An evolutionary algorithm for global induction of regression and model trees

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Abstract: Most tree-based algorithms are typical top-down approaches that search only for locally optimal decisions at each node and does not guarantee the globally optimal solution. In this paper, we would like to propose a new evolutionary algorithm for global induction of univariate regression trees and model trees that associate leaves with simple linear regression models. The general structure of our solution follows a typical framework of evolutionary algorithms with an unstructured population and a generational selection. We propose specialised genetic operators to mutate and cross-over individuals (trees), fitness function that base on the Bayesian information criterion and smoothing process that improves the prediction accuracy of the model tree. Performed experiments on 15 real-life datasets show that proposed solution can be significantly less complex with at least comparable performance to the classical top-down counterparts.

Keywords: evolutionary algorithms; regression trees; model trees; SLR; linear regression; Bayesian information criterion; BIC.

Reference to this paper should be made as follows: Czajkowski, M. and Kretowski, M. (2013) 'An evolutionary algorithm for global induction of regression and model trees', *Int. J. Data Mining, Modelling and Management*, Vol. 5, No. 3, pp.261–276.

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1 Introduction

The most common predictive tasks in data mining (Fayyad et al., 1996) are classification and regression and the decision trees (Murthy, 1998; Rokach and Maimon, 2008) are one of the most widely used prediction techniques. Regression and model trees are now popular alternatives to classical statistical techniques like standard regression or logistic regression (Hastie et al., 2009). They are easy to understand and interpret which makes them particularly useful when the aim of modelling is to understand the underlying processes of the environment. Decision trees are also applicable when the data does not satisfy rigorous assumptions required by more traditional methods (Hastie et al., 2009). We focus on univariate trees since they are a 'white-box' technique and it makes them particularly interesting for scientific modelling. It is easy to find explanation for predictions of univariate regression and model trees.

1.1 Regression and model trees

Regression and model trees may be considered as a variant of decision trees, designed to approximate real-valued functions instead of being used for classification tasks. Main difference between regression tree and model tree is that, for the latter, constant value in the terminal node is replaced by a regression plane. One of the first and most known regression tree solution was presented in the seminal book by Breiman et al. (1984) describing the *CART* system. *CART* finds a split that minimises the sum of squared residuals of the model when predicting and builds a piecewise constant model with each terminal node fitted by the training sample mean. The accuracy of prediction was later improved by replacing single values in the leaves by more advanced models. M5 proposed by Quinlan (1992), induces a model tree that contains at leaves multivariate linear models analogous to piecewise linear functions. *HTL* presented by Torgo (1997) goes further and evaluate linear and non-linear models in terminal nodes. Model trees can also be applied to the classification problems (Kotsiantis, 2010).

Figure 1 An example of univariate decision tree with tests on nominal and continuous-valued features



Note: Depending on the tree type, leaves could contain class (classification tree), continuous value (regression tree) or some kind of model (model tree).

All aforementioned decision trees are built by a process that is known as recursive partitioning. Top-down induction starts from the root node where locally optimal split (test) is searched according to the given optimality measure. Then, the training data is redirected to newly created nodes and this process is repeated for each node until some stopping-rule is violated. Finally, the post-pruning is applied to improve the generalisation power of the predictive model.

1.2 Motivation

Inducing the decision tree by a greedy strategy is fast and generally efficient in many practical problems, but usually produces locally optimal solutions. It can be expected that a more global induction could improve the tree structure and the model prediction. Figure 2 illustrates two simple artificially generated datasets with analytically defined decision borders.





The left dataset *split plane2*, discussed also in Vogel et al. (2007), can be perfectly predictable with regression lines on subsets of the data resulting from a single partition. The equation is:

$$y = \begin{cases} 0 & -4 \le x_1 < -2\\ 0.25x_1 + 0.5 & -2 \le x_1 \le 2 \end{cases}$$
(1)

Most of popular greedy top-down inducers that minimises the residual sum of squares (RSS) like *CART* or standard deviation like *M5* will not find the best partitions (*CART* finds threshold at $x_1 = -0.44$, *M5* at $x_1 = -1.18$). Not optimal partition in the root node usually increases of the tree size and may result the higher prediction error.

