polynomial kernel, $k(x, x') = \langle x, x' \rangle^d$ is commonly used in regression and classification tasks with SVMs. However general polynomial transformations do not verify the kernel trick. — do not verify does not seem like the proper term. 19 Polynomials, one of the most studied subjects in mathematics, generalize linear functions and define, perhaps, the simplest and most used nonlinear mod-

els. Applications include colorimetric calibration [4], explicit formulæ for tur-

bulent pipe flows [5], computational linguistics [6] and more recently analytical techniques for cultural heritage materials [7], liquid epoxy moulding process [8], B-spline surface reconstruction [9], product design [10] or forecasting cyanotoxins presence in water reservoirs [11]. These examples not only illustrate the wide spectrum of applications but, additionally, each one uses, at some point, 27 Genetic algorithms (GA). 28 Evolutionary algorithms, including GA, were, arguably, one of the hottest topics of research in the recent decades and with good reason since they outline an optimization scheme easy to conceptualize and with very broad application. 31 If a nonlinear (or otherwise) model requires parameterization, GAs provide a simple and often effective approach to search for locally optimal parameters. 33 Related research abound and spans from the 1950s seminal work of Nils Aall Barricelli [12] in the Institute for Advanced Study of Princeton to today's principal area of study for thousands of researchers, covered in hundreds of conferences, workshops and other meetings. Perhaps the key impulse to GAs came 37 from John Holland's work and his book "Adaptation in Natural and Artificial Systems" [13]. 39 One interesting variation of genetic algorithms, named *qenetic programming* by John Koza [14], proposes the use of GAs to search the syntactic structure of 41 complex functions. Syntactic structure search is also keen to the central ideas of deep learning [15, 16], a subarea of machine learning actually producing quite 43 promising results (e.g. in [17]). It is also related to the work presented in this paper in the sense that, unlike linear models that have a simple structure, $y = \sum_{i} \beta_{i} x_{i}$, nonlinear (in particular polynomial) models pose an additional structure search problem. 47 The idea of using GAs to find a polynomial regression is not new [18, 19, 20] but still generates original research [21, 22]. The modern formulation of the use of GA to find polynomial models is known as Evolutionary Polynomial Regression (EPR) and systematization can be traced back to the work of Davidson, 51 Savic and Walters [23]. Further developments include multi-objective optimizations [24].

This paper describes an extension of the general EPR method to find a regularized polynomial regression of a given dataset. The optimal regression results come from a cost function that accounts for both the root-mean-square (error) and a regularization factor to avoid overfit by penalysing polynomial complexity.

The next section describes the method's details and is followed by a presentation of some performance results. The last section draws some conclusions and points future research tasks.

62 2. Genetic Algorithms for Polynomials

This section starts with a brief introduction and outline of the evolutionary polynomial regression algorithm, EPR, and proceeds into core details as the encoding used to represent individual polynomial instances in the GA populations and the regularization of the cost function.

A usual representation of polynomials is through expressions of the form

$$p(x_1,\ldots,x_m) = \sum_i \theta_i q_i$$

where each $q_i=\prod_j x_j^{\alpha_{ij}}$ is a monomial, the exponents $\alpha_{ij}\in\mathbb{N}_0$ are non-negative integers and the coefficients $\theta_i\in\mathbb{R}$ are real valued. For example $p\left(x_1,x_2,x_3\right)=2x_1+x_2x_3+\frac{1}{2}x_1^2x_3$ has monomials $q_1=x_1,q_2=x_2x_3$ and $q_3=x_1^2x_3$, exponents $\alpha_{1,1}=1,\alpha_{2,2}=1,\alpha_{2,3}=1,\alpha_{3,1}=2,\alpha_{3,3}=1$ and all other $\alpha_{ij}=0$ and coefficients $\theta_1=2,\theta_2=1$ and $\theta_3=1/2$.

The exponents alone can be organized into a matrix $[\alpha_{ij}]$ that defines the monomial structure of the polynomial. For the example above the matrix representation of the monomials is

$$\begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 1 \\ 2 & 0 & 1 \end{bmatrix} \sim \begin{bmatrix} x_1 \\ x_2 x_3 \\ x_1^2 x_3 \end{bmatrix}$$

72 where each row defines a monomial and each column represents a variable.

Changing the order of the rows doesn't change the polynomial whereas changing

the order of the columns corresponds to changing the respective variables.

