1. Introduction

The task of learning useful models from available data is common in virtually all fields of science, engineering, and finance. The goal of the learning task is to estimate unknown (input, output) dependency (or model) from training data (consisting of a finite number of samples) with good prediction (generalization) capabilities for future (test) data (Cherkassky & Mulier, 2007; Hastie et al., 2003). One of the specific learning tasks is regression – estimating an unknown real-valued function. The process of regression model learning is also called regression modelling or regression model building.

Many practical regression modelling methods use basis function representation – these are also called dictionary methods (Friedman, 1994; Cherkassky & Mulier, 2007; Hastie et al., 2003), where a particular type of chosen basis functions constitutes a “dictionary”. Further distinction is then made between non-adaptive methods and adaptive (also called flexible) methods.

The most widely used form of basis function expansions is polynomial of a fixed degree. If a model always includes a fixed (predetermined) set of basis functions (i.e. they are not adapted to training data), the modelling method is considered non-adaptive (Cherkassky & Mulier, 2007; Hastie et al., 2003). Using adaptive modelling methods however the basis functions themselves are adapted to data (by employing some kind of search mechanism). This includes methods where the restriction of fixed polynomial degree is removed and the model’s degree now becomes another parameter to fit. Adaptive methods use a very wide dictionary of candidate basis functions and can, in principle, approximate any continuous function with a pre-specified accuracy. This is also known as the universal approximation property (Kolmogorov & Fomin, 1975, Cherkassky & Mulier, 2007).

However, in polynomial regression the increase in the model’s degree leads to exponential growth of the number of basis functions in the model (Cherkassky & Mulier, 2007; Hastie et al., 2003). With finite training data, the number of basis functions along with the number of model’s parameters (coefficients) quickly exceeds the number of data samples, making model’s parameter estimation impossible. Additionally the model should not be overly

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complex even if the number of its basis functions is lower than the number of data samples, as too complex models will overfit the data and produce large prediction errors.

To obtain a polynomial regression model that does not overfit (nor underfit) and describes the relations in data sufficiently well, typically the subset selection approach (Hastie et al., 2003; Reunanen, 2006) is used where the goal is from a fixed full predetermined dictionary of basis functions to find a subset which corresponds to a model (a sparse polynomial) with the best predictive performance. This is done via combinatorial optimization. However, for the subset selection approach still the two issues remain – deficiency of adaptation as well as computational inefficiency.

Searching through all the possible combinations of basis functions takes double-exponential runtime as the number of combinations grows exponentially in the number of basis functions of the predetermined dictionary while the number of the basis functions in the dictionary grows exponentially in the number of input variables and “full” model’s degree (Hastie et al., 2003). This makes the exhaustive search through all the combinations impractical. The heuristic greedy search algorithms, such as forward selection (Hastie et al., 2003; Reunanen, 2006), substantially reduce the time and make it practical for not too large number of input variables and not too high degree. Nevertheless, the search time actually is still exponential, hindering their use in problems of larger dimensionality and hindering the removal of the restriction of a fixed degree.

The approach of subset selection assumes that the chosen fixed finite dictionary of the predefined basis functions contains a subset that is sufficient to describe the target relation sufficiently well. However, in most practical situations the required dictionary (and “full” model’s degree) is not known beforehand and needs to be either guessed or found by an additional search loop over the whole model building process, since it will differ from one regression task to another. In many cases, especially when the studied data dependencies are complex and not well studied, this means either a non-trivial and long trial-and-error process or acceptance of a possibly inadequate model.

This chapter presents a sparse polynomial regression model building approach which enables adaptive model building without restrictions on model’s degree and does it in polynomial time instead of exponential time (in the number of input variables, required degree, and target model’s complexity) as well as without the requirement to repeat the model building process. The required basis functions are automatically iteratively constructed using heuristic search specifically for the particular data at hand instead of choosing a subset from a very restricted finite user-defined dictionary (hence the approach is called Adaptive Basis Function Construction, ABFC). The basis function dictionary now becomes infinite and polynomials of arbitrary complexity can be generated bringing the desired flexibility to the model building process.

The remainder of this chapter is organized as follows. The next two sections give brief overview of polynomial regression and the subset selection approach. In Section 4 the ABFC approach is described. Section 5 outlines the related work. The results of the empirical evaluations of the proposed methods and their comparison to other well-known regression modelling methods are presented in Section 6. Section 7 concludes this chapter.
2. Polynomial regression

In standard regression formulation (Vapnik, 1995; Cherkassky & Mulier, 2007; Hastie et al., 2003) the goal is to estimate unknown real-valued function in the relationship

\[ y = G(x) + \varepsilon, \]  

where \( \varepsilon \) is independent and identically distributed random noise with zero mean, \( x = (x_1, x_2, ..., x_d) \) is \( d \)-dimensional input, and \( y \) is scalar output. The estimation is made based on a finite number of samples (training data) provided in form of matrix \( x \) of input values for each sample and vector \( y \) of output values for each corresponding sample. Using the finite number \( n \) of training samples \( (x_j, y_j), j = 1, 2, ..., n \) one wants to build a model \( F \) that allows predicting the output values for yet unseen input values as closely as possible.