Illustrated in Figure 2(b) function is defined as:

$$y = \begin{cases} x_1 + 1 & 0 \le x_1 \le 1\\ -x_1 + 6 & 4 < x_1 \le 5\\ -0.5x_2 + 1.5 & 1 < x_1 \le 4, \ 0 \le x_2 \le 3\\ 3x_2 - 9 & 1 < x_1 \le 4, \ 3 < x_2 \le 5 \end{cases}$$
(2)

It is a little more complex then *split plane2* and many traditional approaches will fail to efficiently split the data as the greedy inducers search only for a locally optimal

solutions. Figure 3 presents the optimal model trees that can be generated by globally induced and greedy top-down algorithms. These two simple artificial problems illustrate general advantage of the global search solutions to greedy algorithms.

Figure 3 Examples of model trees for armchair2, (a) global approach (b) greedy approach



1.3 Related work

Multiple authors have proposed methods to limit negative effects of inducing the decision tree with the greedy strategy. In the *SECRET* authors, Dobra and Gehrke (2002) show that classification approach finds more globally optimal partitions than the *CART* system. Different solution was proposed in *SMOTI* by Malerba et al. (2004) where regression models exist not only in the leaves but also in the upper parts of the tree. Authors suggest that this allows for individual predictors to have both global and local effects on the model tree. A more recent innovation in order to find optimal splits in nodes was presented in *LLRT* by Vogel et al. (2007). *LLRT* allows for a near-exhaustive evaluation of all possible splits in a node, based on the quality of fit of linear regression models in the resulting branches.

In the literature there are also some attempts of applying evolutionary approach for induction of regression trees. In *TARGET*, Fan and Gray (2005) propose to evolve a *CART*-like regression tree with simple genetic operators. Bayesian information criterion (*BIC*) (Schwarz, 1978) is used as a fitness function which penalises the tree for over-parametrisation. Experiments performed on two real datasets suggest that *TARGET* outperforms in the terms of mean squared error the *CART* solution. Much more advanced trees are presented in *GPMCC* where Potgieter and Engelbrecht (2008) make

use of a genetic algorithm to evolve multivariate non-linear models at the leaves. Authors managed to decrease the size of the trees comparing to commercial version of *M5* (Cubist) and neutral network called *NeuroLinear* (Setiono et al., 2002), however their system was outperformed in terms of predictive accuracy.

Currently, there are no sufficient solutions with a good trade-off between predictive performance and a model comprehensibility. Model trees with complex rules at the leaves or ensemble methods generate accurate predictions but are difficult to interpret by a human experts. On the other side, regression trees have lower predictive performance but higher model comprehensibility. Finally, performed experiments suggest that regression and model trees usually built overgrown trees and therefore are more difficult to analyse and interpret.

In this paper, we would like to present an evolutionary algorithm for global induction of regression and model trees. It fills the gap between simple regression trees and advanced but less comprehensible model trees. Previously performed research showed that global inducers are capable to efficiently evolve various types of classification trees: univariate (Kretowski and Grześ, 2005), oblique (Kretowski and Grześ, 2006) and mixed (Kretowski and Grześ, 2007). In our last paper, we applied a similar approach to obtain accurate and compact regression trees (Kretowski and Czajkowski, 2010) called *GRT* and we did preliminary experiments with the model trees (Czajkowski and Kretowski, 2010) called *GMT*. Our work covers the induction of univariate regression trees and model trees with simple linear models at the leaves. Proposed solution denoted as *GMT* 2.0 improved our previous solutions in almost every step of evolutionary algorithm. Starting with more heterogeneity population, additional genetic operators and new fitness function, that extends the BIC. We also introduced the smoothing process that could improve the prediction accuracy of the model tree.