This partial representation of polynomials makes the problem of structure search very clear: except for the trivial cases, the number of possible monomials given n variables and a maximum joint degree d grows exponentially with either n or d. But more importantly, by separating the set of monomials from the coefficients, the polynomial regression problem can be naturally split into two subproblems:

- 1. For a given set of monomials $Q = \{q_1, \dots, q_k\}$ find the regression coefficients $\Theta = \{\theta_1, \dots, \theta_k\}$ that minimize the error on a given dataset;
- 2. Find the fittest set of monomials, *i.e.* the polynomial that minimizes the error on the same dataset;
- More precisely, concerning the first problem, let \mathcal{D} be a dataset with n obser-
- vations of variables Y, X_1, \ldots, X_m and $\mathcal{Q} = \{q_1, \ldots, q_k\}$ a set of k monomial
- expressions over X_1, \ldots, X_m . Define the hypothesis¹

$$h_{\Theta,\mathcal{Q}}(x_1,\ldots,x_m) = \sum_{j=1}^k \theta_j q_j|_{X_i=x_i,\forall 1 \le i \le m}$$
 (1)

and let the error (as "cost") be

$$J_{\text{fit}}(\Theta; \mathcal{Q}, \mathcal{D}) = \sqrt{\frac{1}{n} \sum_{i=1}^{n} \left(y^{(i)} - h_{\Theta, \mathcal{Q}} \left(x_1^{(i)}, \dots, x_m^{(i)} \right) \right)^2}$$
(2)

the usual root-mean-square (error) function. Now the first problem can be stated as: Given a dataset \mathcal{D} and a set of monomials \mathcal{Q} find parameters Θ that

91 minimize the cost $J_{fit}(\Theta; \mathcal{Q}, \mathcal{D})$.

This is a simple linear regression problem obtained by expanding $\mathcal D$ with

- columns that replicate the monomials in \mathcal{Q} . The resulting dataset, $\mathcal{D} \cup \mathcal{Q}(\mathcal{D})$,
- adds the monomial transformations in \mathcal{Q} to the original dataset \mathcal{D} . An alter-
- native formulation would just replace \mathcal{D} by $\mathcal{Q}(\mathcal{D})$. It turns out that the first

¹The expression " $q|_{X=x}$ " reads "q with all instances of X replaced by x."

Algorithm 1 This EPR algorithm uses linear regression for the calculation of the error J and the space of polynomials is searched in the GAs iteration step. At exit the error of the fittest instance is bounded by ϵ or the maximum number of allowed iterations.

```
function \text{EPR}(D, pop_0, \epsilon, maxiter)
pop \leftarrow pop_0; \ err \leftarrow 1.0 + \epsilon
\text{while } err > \epsilon \land iterations < maxiter \ \mathbf{do}
pop \leftarrow \text{ITERATEGA}(pop)
pop \leftarrow \text{SORT}(pop, key = J) \qquad \triangleright \text{Sort population by regression error}
err \leftarrow J \left( \text{First}(pop) \right)
\text{end while}
\text{return First}(pop)
end function
```

formulation is a special case of the second (by including the variables in the monomial set) and has better error performance — what is not surprising because it uses more features. — How is this known??? Is there a reference or some experimental results backing this claim or is it simply intuition???

The second problem is treated in the GA setting: Let \mathcal{D} be a dataset as above and \mathcal{P} a set of polynomials. For each polynomial $p \in \mathcal{P}$ let \mathcal{Q}_p be the set of monomials in p (without the coefficients) and define the (anti) fitness — say what? Do you mean something like a minimization based fitness?? Anti fitness seems odd.

$$\phi_p = \min_{\Theta} J_{\text{fit}}(\Theta; \mathcal{Q}_p, \mathcal{D})$$
 (3)

by solving the first problem. With a fitness of every instance, the GA genetic operators (usually mutation and crossover) evolve the population \mathcal{P} until a reasonable approximation of a local minimum is found. The properties of GAs and linear regression entail that Algorithm 1 converges to a polynomial that is a local minimum of the fitness function — GAs are global optimization techniques (in spite of what wikipedia states!), encapsulated in the error function J_{fit} .

Subsection 2.1 describes the encoding of individual polynomial instances as chromosomes and other parameters used in the GA implementation. The regularization of the cost function is discussed in subsection 2.2.

2.1. Polynomial Encoding

122

123

124

125

126

127

128

129

The specific encoding (representation) of a set of monomials is an important aspect in the implementation of EPR. The choice described below permits
active and inactive monomials for regression purposes — is this an original contribution???? Otherwise it needs a reference.. The active (or inactive) state of a
monomial might change through mutation or crossover. This simple mechanism
enhances variation in the complexity of polynomial expressions by evolutionary
operations.