Generally, a linear regression model may be defined as a linear expansion of basis functions:

\[ F(x) = \sum_{i=1}^{k} a_i f_i(x), \]  

where \( a = (a_1, a_2, ..., a_k)^T \) are model’s parameters, \( k \) is the number of basis functions included in the model (equal to the number of model’s parameters), and \( f_i(x), i = 1, 2, ..., k \) are the included basis functions of the input \( x \). As the model is linear in the parameters, the estimation of its parameters is typically done using the Ordinary Least-Squares (OLS) method (Hastie et al., 2003) minimizing the squared-error:

\[ a = \arg \min_{a} \sum_{j=1}^{n} (y_j - F(x_j))^2. \]  

The basis function representation enables moving beyond pure linearity, by defining nonlinear transformations of \( x \) while still working with linear models (and employing OLS). For example, for \( d = 1 \) a polynomial model of fixed degree \( p \) can be defined as follows:

\[ F(x) = \sum_{i=0}^{p} a_i x^i. \]  

Generally for a given \( d \) and \( p \) the total number of basis functions in a “full” polynomial, i.e. the total number of basis functions in the dictionary, is

\[ m = \prod_{i=1}^{p} (1 + d / i). \]
3. Subset selection

Models which are too complex (i.e. that fit the training data too well causing overfitting) or too simple (i.e. that fit the data poorly causing underfitting) provide poor predictive performance for the future data. The most popular approach of controlling model’s complexity is subset selection. The goal of subset selection is from a fixed full predetermined dictionary of basis functions to find a subset that provides the best predictive performance of the model (Hastie et al., 2003; Reunanen, 2006). Now in addition to the estimation of model’s parameters, the structure of the model itself needs to be found.

The total number of possible subsets from a dictionary of size \( m \) is \( 2^m \). This means that searching through all the possible subsets is in most cases impractical. Hence in subset selection heuristic search algorithms are used. They efficiently traverse the space of subsets, by adding and deleting basis functions of the model, and use model evaluation measure to direct the search into areas of increased performance. The typical examples of heuristic search algorithms are the greedy hill-climbing algorithms – Forward Selection (also known as Sequential Forward Selection, SFS) and Backward Elimination (also known as Sequential Backward Selection, SBS) (Hastie et al., 2003; Reunanen, 2006). However, there exist also more recently developed search strategies, such as Beam Search, Floating Search, Simulated Annealing, Genetic Algorithms etc. (Reunanen, 2006; Pudil et al., 1994; Russel & Norvig, 2002).

Summarizing (Russel & Norvig, 2002; Molina et al., 2002; Kohavi & John, 1997), in order to characterize a heuristic search problem one must define the following: 1) initial state of the search; 2) available state-transition operators; 3) search strategy; 4) evaluation measure; 5) termination condition. Note that in the context of model building the “initial state” is also called “initial model” and the “state-transition operators” are also called “model refinement operators”.

In the subset selection approach for polynomial regression, typically the initial state is the model that corresponds to the empty subset, the subset with only the intercept term in it, full set of all the defined basis functions, or a randomly chosen subset. The typical basic state-transition operators are addition and deletion of a basis function. The typical search strategy is the hill-climbing (Russel & Norvig, 2002) which, in combination with the empty (or sufficiently small) subset as initial state and the addition operator, becomes SFS, but, in combination with the full subset as initial state and the deletion operator, becomes SBS. As the evaluation measures classically the statistical significance tests are used (Hastie et al., 2003; Dreyfus & Guyon, 2006). However, in model building currently two other strategies predominate (Cherkassky & Mulier, 2007; Dreyfus & Guyon, 2006): employment of complexity penalization criteria (also known as analytical criteria), e.g., the well-known Akaike’s Information Criterion, AIC (Akaike, 1974; Burnham & Anderson, 2002), and the resampling techniques, e.g., Hold-Out, Cross-Validation (CV), and Bootstrap (Kohavi, 1995; Hastie et al., 2003; Dreyfus & Guyon, 2006). The termination condition typically corresponds to finding of a state in that none of the state-transition operators can lead to a better state (i.e. a local minimum).