2 Evolutionary induction of model trees

The GMT 2.0 general structure follows a typical framework of evolutionary algorithms (Michalewicz, 1996) with an unstructured population and a generational selection.

2.1 Representation

Model trees are represented in their actual form as classical univariate trees with a simple linear model at each leaf. Each test in a non-terminal node concerns only one attribute (nominal or continuous valued). In a case of a continuous-valued feature typical inequality tests are applied. As potential splits only precalculated candidate thresholds (Fayyad and Irani, 1992) are considered. A candidate threshold for the given attribute is defined as a midpoint between such a successive pair of examples in the sequence sorted by the increasing value of the attribute, in which the examples are characterised by different predicted values. Such a solution significantly limits the number of possible splits and focuses the search process. For a nominal attribute at least one value is associated with each branch. It means that an inner disjunction is built into the induction algorithm.

A simple linear model is calculated at each terminal node of the model tree using standard regression technique (Press et al., 1988). A dependent variable y is modelled as a linear function of single variable x:

$$Y = \alpha + \beta * x,\tag{3}$$

where x is one of the independent variables, α is the intercept and β is the slope of the regression line that minimises the sum of squared residuals of the model.

In every node information about learning vectors associated with the node is stored. This enables the algorithm to perform more efficiently local structure and tests modifications during applications of genetic operators.

2.2 Initialisation

Initial individuals are created by applying the classical top-down algorithm, similar to the *CART* and *M5* approaches. At first, we learn a standard regression tree that has mean of dependent variable values from training objects at each leaf. The recursive partitioning is finished when all training objects in a node are characterised by the same predicted value (or it varies only slightly, default: 1%) or the number of objects at node is lower than the predefined value (default value: 5). Additionally, user can set the maximum tree depth (default value: 10) to limit initial tree size. Next, a simple linear model is calculated at each terminal node of the model tree.

Traditionally, the initial population should be generated randomly to cover the entire range of possible solutions. Due to the large solution space the exhaustive search may be infeasible. Therefore, while creating initial population we search for a good trade off between a high degree of heterogeneity and relatively low computation time. We propose several strategies:

- initial individuals are created by applying the classical top-down algorithm to randomly chosen subsamples of the original training data (10% of data, but not more than 500 examples)
- for each individual only tests based on the random subset of attributes (default 50% of attributes) are applied
- at each individual for all non-terminal nodes one of the four test search strategies is randomly chosen:
 - 1 *least squares (LS)* function reduces node impurity measured by sum of squares proposed in *CART*
 - 2 *least absolute deviation (LAD)* function reduces the sum of absolute deviations. It has greater resistance to the influence of outlying values to *LS*
 - 3 *mean absolute error (MAE)* function which is more robust and also less sensitive to outliers to LS
 - 4 *dipolar*, where a dipol (a pair of feature vectors) is selected and then a test is constructed which splits this dipole. First instance that constitutes dipol is randomly selected from the node. Rest of the feature vectors are sorted decreasingly according to the difference between dependent variable values to the firstly chosen instance. To find a second instance that constitutes dipol we applied mechanism similar to the ranking linear selection (Michalewicz, 1996).

- one of three search strategies of predicted variable used in linear model at the leaves is applied:
 - 1 *optimal*: finds the locally optimal model that minimises the sum of squared residuals. It is the most time-consuming search strategy as it must calculate simple linear regression model for each attribute
 - 2 *random*: finds the simple linear model from training objects in this leaf on the randomly chosen independent variable
 - 3 *none*: the fastest strategy. No attribute is used to build the simple linear model, therefore each terminal node contains the sample mean.

Additionally, user can set the size of the population (default value: 50).