Let $\{q_1, \ldots, q_k\}$ be a set of monomials over the variables X_1, \ldots, X_m . The encoding of that set using d bits per exponent is a binary list such that

- 1. the initial segment of k bits defines the active state of each monomial;
- 2. the remaining bits are split into k segments of size $m \times d$, each representing a monomial;
- 3. the bits in each monomial segment are split into m sub-segments of size d. The j^{th} sub-segment is the binary representation of the degree of the variable X_j in the enclosing monomial segment;

This encoding can also be viewed as the flattening of the binary exponents in the matrix representation prefixed by the activation segment. The set $\{x_1^3x_3, x_3^7, x_1x_2\}$ (with m=3) has matrix representation

$$\begin{bmatrix} 3 & 0 & 1 \\ 0 & 0 & 7 \\ 1 & 1 & 0 \end{bmatrix} = \begin{bmatrix} 011 & 000 & 001 \\ 000 & 000 & 111 \\ 001 & 001 & 000 \end{bmatrix}_{(2)}$$

where the right matrix is in binary form using d=3 bits. An encoding of this set of monomials with the extra monomial $x_1^6x_2^2x_3^5$ inactive, setting k=4, would be

where, for reading purposes, semicolons separate segments and commas separate variables. The first k=4 bits inform that the first, second and third monomials 135 are active while the fourth is not.

While each valid encoding represents a set of monomials the map is not bijective: each set of monomials has multiple encodings, for example by changing d or the order of monomial segments. However, considering the EPR task, this is a minor issue and a bijective map would add computational complexity and negative impact to the algorithm's performance — The encoding example should include the inactive term so that both examples are consistent (lines 113-115). Is there a reference some experimentation to back this up?.

There is one final remark concerning this encoding method. As it is, the activation segment can become all zeros, representing the empty set of monomials. This situation can be avoided with a simple hack: Given an encoding, the first monomial is always considered active, thus restricting the syntactic form of encodings to binary strings starting with 1. In practice, this means that the implementation of the encoding can omit the first bit.

2.2. Cost Function 150

133

134

136

138

139

140

141

143

144

145

146

148

149

151

152

153

154

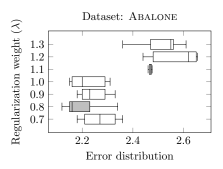
155

157

The polynomial regression error considered so far accounts for the ability to predict the transformed testset. A known problem of using a cost function based only in the dataset error (and of polynomial regressions in general) is the tendency to overfit training data. Excessive variance of the estimation method can be reduced by regularizing the error function with a penalty factor. Thus, to reduce polynomial complexity and variance by regularizing the size of the monomial set the error function from equation 2 is multiplied by a factor λ^k

$$J_{\text{reg}}(\Theta, \lambda; \mathcal{Q}, \mathcal{D}) = \lambda^k J_{\text{fit}}(\Theta; \mathcal{Q}, \mathcal{D})$$
 (4)

where k is the number of monomials in the polynomial. When $\lambda > 1$ polyno-158 mials with more monomials are penalized. The regularized extension of EPR is 159 denoted by Evolutionary Polynomial Regression with Regularization (EPRR).



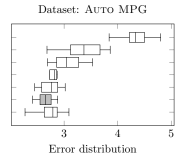


Figure 1: Error distribution by regularization exponent for two common datasets. The box plots summarize error values of ten simulations for each value of λ . The smallest overall error, in grey, is achieved in both datasets when $\lambda=0.8$. Performance of the non-regularized EPR is plotted in the line $\lambda=1$.

A simple exploration on the effect of the regularization parameter is depicted in Figure 1 where it is possible to observe that the typical inflection point lies around $\lambda=0.8$. This value, favoring "larger" polynomials is justified by the balance of the data's non-linearity and polynomial complexity: below $\lambda=0.8$, even penalized, larger monomial sets achieve better error performance than smaller ones while above that value the size of the monomial set is excessive—please clarify this paragraph and justify this claim. Within this tension the overall error results reduced when compared to the non-regularized EPR version.

2.3. Genetic Algorithm Parameterization

- Please indicate all these values in detail
- You should contrast obtained results in a table. Results are hard to judge from Fig 3 alone.
 - The number of 250 for iterations seems premature specially when one looks at Fig 4. Stabilize? Really? This only seems valid in one case of lambda.