In polynomial regression, increase in the model’s degree leads to exponential growth of the number of basis functions in the dictionary, i.e. \( O(m) = O(d^n) \) (Cherkassky & Mulier, 2007; Hastie et al., 2003) and to double-exponential growth of the number of all possible subsets (or the number of states in the state space): \( O(2^m) = O(2^{d^2}) \). When using one or both of the
two basic state-transition operators, the order of the branching factor of a state in the state space in the very first iteration of the search is already equal to the number of basis functions in the dictionary, i.e. it also increases exponentially. Assuming that the “best” model found in the search process includes a total of \( k_* \) basis functions and that in each iteration the number of basis functions of the current model is increased by 1, the total number of evaluated models (subsets) is of order

\[
O\left( \sum_{i=1}^{k_*} d^p \right) = O(d^p k_*). \tag{6}
\]

Hence for larger values of \( d \) and \( p \) (e.g., when \( m \) reaches thousands) subset selection is rendered impractical. Additionally, because of the branching factor’s direct dependence on the number of basis functions in the dictionary, the idea of unrestricted degree (i.e. dictionary of infinite size) is hardly applicable. The computational problem could be somewhat reduced by choosing a sufficiently small but useful value of \( p \) before the actual model building is performed. However, generally the required maximal degree is not known beforehand and needs to be either guessed or found by additional search loop over the whole model building process, since it will differ from one regression task to another, which means either a non-trivial and long trial-and-error process or acceptance of a possibly inadequate model.

4. Adaptive Basis Function Construction

This section introduces Adaptive Basis Function Construction – a possible alternative to the classical subset selection approach. The goal of the ABFC approach is to overcome some of the limitations associated with the subset selection, outlined in the previous section. The ABFC approach is developed for sparse polynomial regression model building without restrictions on model’s degree, enables model building in polynomial time, and does not require repetition of the building process (in contrast to the subset selection approach). The required basis functions are automatically adaptively constructed specifically for data at hand, without using a restricted fixed finite user-defined dictionary. The dictionary in the ABFC is infinite and polynomials of arbitrary complexity can be constructed.

4.1 The models and the basis functions

Generally, a basis function in a polynomial regression model can be defined as a product of original input variables each with an individual exponent:

\[
f_i(x) = \prod_{j=1}^{d} x_j^{r_{ij}}, \tag{7}
\]

where \( \mathbf{r} \) is a \( k \times d \) matrix of nonnegative integer exponents such that \( r_{ij} \) is the exponent of the \( j \)th variable in the \( i \)th basis function. Note that, when for a particular \( i \)th basis function \( r_{ij} = 0 \) for all \( j \), the basis function is the intercept term.
Given a number of input variables $d$, matrix $r$, with a specified number of rows $k$ and with specified values for each of its elements, completely defines the structure of a polynomial model with all its basis functions. The set of basis functions, included in a model, is then

$$f = \left\{ \prod_{j=1}^{d} x_j^{r_{ij}} \mid i = 1, 2, \ldots, k \right\}. \quad (8)$$

For example, if $d = 3$ and $k = 4$, then the matrix

$$r = \begin{bmatrix} 0 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 1 & 3 \\ 1 & 1 & 1 \end{bmatrix} \quad (9)$$

corresponds to the set

$$f = \{x_1 x_2 x_3, x_1 x_2 x_3, x_1 x_2 x_3, x_1 x_2 x_3\} = \{1, x_1, x_2 x_3, x_1 x_2 x_3\}, \quad (10)$$

which in turn corresponds to the model

$$F(x) = a_1 + a_2 x_1 + a_3 x_2 x_3 + a_4 x_1 x_2 x_3. \quad (11)$$

Formally, the problem of finding the best set of basis functions can be defined as finding the best matrix $r$ with the best combination of nonnegative integer values of its elements:

$$r^* = \arg \min_r \left\{ \left\{ \prod_{j=1}^{d} x_j^{r_{ij}} \mid i = 1, 2, \ldots, k \right\} \right\}, \quad (12)$$

where $J(.)$ is an evaluation criterion that evaluates the predictive performance of the regression model which corresponds to the set of basis functions.

As neither the upper bounds of $r$ elements’ values nor the upper bound of $k$ are defined, it is possible to generate sparse polynomials of arbitrary complexity, i.e. of arbitrary number of basis functions each with an arbitrary exponent for each input variable. This also means that the searchable state space is infinite.

### 4.2 The search process

Finding the “best” structure of matrix $r$ requires search. In this section the five components (outlined in Section 3) of a heuristic search problem are analyzed in the context of the ABFC approach.

**Initial state.** In ABFC, the state space is infinite therefore a natural initial state of the search is the state that corresponds to the simplest model located in the space. In the current study it is assumed that the simplest model is the one with a single basis function corresponding to
the intercept term. However, also other models could be used as initial states, e.g., an empty model (without any basis functions), a first degree “full” polynomial, or a small randomly generated model. Note that in the current study the basis function corresponding to the intercept term stays in the model at all times and is not allowed to be modified or deleted.

*State-transition operators.* Using efficient state-transition operators is vital for the search process to be efficient. The employed state-transition operators are the main methodological difference between the subset selection approach and the ABFC approach. Generally, there are two different basic types of modifications to an existing polynomial model: complication and simplification (Jekabsons & Lavendels, 2008a). In the subset selection approach, these are the addition and deletion operators. The addition operator makes the model more complex (by adding a new basis function) but the deletion operator makes it simpler (by deleting an existing basis function).