2.3 Genetic operators

To maintain genetic diversity, we have proposed two specialised genetic operators corresponding to the classical mutation and cross-over. Each evolutionary iteration starts with randomly choosing the operator type where default probability to select mutation equal 0.8 and cross-over 0.2. Both operators have influence on the tree structure, tests in non-terminal nodes and models at the leaves. After each operation it is usually necessary to relocate learning vectors between parts of the tree rooted in the altered node. This can cause that certain parts of the tree does not contain any learning vectors and has to be pruned. Modifying a leaf makes sense only if it contains objects with different dependent variable values.

2.3.1 Cross-over

Cross-over solution starts with selecting positions in two affected individuals. In each of two trees one node is chosen randomly. We have proposed three variants of recombination (Czajkowski and Kretowski, 2010):

- tests associated with the nodes are exchanged (only when non-terminal nodes are chosen and the number of outcomes are equal)
- subtrees starting in the selected nodes are exchanged
- branches which start from the selected nodes are exchanged in random order (only when non-terminal nodes are chosen and the number of outcomes are equal).

2.3.2 Mutation

Mutation solution starts with randomly choosing the type of node (equal probability to select leaf or internal node). Next, the ranked list of nodes of the selected type is created and a mechanism analogous to ranking linear selection is applied to decide which node will be affected. Depending on the type of node, ranking take into account:

• Location (level) of the internal node in the tree – it is evident that modification of the test in the root node affects whole tree and has a great impact, whereas mutation of an internal node in lower parts of the tree has only a local impact. Therefore, internal nodes in lower parts of the tree are mutated with higher probability.

• Absolute error – worse in terms of prediction accuracy leaves and internal nodes are mutated with higher probability (homogenous leaves are not included).

We have proposed new variants of mutation for internal node:

- node can be transformed (pruned) into a leaf
- tests between father and son exchanged
- mutation between subtrees that replaces all subtrees with randomly chosen one
- test in node reinitialised by new random or new dipolar one (described in Section 2.2)
 - 1 shifting the splitting threshold at continuous-valued feature
 - 2 re-grouping nominal feature values by adding, merging branches or moving value between them.

and for the leaf:

- transform leaf into an internal node with a new dipolar test
- replace simple linear model by a new one that is recalculated on a random predictor variable
- remove predictor variable and leave mean value at the leaf.

2.4 Selection and termination condition

Ranking linear selection is applied as a selection mechanism. Additionally, in each iteration, single individual with the highest value of fitness function in current population in copied to the next one *(elitist strategy)*.

Evolution terminates when the fitness of the best individual in the population does not improve during the fixed number of generations (default value: 1,000). In case of a slow convergence, maximum number of generations is also specified (default value: 5,000), which allows to limit the computation time.

2.5 Fitness function

Specification of a suitable fitness function is one of the most important and sensitive element in the design of the evolutionary algorithm. It measures how good a single individual is in terms of meeting the problem objective and drives the evolutionary search process. Direct minimisation of the prediction error measured on the learning set usually leads to the overfitting problem. In a typical top-down induction of decision trees (Rokach and Maimon, 2008), this problem is partially mitigated by defining a stopping condition and by applying a post-pruning (Esposito et al., 1997).

Fitness is computed for all members of the population after each generation. In our previous work (Czajkowski and Kretowski, 2010), we used Akaike's (1974) information criterion (AIC) as a fitness function. This measure of the goodness of fit worked also as a penalty for increasing the tree size. AIC is given by:

$$Fit_{AIC}(T) = -2 * ln(L(T)) + 2 * k(T),$$
(4)

where L(T) is the maximum of the likelihood function of the tree T and k(T) is the number of model parameters in the tree. Log(likelihood) function L(T) is typical for regression models (Gagne and Dayton, 2002) and can be expressed as

$$ln(L(T)) = -0.5n * [ln(2\pi) + ln(SS_e(T)/n) + 1],$$
(5)

where $SS_e(T)$ is the sum of squared residuals of the tree T and n is the number of observations. The term k(T) can also be viewed as a penalty for over-parametrisation. This complexity penalty term was set to Q(T) + 1 in where Q(T) is equal to the number of terminal nodes in model tree T.