In general GAs offer many possibilities with respect to the choice of genetic operators and respective application rates, population evolution, *etc*. The results found here where obtained using the package genalg [25] with default

parameters, standard operators (crossover and mutation) and population evolution with 20% elitism between generations.

3. Experimental Results

180

201

202

Here is described the experiment setup used to gather and summarize the empirical evidence that supports this comparative study of EPR and EPRR. Evaluation is focused in error distribution and, besides EPR and EPRR, also uses several common regression methods and datasets easily accessible in R, the free software environment for statistical computing and graphics [26]². A small consideration on the convergence speed concludes this section.

3.1. Regression Methods and Datasets

The EPRR method is ranked against several well-known learning algorithms for regression, namely: non-regularized EPR, Linear Regression, Support Vector Machines [27], Regression Trees [28] and Conditional Inference Trees [29, 30, 31]. To achieve better error results the SVM and Regression Tree parameters are tuned in each dataset.

The performance of each method is evaluated on several common datasets.
From each dataset 70% of the observations are reserved for training purposes and the remaining observations used to estimate the error. To enhance the robustness of results this process is repeated 25 times, each time with a different shuffling of the samples in the train and test sets. Some datasets with attribute values of different magnitudes have a pre-processing scaling transformation. The box plots in figures 2 and 3 resume the test set error distributions over these different runs.

One of the used datasets, ARTIFICIAL, has a special role: it is used to test if EPRR is able to discover a polynomial model. The idea of this test is to generate a polynomial dependent variable and measure the EPRR error after

²The datasets and R code used to produce the results and plots in this paper are available online at https://github.com/jpneto/GenAlgPoly.

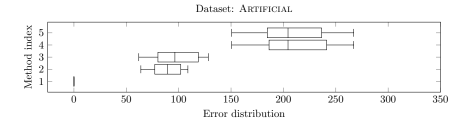


Figure 2: Testing polynomial discovery. The dataset is generated from a polynomial expression and, as shown, EPRR finds the exact generator structure: in line 1, the error box is centered in 0 and has width 0. The regression methods depicted are: 1. EPRR, 2. Linear Regression, 3. SVM, 4. Regression Trees and 5. Conditional Inference Trees

fitting the dataset. The genetic algorithm parameterization for this dataset uses a population with size n=100 and evolves for 50 generations. For the remaining datasets the population has size n=300 and evolves for 100 generations.

ARTIFICIAL is a polynomial dataset with four numeric features, $x_1, \dots x_4$,
where x_1, x_3 are outcomes from Poisson random variables, and x_2, x_4 from
Normal random variables. The dependent variable is given by the polynomial expression $y = x_2 x_4^2 + x_1^2 x_3 + 5$. The dataset includes n = 50 observations;

Housing concerns the task of predicting housing values in areas of Boston.

There are n=506 observations of m=13 continuous attributes and one dependent variable, the median value of owner-occupied homes in thousands of USD;

Abalone is used to predict the age of a abalone shell using m=8 numeric attributes concerning several physical measurements. There are n=4177 observations;

Auto MPG gathers fuel consumption in miles per gallon, based on two discrete and five continuous attributes (m=7). There are n=398 observations;

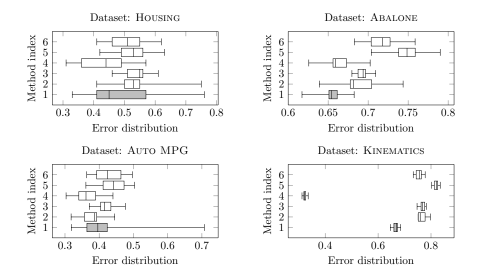


Figure 3: Summary results for different regression methods on diverse datasets. Although EPRR not always achieves the smallest expected error, performance is on-par with more sophisticated methods. The regression methods depicted in these figures are: 1. EPRR; 2. EPR; 3. Linear Regression; 4. SVM; 5. Regression Trees; 6. Conditional Inference Trees;

KINEMATICS results from a realistic simulation of the forward kinematics of an 8 link robot arm. The task is to predict the distance of the end-effector from a target using m=8 continuous attributes. There are n=8192 observations;

226 3.2. Convergence speed

222

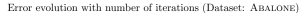
223

224

225

Since this work is oriented to the error of the EPRR model it is necessary to assess how this depends on the number of generations of the GA. As illustrated in Figure 4, the error quickly drops during the initial 50 to 100 generations.