In the ABFC, the two standard operators from subset selection are replaced with other operators that not only add or delete basis functions but also work on the level of individual exponents, modifying the existing basis functions and creating modified copies of them. The basic idea is to use an operator that adds only the simplest (i.e. linear) basis functions which serve as a basic material for further construction of more complex functions using other operators. In this manner there is no need for an operator that explicitly tries to add basis functions of each possible combination of exponent values (as the addition operator in the subset selection). Hence the branching factor of the state space stays not only finite but also relatively small while the state space itself is infinite.

In this study, a set of the following four state-transition operators for the polynomial regression model building are proposed. Operator1: Addition of a new linear basis function with one of its exponents set to one and all the others set to zero. Operator2: Addition of an exact copy of an already existing (in the current model) basis function with one of its exponents increased by 1. Operator3: Decreasing of one of the exponents in one of the existing basis functions by 1. Operator4: Deleting of one of the existing basis functions. Figure 1 gives examples of the operators operating on a simple matrix.

The set of the four state-transition operators is sufficient to generate any polynomial model definable by the matrix $r$. Their use can also be viewed as a piece of application-domain knowledge. While starting the search from the simplest model, the complication operators (the first two) do the main job – they “grow” the model. The simplification operators (the last two), on the other hand, work as “purifiers” – they decrease the unnecessarily high exponents and delete the unnecessary basis functions. Without the use of simplification operators, a regression model may contain unnecessarily high exponents and include too...
many unnecessary basis functions, at the same time preventing truly necessary modifications (this is also known as the nesting effect (Pudil et al., 1994)) and increasing overfitting. Additionally, for all the state-transition operators a special care is taken to prevent basis function duplicates in the resulting model as well as to preserve the intercept term.

The initial state and the state-transition operators together form a state space. Figure 2 shows a small example of a state space in ABFC when the number of input variables is three and all the four state-transition operators are used. Each state represents a set of basis functions included in the regression model. The ordering of the states in the space is such that the simplest models and the simplest basis functions are reached first and, as the search goes on, increasingly complex models and basis functions can be reached.

![State Space Example](image)

Fig. 2. A small example of the first three layers of a state space in ABFC when \( d = 3 \) (the space is infinite in the direction of more complex models)

In the Section 3, it is stated that in the subset selection approach the branching factor of a state in the state space increases exponentially with respect to the number of input variables \( d \) and pre-specified maximal degree \( p \). In ABFC, the branching factor of the current state in the state space depends on \( d \) and on the number of basis functions \( k \), already included in the current model. The upper bound of the number of possible modifications to a model using Operator1 is equal to \( d \); using Operator2 and Operator3 it is equal to \( dk \); and using Operator4 it is equal to \( k \). So the upper bound of the branching factor is of order \( O(d + 2dk + k) = O(dk) \) that is linear in respect to both \( d \) and \( k \).

**Search strategy.** Most of the heuristic search algorithms of the hill-climbing type can be divided in two categories: those that assume the model state-transition operators to be of either or both the forward and the backward type (e.g., SFS, SBS, and Floating Search algorithms) and those that do not distinguish between the two types (e.g., Steepest Descent Hill-Climbing and Simulated Annealing). The four operators proposed in this study are naturally divided in forward (complication) and backward (simplification) operators; therefore in ABFC both categories of the search algorithms can be applied.

On the other hand, non-hill-climbing search algorithms, e.g., Genetic Algorithms and the like, employ completely different kind of operators (i.e. Crossover and Mutation). While they could be adapted to work with the infinite dictionary of basis functions, their major disadvantage is that, in contrast to the simple hill-climbing algorithms, they are not generally biased towards simpler models. In large state spaces they often spend most of the time exploring too complex models while the “best” ones are in fact mostly the relatively simple ones.
**Evaluation measure.** The proposed state-transition operators allow using the same methods for model evaluation and comparison as those used in subset selection. However, note that the model complexity penalization criteria, in contrast to the resampling techniques, usually require substantially lower computational resources as well as are less noisy creating less local minima in the state space.

**Termination condition.** Many different termination conditions can be used to terminate the search process. Some of most widely used ones are the following: a) a user pre-specified number of iterations is reached; b) a user pre-specified size of the model is reached; c) using the available state-transition operators the model could not be improved any further (evaluated by the chosen evaluation measure). The first two termination conditions require the user to set a hyperparameter value. This is a non-trivial task as usually the required information is not available. Adjusting such a hyperparameter may also require too large amounts of computational resources. In this study, the termination condition listed here as the last (c) is employed.

### 4.3 A concrete practical model building method

This section proposes Floating Adaptive Basis Function Construction (F-ABFC) – a concrete practical polynomial regression model building method, which is a special case of the ABFC approach.