We tested also a measure introduced by Schwarz (1978) called *BIC* that seems to be more appropriate for the regression and model trees. In this information criterion, which is similar to *AIC*, the penalty for increasing model size depends on the n – number of observations in the data:

$$Fit_{BIC}(T) = -2 * ln(L(T)) + ln(n) * k(T).$$
(6)

However, performed research reveal that both information criteria in their base form were not able to find an optimal structure of GMT 2.0. Measures worked sufficiently good when the probability of mutation for leaves to transform into internal nodes was very low or equal zero. Higher probability of transforming leaves into the internal nodes caused rapid increase of size and error of the searched structure. However, not including this mutation operator strongly limits variants of the evolution of the tree structure.

In this paper, we propose a new fitness function which extends the *BIC*. The number of independent parameters in the complexity penalty term k(T) for *GMT* 2.0 was set to 2(Q(T) + W(T)) where W(T) is the number of attributes in the linear models at the leaves (equal 1 for model node or 0 for regression node). High value of penalty term, compared to our previous solution or the *TARGET* system allow *GMT* 2.0 to induce significantly smaller trees.

Performed research in determining appropriate value of the penalty term k(T) suggests that the modification of the number of model parameters in the tree is only a partial solution. Higher value of k(T) impact data with high and low value of likelihood function in a different way and therefore it is not universal. Complexity penalty term has the highest effect when the sum of squared residuals $SS_e(T)$ of the tree is high because of the logarithm function. Small value of fraction $SS_e(T)/n$ results in high value of likelihood function which makes $Fit_{BIC}(T)$ less sensitive to the penalty term k(T). To obtain fitness function that is not sensitive to the various values of the likelihood function, we multiplied the $Fit_{BIC}(T)$ by, as we call it, the *tree size factor*. This is additional complexity penalty tries to balance the penalty for the small and large datasets. The *tree size factor* is denoted as $\varrho(T)$ and can be expressed as:

$$\varrho(T) = \frac{n + Q(T)}{n - Q(T)}.\tag{7}$$

Therefore, the complete fitness function equation for GMT 2.0 is given by:

$$Fit_{GMT\ 2.0}(T) = \begin{cases} Fit_{BIC}(T) * \varrho(T) \text{ when } Fit_{BIC}(T) \ge 0\\ Fit_{BIC}(T) * \frac{1}{\varrho(T)} \text{ when } Fit_{BIC}(T) < 0 \end{cases}$$

$$(8)$$

The best single individuals are the ones with the lowest $Fit_{GMT 2.0}(T)$ value. The *tree size factor* $\rho(T)$ increases the value of the fitness function and depends on the number of observation and leaves.

2.6 Smooting

In M5 algorithm, Quinlan (1992) proposed the smoothing process to improve the prediction accuracy of the tree-based models. The smoothing process modify the predicted by a model at the appropriate leaf, value of each case to reflect the predicted values at nodes along the path from that leaf to the root. It requires to generate additional linear models for every internal node of the tree. In the GMT 2.0 we developed the form of smoothing that is similar to the one in M5 algorithm. At first, predicted value for a test instance is computed by the leaf model. Then, this value is smoothed and updated along the path back to the root by linear models for each nodes. Let $P(T_i)$ denote the predicted value at T_i subtree of tree T, then:

$$P(T) = \frac{n_i * P(T_i) + k * M(T)}{n_i + k},$$
(9)

where n_i is the number of training cases at T_i , M(T) is the predicted value recalculated from the linear model at T and k is a smoothing constant (default 10).

Figure 4 illustrates the smoothing process for the test instances at the leaf with linear model denoted as LM4. If there were no smoothing process, the predicted value P(T) for a tested instance would be equal the value calculated from the model LM4. However, with the smoothing process turned on, the models that are on the path from the leaf to the root (LM5 and LM6) have influence on the final predicted value P(T).