Then, it proceeds slower achieving better solutions only with marginal error reduction.



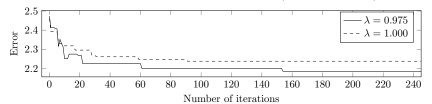


Figure 4: Learning curve: Error progress for the Abalone dataset during a single execution of the genetic algorithm. The figure shows the fitness evolution for two different regularization values. The population for both consists of 200 polynomials. The error values seem to stabilize around iteration 250.

4. Conclusion and Future Work

234

236

237

240

241

242

243

244

251

Of the regression methods considered SVM achieves the best results in three 233 out of four datasets. However SVM and Conditional Inference Trees are pretrained, having parameters tuned for each particular dataset unlike EPRR, that 235 runs with the same parameterization on all datasets. Even so it is the best estimator for the ABALONE dataset and in the remaining datasets it outperforms most of the other estimators. 238

Comparing EPR and EPRR — the main article's topic — the regularized version achieves much better results at Abalone and especially Kinematics. On the Housing dataset errors are improved wrt EPR in a difference in means, resulted in a 95% HDI (Highest Density Interval) equal to [0.001, 0.119] which, while borderline, achieves statistical significance. Only in the Auto MPG dataset EPR achieves better results, even if not that different from EPRR.

For complexity considerations EPR and EPRR demand some processing 245 time. On a quad-core computer, processing the Kinematics dataset (with near 246 8K observations) takes approximately 5 minutes. Probably processing time can 247 be reduced by one to two orders in magnitude if the algorithm is implemented with computational speed in mind. However, speed optimization is not the focus of this article. 250

A cross-validation procedure can be implemented to refine the appropriate

parameter values to achieve better errors. Namely, the regularization parameter, λ , can be tested with several values, instead of being fixed at 0.8. Other parameters like mutation chance or the amount of elitism can also be tested. However, these type of tests need a low-level, fast implementation of EPR and are postponed to future investigation.

Acknowledgements

The authors are grateful to the Fundação para a Ciência e Tecnologia (FCT)
and the R&D laboratory LabMAg for the financial support given to this work,
under the strategic project PEST-OE/EEI/UI0434/2011.

Datasets used herein are selected from Luís Torgo's data repository, http://www.dcc.fc.up.pt/~ltorgo/Regression/DataSets.html. Most can also be found in the UCI ML repository at http://archive.ics.uci.edu/ml/.

The authors wish to thank professor André Falcão for motivation and useful discussions around the article.

- [1] B. Schölkopf, A. Smola, K.-R. Müller, Kernel principal component analysis,
 in: Artificial Neural Networks ICANN'97, Springer, 1997, pp. 583–588.
- ²⁶⁸ [2] Z. Liang, Y. Lee, Eigen-analysis of nonlinear pca with polynomial kernels.
- [3] Y. Bao, Z. Hu, T. Xiong, A pso and pattern search based memetic algorithm
 for syms parameters optimization, Neurocomputing 117 (2013) 98–106.
- [4] L. Mendes, P. d. Carvalho, Adaptive polynomial regression for colorimetric
 scanner calibration using genetic algorithms, in: Intelligent Signal Processing, 2005 IEEE International Workshop on, IEEE, 2005, pp. 22–27.
- [5] J. Davidson, D. Savic, G. Walters, Method for the identification of explicit
 polynomial formulae for the friction in turbulent pipe flow., Journal of
 Hydroinformatics 1 (1999) 115–126.