The search procedure of the F-ABFC starts with the simplest model (with only the intercept term included) and uses the Floating Search strategy (hence the name of the method), in particular the Sequential Floating Forward Selection algorithm, SFFS (Pudil et al., 1994), together with the set of the four state-transition operators proposed in the previous section.

In SFFS, the search process consists of two phases – the forward phase and the backward phase. In each iteration of the search, the forward phase is done only once but the number of times the backward phase is performed is determined dynamically. In the forward phase, all the models, which can be generated using the complication operators on the current best model, are evaluated and, if there is improvement over the current best model, the best of the new models is chosen as the new current best model and the search proceeds to the second phase. If there is no improvement, the whole search procedure is stopped. In the backward phase, on the other hand, all the models, which can be generated using the simplification operators on the current best model, are evaluated. In this phase ever simpler models are repeatedly generated and the phase is ended only when, using the available simplification operators, it is impossible to generate a model which is better than the current best one. After the second phase, the search process always proceeds to the next iteration (starting again with the first phase).

According to the studies of many researchers, the Floating Search algorithms, including SFFS, are some of the most efficient heuristic search algorithms for deterministic combinatorial optimization in terms of both required computational resources and quality of the results (Ferri et al., 1994; Jain & Zongker, 1997; Jain et al., 2000; Zongker & Jain, 1996; Pudil et al., 1994; Kudo & Sklansky, 2000; Reunanen, 2006). SFFS also does not have any adjustable hyperparameters, has a tendency to generate simpler models than many other algorithms, and is very simple to implement.

As in (Jekabsons & Lavendels, 2008a; Jekabsons, 2008), to evaluate the predictive performance of a newly generated model, to perform model comparisons, and to steer the
search in direction of the most promising models, in F-ABFC the Corrected Akaike’s Information Criterion, AICC (Hurvich & Tsai, 1989) is used. AICC is defined as follows:

\[
AICC = n \ln(MSE) + 2k + \frac{2k(k+1)}{n-k-1},
\]

where MSE is the Mean Squared Error of the model of interest in the training data. AICC evaluates model’s predictive performance as a trade-off between its accuracy in the training data (the first term of (13)) and its complexity (the last two terms of (13)). Calculation of the AICC for a single model requires a single estimation of model’s parameters using OLS and calculation of MSE in training data. The “best” model is that whose AICC value is the lowest.

The AICC is an improvement over the classical AIC (Akaike, 1974) with the third term in (13) added as a correction term intended for working with small-sized data sets. For problems with relatively small \( n \), AICC is suited better than AIC but converges to AIC as \( n \) becomes large (Hurvich & Tsai, 1989). AIC and AICC theoretical justification is based on the relationship between the Kullback-Leibner information and the maximum likelihood principle (Burnham & Anderson, 2002). Note that AIC as well as AICC does not assume that the “true model” (which was presumably used to generate the data) is one of the candidates (Burnham & Anderson, 2002).

In (Jekabsons & Lavendels, 2008b), an issue of the F-ABFC is stated, that, because the branching factor of the ABFC’s state space increases very slowly together with \( d \) and \( k \), in special cases when the data is of low dimensionality (e.g., \( d \leq 4 \)) and/or the existing structure in the data is very complex (i.e. a very complex model is required) the search algorithm may get stuck in a local minimum too early in the search returning a too simple and underfitted model.

As a remedy for this, here an additional recursion of the state-transition operators is proposed introducing one hyperparameter for the F-ABFC. The idea is to recursively create additional regression models from models already created from the current best model using the same state-transition operators with which they were initially created. This essentially means that if, for example, the recursion depth is set to 2, Operator1 will create not only linear basis functions but also basis functions of the second degree, Operator2 will create not only copies of basis functions with degree increased by 1 but also by 2, and Operator3 will not only try to decrease degrees by 1 but also by 2. However, as still none of the operators add more than one basis function to the model at a time, for the Operator4 the recursion is not used.

The recursion of the operators reduces the number of local minima in the state space which is especially important near the starting-point of the search (the initial model) and enables the search algorithm to find a much better model.

Presence of such a “recursion depth” hyperparameter is a disadvantage as now a user intervention might be required. However, for larger dimensionalities of the input space (when also the increased computational resources are required) it is reasonable to completely disable the recursion (by setting the hyperparameter equal to 1) as with large dimensionalities the branching factor increases sufficiently fast and the problem of too early local minima diminishes.
Figure 3 shows pseudo-code of F-ABFC’s search procedure. Note that in practical implementations of F-ABFC maintaining the set of the newly generated models (“MODELS”) is not required as a single model can be created, evaluated, and, if it turns out not to be an improvement, immediately discarded.