According to Quinlan (1992) smoothing has most effect when some models were constructed for few training cases or when the models along the path predict instances very differently. However trees that adapt smoothing differs from the classical univariate model trees. Each test instance is predicted not only by single model at proper leaf but also by different linear models generated for every internal node up to the root node. Therefore smoothing affects the simplicity of the solution making it more difficult to understand and interpret.



Figure 4 The smoothing process for the test instances at the leaf with linear model denoted as LM4

3 Experimental validation

Two sets of experiments were performed – one for the regression trees and second for the model trees. GMT 2.0 was validated on several real-life datasets. Obtained results were compared with our previous solutions (Kretowski and Czajkowski, 2010; Czajkowski and Kretowski, 2010) and popular regression and model trees that are available in the *Weka* system (Hall et al., 2009).

3.1 Setup

To assess the performance of the proposed system in solving real-life problems, several datasets from UCI machine learning repository (Blake et al., 1998) and provided by Torgo (2010) were analysed. Table 1 presents the details of each dataset. All results presented in this paper correspond to averages of 20 runs and were obtained by using test sets (when available) or by 10-fold cross-validation. Root mean squared error (RMSE) is given as the prediction error measure of the tested systems. The number of nodes is given as a complexity measure (size) of regression and model trees. Each tested algorithm runs with default values of parameters through all datasets.

Table 1	Charac	teristics	of	the	real-life	datasets

	Dataset	Number of features		
Name	Symbol	Number of instances	Numeric	Nominal
Abalone	AB	4,177	7	1
Ailerons	AI	13,750	40	0
Auto-Mpg	AM	392	4	3
Auto-Price	AP	159	14	1
Delta Ailerons	DA	7,129	5	0
Delta Elevators	DE	9,517	6	0
Elevators	EL	16,599	18	0
Housing	HO	506	13	0
Kinemaics	KI	8,192	8	0
Machine CPU	MC	209	6	0
Pole	PO	15,000	48	0
Pyrimidines	PY	74	27	0
Stock	ST	950	9	0
Triazines	TR	186	60	0
Wisconsin Cancer	WC	194	32	0

3.2 Regression trees

Regression trees are used for analysis that require simple predictions based on a few logical if-then conditions. However, most solutions induce overgrown regression trees which are difficult to analyse. Domain experts need solutions that are smaller and therefore easier to understand. Our main goal in this set of experiments was to decrease the tree size of our previous solution called *GRT* without significant increase of prediction error. It is expected that changes in the complexity penalty term k(T) influence not only the tree size but also the prediction error of *GMT* 2.0. For a comparison purpose, we have tested four regression tree systems:

- $GMT \ 2.0_{reg}$ proposed solution, set to work as a regression tree.
- *GRT* one of the predecessors of *GMT* 2.0. Globally induced regression tree proposed in Kretowski and Czajkowski (2010).
- *REPTree* popular top-down inducer. *REPTree* builds a regression tree using variance and prunes it using reduced-error pruning (with backfitting).
- $M5_{reg}$ state of art model tree proposed by Quinlan (1992), set to work as a regression tree.

Dataset	GMT 2.0		GRT		_	REPTree			$M5_{reg}$		
Duiusei	RMSE	Size	RMSE	Size		RMSE	Size	-	RMSE	Size	
AB	2.33	3.5	2.31	51		2.35	201		2.28	36	
AI	0.000213	19	0.000217	27		0.000203	553		0.000199	166	
AM	3.96	2.1	3.57	45		3.6	94		3.49	19	
AP	2542	2.0	2618	13		2760	32		2543	8.0	
DA	0.000179	7.4	0.000179	82		0.000175	291		0.00176	74	
DE	0.00150	7.9	0.00148	78		0.00150	319		0.00148	59	
EL	0.00435	22	0.00443	32		0.00398	503		0.00413	189	
HO	4.51	6.5	4.17	32		4.84	41		4.72	26	
KI	0.194	25	0.194	34		0.191	819		0.182	264	
MC	74.2	2.7	63.9	15		92.34	15		64.8	10	
PO	9.91	19	10.32	25		8.25	223		9.32	139	
PY	0.104	4.1	0.101	10		0.135	1.0		0.135	1.0	
ST	1.41	13	1.33	39		1.19	137		1.14	88	
TR	0.142	4.9	0.139	14		0.152	7.0		0.140	5.0	
WC	33.3	1.9	39.2	16		35.9	9.0		35.0	3.0	