- ²⁷⁷ [6] L. Sánchez, J. Otero, I. Couso, Obtaining linguistic fuzzy rule-based regres-²⁷⁸ sion models from imprecise data with multiobjective genetic algorithms, ²⁷⁹ Soft Computing 13 (5) (2009) 467–479.
- [7] L. Cséfalvayová, M. Pelikan, I. Kralj Cigić, J. Kolar, M. Strlič, Use of genetic algorithms with multivariate regression for determination of gelatine in historic papers based on FT-IR and NIR spectral data, Talanta 82 (5) (2010) 1784–1790.
- [8] K. Y. Chan, T. S. Dillon, C. K. Kwong, Modeling of a liquid epoxy molding process using a particle swarm optimization-based fuzzy regression approach, Industrial Informatics, IEEE Transactions on 7 (1) (2011) 148–158.
- [9] A. Gálvez, A. Iglesias, J. Puig-Pey, Iterative two-step genetic-algorithm based method for efficient polynomial b-spline surface reconstruction, Information Sciences 182 (1) (2012) 56–76.
- [10] K. Y. Chan, C. Kwong, T. S. Dillon, Development of product design models using fuzzy regression based genetic programming, in: Computational Intelligence Techniques for New Product Design, Springer, 2012, pp. 111–128.
- ²⁹⁴ [11] P. J. García Nieto, J. Alonso Fernández, F. de Cos Juez,
 F. Sánchez Lasheras, C. Díaz Muñiz, Hybrid modelling based on
 support vector regression with genetic algorithms in forecasting the cyanotoxins presence in the trasona reservoir (northern spain), Environmental
 research.
- [12] N. A. Barricelli, Numerical testing of evolution theories. part i: Theoretical introduction and basic tests, Acta Biotheoretica 16 (1-2) (1962) 69–98.
- [13] J. H. Holland, Adaptation in natural and artificial systems: An introductory analysis with applications to biology, control, and artificial intelligence., U Michigan Press, 1975.

- [14] J. R. Koza, Genetic Programming: vol. 1, On the programming of comput ers by means of natural selection, Vol. 1, MIT press, 1992.
- [15] Y. Bengio, Learning deep architectures for AI, Foundations and trends in
 Machine Learning 2 (1) (2009) 1–127.
- ³⁰⁸ [16] Y. Bengio, A. Courville, P. Vincent, Representation learning: A review and new perspectives.
- [17] D. Tarlow, I. Sutskever, R. S. Zemel, Stochastic k-neighborhood selection for supervised and unsupervised learning, Journal of Machine Learning Research.
- [18] K. Maertens, J. De Baerdemaeker, R. Babuška, Genetic polynomial regression as input selection algorithm for non-linear identification, Soft Computing 10 (9) (2006) 785–795.
- [19] T.-L. Yu, W.-K. Lin, Optimal sampling of genetic algorithms on polynomial
 regression, in: Proceedings of the 10th annual conference on Genetic and
 evolutionary computation, ACM, 2008, pp. 1089–1096.
- [20] C.-H. Wu, G.-H. Tzeng, R.-H. Lin, A novel hybrid genetic algorithm for kernel function and parameter optimization in support vector regression, Expert Systems with Applications 36 (3) (2009) 4725–4735.
- [21] M. Hofwing, N. Strömberg, M. Tapankov, Optimal polynomial regression models by using a genetic algorithm, in: Proceedings of the Second International Conference on Soft ComputingTechnology in Civil, Structural and Environmental Engineering Conference, (Crete, Greece), 2011009, 2011.
- [22] B. Cetisli, H. Kalkan, Polynomial curve fitting with varying real powers,
 Electronics and Electrical Engineering 112 (6) (2011) 117–122.
- [23] J. Davidson, D. A. Savic, G. A. Walters, Symbolic and numerical regression:
 Experiments and applications, Information Sciences 150 (1) (2003) 95–117.

- [24] O. Giustolisi, D. Savic, Advances in data-driven analyses and modelling
 using epr-moga, Journal of Hydroinformatics 11 (3-4) (2009) 225–236.
- [25] E. Willighagen, genalg: R based genetic algorithm (2012).
- [26] R Core Team, R: A language and environment for statistical computing.
 URL http://www.R-project.org/
- [27] D. Meyer, E. Dimitriadou, K. Hornik, A. Weingessel, F. Leisch, e1071:
 Misc functions of the department of statistics (e1071), tu wienR package
 version 1.6-1.
- URL http://CRAN.R-project.org/package=e1071
- ³³⁹ [28] T. Therneau, B. Atkinson, B. Ripley, rpart: Recursive partitioning R pack-³⁴⁰ age version 4.1-1.
- ³⁴¹ [29] T. Hothorn, K. Hornik, A. Zeileis, Unbiased recursive partitioning: A conditional inference framework, Journal of Computational and Graphical Statistics 15 (3) (2006) 651–674.
- [30] C. Strobl, A.-L. Boulesteix, A. Zeileis, T. Hothorn, Bias in random forest
 variable importance measures: Illustrations, sources and a solution, BMC
 Bioinformatics 8 (25).
- URL http://www.biomedcentral.com/1471-2105/8/25
- [31] C. Strobl, A.-L. Boulesteix, T. Kneib, T. Augustin, A. Zeileis, Conditional variable importance for random forests, BMC Bioinformatics 9 (307).
- URL http://www.biomedcentral.com/1471-2105/9/307