![Pseudo-code of F-ABFC’s search procedure](image)

**Fig. 3.** Pseudo-code of F-ABFC’s search procedure

In (Jekabsons & Lavendels, 2008a), a version of F-ABFC was developed that slightly differs from the one proposed here in that the method used one additional state-transition operator and the “recursion depth” hyperparameter was not introduced. The paper (Jekabsons & Lavendels, 2008a) empirically demonstrated the computational and predictive performance advantages of F-ABFC comparing to subset selection and a number of other popular regression modelling methods. F-ABFC advantages in real-world practical applications are demonstrated in (Kalnins et al., 2008a; Kalnins et al., 2009b) where it is applied for modelling bending and buckling behaviour of different composite material structures.
4.4 Computational considerations
Assuming that the “best” model found by the F-ABFC search procedure includes a total of \( k_* \) basis functions and in each iteration the number of basis functions in the current model is increased by 1, the total number of evaluated models is of order

\[
O \left( \sum_{i=1}^{k_*} d_i \right) = O \left( d_{k_*} \sum_{i=1}^{k_*} i \right) = O \left( d_{k_*} \frac{k_* (k_* + 1)}{2} \right) = O \left( d_{k_*}^3 + d_{k_*}^2 \right) = O \left( d_{k_*}^3 \right).
\]  

Consequently, relatively to the typical subset selection methods, the efficiency of the F-ABFC increases together with the increase in the number of input variables and in the required nonlinearity of the model (the value of \( p \)) but decreases together with the increase in the complexity \( k_* \) of the “best” found model. Moreover, the relative efficiency of the subset selection additionally substantially decreases in the common case when the required value of \( p \) is unknown and needs to be found by trying different values.

Using F-ABFC together with OLS, the associated linear least-squares fitting, required for a single model to be evaluated, demand computations of order \( O(nk^2 + k^3) \), where \( nk^2 \) operations are required for filling a \( k \times k \) matrix and \( k^3 \) operations are required for solving a linear equation system (Hastie et al., 2003). However, none of the proposed state-transition operators operate on more than one basis function of a model at a time meaning that, each time the parameters of a newly created model are calculated, only one row and one column of the \( k \times k \) matrix will change. Recalculating only the elements of the corresponding row and column, reduces the order of the computations to \( O(nk^3) \). Moreover, as the Operator4 does not modify any basis function (only deletes one), the order of the computations for this particular operator reduces further to \( O(k^3) \).

Yet it must be noted that the F-ABFC can still become computationally rather demanding, especially when the number of input variables and/or the number of samples in the training data gets very large. This is the price to pay for the high flexibility of the method.

4.5 Convergence of the search process
The F-ABFC’s search algorithm is cycle-free because a new model is allocated to “BestModel” (Figure 3) only if it is better than the old one (according to AICC). Moreover, as the AICC criterion tries to estimate model’s true predictive performance, the algorithm will seek for the best trade-off between too simple and too complex models and will stop somewhere in-between them. Additionally there is also a hard bound – the number of basis functions in a model will never exceed the number of samples in the training data as otherwise the OLS cannot estimate model’s parameters.

It should also be noted that, although the state space of F-ABFC is infinite, in practice the models of the best predictive performance are normally located in the part of the space that is relatively near to the initial state where all the models (and their basis functions) are relatively simple and do not yet neither overfit the data nor have basis functions more than samples in the training data. This also means that really only a small finite fraction of the whole infinite state space must be explored.
4.6 Selection bias, selection instability, and model averaging

There are two issues that to some extent plague all the methods of model building (including subset selection and ABFC), especially when working with relatively little data – selection bias and selection instability (also called selection variance). While the issues are attributable to virtually any model building method, they are commonly ignored frequently resulting in models of lower predictive performance.

Selection bias occurs when in the search procedure one uses the same data to compute model’s parameters, to perform model building (i.e. evaluation of candidate models, selection of the best one, and steering the search in direction of the most promising models), and to select the final “best” model which will be returned as the result of the model building process (Reunanen, 2003; Reunanen, 2006; Loughrey & Cunningham, 2004; Jekabsons, 2008). The problem is that the more candidates are visited during the search, the greater the likelihood of finding a model that has high accuracy in the training set while having a very low predictive performance (accuracy in the test set) (Reunanen, 2003; Reunanen, 2006; Kohavi & John, 1997; Loughrey & Cunningham, 2004). The random fluctuations in the data will improve the evaluations of some models more than others.

The problem is relevant regardless of the model evaluation measure used – statistical significance tests, complexity penalization criteria, or resampling techniques. In addition, the selection bias occurs even when performing model evaluation using completely independent validation data set (Kohavi & John, 1997; Reunanen, 2006). In any case, the more intensive (relative to the number of samples) is the search process, the larger is the selection bias, and, the larger is the noise in the data, the potentially larger is the harm (in terms of overfitting) done by the selection bias.