Table 2 Obtained results for the regression trees

Table 2 presents results for the regression trees. GMT 2.0 alike GRT managed to induce significantly smaller trees compared to the tested counterparts. This can be especially noticed on large datasets. Almost all GMT 2.0 trees are smaller and therefore easier to analyse and interpret. The only exception appears in the *Pyrimidines (PY)* dataset where globally induced trees are little more complex to the tested counterparts however have significantly higher prediction accuracy. This suggests that greedy algorithms like $M5_reg$ and REPTree underfitted to the training data and did not capture the underlying structure.

The average prediction error of GMT 2.0 is similar to the GRT and REPTree however it is slightly worse than $M5_{reg}$. Comparing to our previous solution, GMT 2.0 managed to significantly decrease tree size in all datasets for over 70% (average). In the same time, in 7 out of 15 datasets the prediction error RMSE for GMT 2.0 decreased, compared to GRT or stayed on the same level. Lack of improvement on some datasets may be explained by the significantly smaller trees induced by the GMT 2.0. There is usually a trade-off between the predictive performance and the model comprehensibility. Additional experiments showed that the GMT 2.0 with lower value of the parameter k(T) managed to induce larger but much more accurate regression trees. Modification of this penalty term allows to fine tune the decision tree algorithm.

3.3 Model trees

Model trees which are an extension of the regression trees, usually have higher performance in the terms of the accuracy prediction. However, model trees like *HTL* (Torgo, 1997) or *SMOTI* (Malerba et al., 2004) build complex models at the leaves that reduces simplicity of the predictions. Therefore in this set of experiments we focus on comparing the model trees with simple linear regression models at the leaves:

- GMT 2.0 proposed solution with no smoothing.
- *GMT* one of the predecessors of *GMT* 2.0. Globally induced model tree with simple linear regression models at the leaves proposed in Czajkowski and Kretowsk (2010).
- $M5_{slr}$ state of art model tree system proposed by Quinlan (1992), set to work with simple (instead of multivariate) linear regression model at the leaves.

We may observe from Table 3 that GMT 2.0 alike GMT managed to induce significantly smaller trees to the tested counterparts. Research showed that the sizes of induced GMT and GMT 2.0 trees are similar. However, in this set of experiment we focus on improving of the GMT prediction power. We managed to reduce RMSEerror, comparing to our previous solution, in 14 out of 15 datasets for GMT 2.0. Comparing to the $M5_{slr}$, proposed solution managed to not only significantly decrease tree size but also reduce the prediction error in most of the datasets.

Table 3	Comparison	results fo	r the	model	trees	with	simple	linear	regression	models	at t	he	leaves
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Dataset	GMT 2	.0	GMT		$M5_{slr}$		
Duiusei	RMSE	Size	RMSE	Size	RMSE	Size	
AB	2.24	6.7	2.30	7.7	2.24	35	
AI	0.000200	24	0.000207	18	0.000194	109	
AM	3.23	4.7	3.43	9.9	3.35	11	
AP	2328	2.9	2507	3.7	2183	6	
DA	0.000173	13	0.000178	11	0.000170	46	
DE	0.00147	8.6	0.00150	9	0,00148	40	
EL	0.00397	40	0.00444	13	0.00386	174	
НО	4.21	6.6	4.32	9.1	4.36	21	
KI	0.183	34	0.196	20	0,178	196	
MC	63.4	6.1	67.5	3.8	89.5	7.0	
PO	9.37	67	12.41	12	10.28	108	
PY	0.103	4.4	0.109	4.5	0.118	3.0	
ST	1.22	18	1.63	7.1	1.08	64	
TR	0.142	4.9	0.141	4.7	0.141	4.0	
WC	32.7	1.0	34.3	3.1	33.8	1.0	