While the deterministic search algorithms of the hill-climbing type (including the SFFS algorithm of the F-ABFC) are usually less intensive and consequently more robust against overfitting than, for example, Simulated Annealing or Genetic Algorithms (Loughrey & Cunningham, 2004; Guyon & Elisseeff, 2003), the problem of selection bias remains relevant. The second issue, selection instability, is related to the fact that small perturbations of the data (deleting or adding samples, adding noise, rescaling the values) can lead the model building process to vastly different models. This is because the large variability of estimates of the evaluation methods can lead to different local minima (Breiman, 1996; Kotsiantis & Pintelas, 2004; Guyon & Elisseeff, 2003; Cherkassky & Mulier, 2007). This variance is undesirable because variance is often the symptom of a “bad” model that does not generalize well and because the model may be failing to capture the “whole picture” (Guyon & Elisseeff, 2003).

One of the ways to reduce both the selection bias and the selection instability, is to employ model combining (also called model ensembling or averaging) techniques (Breiman, 1996; Opitz & Maclin, 1999; Cherkassky & Mulier, 2007; Jekabsons, 2008). While a typical model building process usually consists in choosing only one best description for the data discarding the remainder, combining a number of models in some reasonable manner appears more reliably accurate as this can have the effect of smoothing out erratic models that overfit the data and gain more stability in the modelling process.

A typical model combination procedure consists of a two-stage process (Cherkassky & Mulier, 2007). In the first stage, a number of different models are constructed. The parameters of these models are then held fixed. In the second stage, these individual models are linearly combined to produce the final model.
Both stages can be done in different ways. In this study, to increase the predictive performance of models built by the F-ABFC, a CV-type resampling of the training data together with unweighted model averaging (Opitz & Maclin, 1999; Duin, 2002) is employed. As this resampling and model averaging works on top of the F-ABFC, the method is called Ensemble of Floating Adaptive Basis Function Construction (EF-ABFC). During resampling, the whole training data is randomly divided into \( v \) disjoint subsets (\( v \) typically being equal to 10). Then \( v \) overlapping training data sets are constructed by dropping out a different one of these \( v \) subsets. Such procedure is also employed to construct training sets for \( v \)-fold CV, so model ensembles constructed in this way are also called cross-validated committees (Parmanto et al., 1996).

Combining models via simple unweighted averaging requires them to be not too underfitted as well as not too overfitted (Duin, 2002). To lower the overfitting, in each CV iteration the unused 10th data subset is used as a validation data set for “re-evaluation” (using MSE) of the best models of each F-ABFC iteration and for selection of the one “final best” model from any iteration. Note that this validation set is never used for model evaluation during the search. Instead it is used strictly only for the “re-evaluation” and “re-selection” after the F-ABFC search process has already ended. Also note that as an evaluation measure in the search algorithm still the AICC is applied. This “re-evaluation” using the validation data set can detect whether the search process at some iteration may have started to generate overfitted models and select a model of some earlier iteration that is (hopefully) not (or at least less) overfitted (see Figure 4).

![Fig. 4. An example of how a less overfitted model is selected using “re-evaluation” in validation set. Note that here starting from the 35th iteration the AICC values also start to increase (in contrast to the training error which always decreases) however this might be too late due to selection bias](image)

The so far described process produces \( v \) models built by \( v \) independent F-ABFC runs each using a different combination of CV-partitioned data subsets. Next, the \( v \) models from the \( v \) CV iterations are combined using the unweighted model averaging. Note that prior to combining, all the models are re-fitted to the whole training data set (without the CV partitioning). This is done to compensate for the smaller training sets used during the individual model building.

Model combining by unweighted model averaging consists in taking an unweighted average of predictions of all the models:
where $F_i$ is $i$th individual model from the $i$th CV iteration and $F_{comb}$ is the combined model. For polynomial regression this simply means summation of all the polynomials and then a division of all the parameters of $F_{comb}$ (that is also a polynomial) by $v$. Note that the parameter values of $F_{comb}$ will not necessarily be optimal in the sense of the least-squares loss (in fact they will be optimal only in special cases, e.g., when all $F_i$'s are identical).

The employed model combining method is similar to Bagging (bootstrap aggregating (Breiman, 1996)) where the training set is bootstrapped (usually to build varied decision trees), and the unweighted average of the resulting models is taken.

Figure 5 gives an outline of the EF-ABFC model building process when the number of CV folds $v$ is three. Note however that for all the practical applications of this study $v = 10$ is used. This is because too small number of models in ensemble will yield too little diversity hindering the models to correct each others errors, but, on the other hand, using too many models will yield no further improvement (Breiman, 1996; Opitz & Maclin, 1999; Kotsiantis & Pintelas, 2004; Parmanto et al., 1996). Moreover, too large number of CV folds can yield unreliable validation MSE estimates for the selection of the individual final best models, as then the individual validation sets may be too small.