3.4 Smoothed model trees

The smoothing process often improves the prediction accuracy of the tree-based models. Table 4 illustrates the impact of the smoothing function on GMT 2.0 and $M5_{slr}$ solutions. We may observe that both algorithms managed to slightly improve the prediction accuracy on most of the datasets. Low impact of smoothing function and weaker improvement of *RMSE* on proposed solution comparing to the $M5_{slr}$ smot may result from smaller and more optimal GMT 2.0 tree structure that cannot be so efficiently adjusted.

Dataset	GMT 2.0	smot	$M5_{slr}$ st	not
	RMSE	Size	RMSE	Size
AB	2.23	6.7	2.21	35
AI	0.000200	24	0.000186	109
AM	3.18	4.7	3.22	11
AP	2243	2.9	2282	6
DA	0.000172	14	0.000169	46
DE	0.00146	8.8	0.00147	40
EL	0.00393	39	0.00366	174
НО	4.07	6.9	4.07	21
KI	0.182	34	0.172	196
MC	62.6	5.8	87.6	7.0
PO	9.37	61	9.61	108
PY	0.093	4.3	0.115	3.0
ST	1.22	17	1.22	64
TR	0.132	5.0	0.137	4.0
WC	32.7	1.0	33.5	1.0

 Table 4
 Comparison results for the smoothed model trees with simple linear regression models at the leaves

Table 4 results shows that smoothing process may have also a negative impact on the final prediction. In *stock* and *auto-price* dataset, the *RMSE* calculated for the $M5_{slr}$ smot has increased. None of this happen to GMT 2.0 smot.

3.5 Calculation time

As with most evolutionary algorithms, calculation time of the proposed approach is more time consuming than the classical top-down inducers. Performed experiments with the dual-core CPU 1.66 GHz with 2 GB RAM on the dataset *Elevators* (16,559 instances, 18 attributes) showed that time:

- for regression trees: $M5_{reg}$ equal 7 seconds, GMT 2.0 equal 1.5 minutes
- for model trees: $M5_{slr}$ equal 11 seconds, GMT 2.0 equal 33 minutes
- for smoothed model trees: no relevant time differences.

Difference between $M5_{slr}$ and GMT 2.0 is caused by the evolutionary evaluation of linear models at the leaves. However, proposed solution is scalable and can manage large datasets.

4 Conclusions

Regression trees and model trees with simple linear models at the leaves are important 'white box' solutions. In this paper, we propose a new global approach to the model tree learning and compare it with classical top-down inducers. The structure of the GMT 2.0 tree, tests in non-terminal nodes and models at the leaves are searched in the same time by specialised evolutionary algorithm.

Experimental results show that the globally evolved regression models are highly competitive compared to the top-down-based counterparts, especially in the term of

tree size. GMT 2.0 managed to significantly improve our previous solution: GRT regression trees in the term of size and GMT in the predictive accuracy.

Proposed solution may be applied to the problems that are primarily concerned with the regression of an outcome onto a single predictor. As an example the original genetic epidemiology problem required only consideration of simple linear regression models like (Shannon et al., 2002) to locate genes associated with a quantitative trait of interests. GMT 2.0 is constantly improved. We plan to introduce oblique tests in the non-terminal nodes and more advance models at the leaves. We also plan to parallelise the evolutionary algorithm in order to speed-up its execution time.

Acknowledgements

This work was supported by the grant S/WI/2/08 from Bialystok University of Technology.

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