![Diagram of the EF-ABFC model building process](image)

Fig. 5. An outline of the EF-ABFC modelling process when $v = 3$: (a) search for the best model according to AICC using F-ABFC; (b) select the one final best model according to MSE in validation data set; (c) re-fit the model (recalculate its parameters) using the whole training data; (d) combine the models

In recent literature, there is ever growing confidence that model ensembles often perform better than individual models and consistently reduce prediction error (Breiman, 1996; Opitz & Maclin, 1999; Kotsiantis & Pintelas, 2004; Jekabsons, 2008). However, model ensembles are not always the best solutions (Kotsiantis & Pintelas, 2004): if there is too little...
data, the gains achieved via an ensemble may not compensate for the decrease in accuracy of individual models, each of which now sees an even smaller training set. On the other end, if the data set is sufficiently large, even a single flexible model can be quite adequate. Using large data sets also substantially decreases potential selection bias, so superiority of EF-ABFC over F-ABFC in such situations is expected to diminish.

The most significant disadvantage of the EF-ABFC compared to F-ABFC is that it requires larger computational resources. However, the fact, that before the model combining the \( \nu \) models are built completely separately, allows for an easy parallelization of the process dividing the execution time by \( \nu \). In this study however the parallelization is not done.

The paper (Jekabsons, 2008) empirically demonstrated the computational and predictive performance advantages of EF-ABFC comparing to subset selection and a number of other popular regression modelling methods. EF-ABFC advantages in real-world practical applications are demonstrated in (Kalnins et al., 2008b; Kalnins et al., 2009a) where it is applied for modelling bending and buckling behaviour of different composite material structures.

### 4.7 Remarks

This section covers various aspects (extensions, limitations, etc.) of the ABFC not discussed in the previous sections.

#### 4.7.1 Incorporating domain knowledge

The ABFC methods attempt to model arbitrary dependencies in data with little or no knowledge of the system under study. In problems of moderate and large dimensionality the user usually is not required to tune any hyperparameters. However, if there is sufficient additional domain knowledge outside the specific data at hand, it may be appropriate to place some constraints on the final model. If the knowledge is fairly accurate, such constraints can improve the accuracy while saving computational resources.

For example, the constraints might be one or more of the following: 1) limiting the maximal degree of all the basis functions (similarly as in the subset selection), i.e. \( 0 \leq \sum_{j=1}^{d} r_{ij} \leq p \) for all \( i \); 2) limiting the maximal value of exponents for each particular input variable in all the basis functions, i.e. \( 0 \leq r_{ij} \leq p_{j} \) for all \( i \), where \( p_{j} \) is maximal exponent of the \( j \)th variable; 3) restricting contributions of specific input variables that are not likely to interact with others so that those variables can enter the model in basis functions only solely – with exponents of all other variables fixed to zero. These constraints, as well as far more sophisticated ones, can be easily incorporated in the ABFC. However, note that in all the experiments described in this chapter no constraints are used.

#### 4.7.2 Robustness

The ABFC methods described in this study estimate model parameters via minimization of the squared-error loss, i.e. using OLS. However, while the squared-error loss is the most commonly used, it is known that it loses its robustness against grossly outlying samples as well as in very sparse high-dimensional data sets (Cherkassky & Ma, 2002).

One solution of this problem is to use a more robust loss function. The squared-error loss in ABFC is not fundamental. Any other loss function can be used to estimate the parameters...
and to evaluate the models by simply replacing the routine “PerformOLSandCalculateAICC” of the search procedure (Figure 3) with a more robust one. Note that while this would make the methods more robust, the computational advantage of OLS would be lost. In any case, gross outliers (in output variable as well as input variables) that can be detected through a preliminary data analysis should be considered for removal before applying ABFC.

4.7.3 Other types of basis functions
The ABFC methods described in this study can generate regression models with basis functions of only nonnegative integer exponents. However, in principle the exponents can also be allowed to take negative or even fractional values. Appropriate adaptation of the state-transition operators can enable generating such models. Keeping the same initial model as before, the search now could go in direction of both positive and negative exponents.

4.7.4 Integrating ABFC into other modelling methods
The result of running an ABFC procedure is a simple polynomial regression model. Such models are also utilized as “sub-models” in a number of other regression modelling methods. For example, the ABFC methods can be used in Polynomial Neural Networks (usually induced by Group Method of Data Handling) (Nikolaev & Iba, 2006) for adaptation of each individual neuron’s functional form and degree. The methods also can serve for generation of local regression models in Locally-Weighted Regression (also called Moving Least Squares) (Cleveland & Devlin, 1988; Kalnins et al., 2008b; Kalnins et al., 2005) adaptively generating a model each time a query is received. ABFC can also induce piecewise polynomial models for appropriately partitioned data sets.

The polynomial basis functions can also be viewed as nonlinear transformations (or features) of the original input variables. In this manner the ABFC methods can also be viewed as methods for automatic adaptive feature construction. For example, the constructed features can further serve as inputs for Support Vector Machines (Vapnik, 1995; Smola & Scholkopf, 2004) similarly to the features constructed using genetic algorithm in (Rithoff et al., 2002).

All these applications of ABFC can make the original methods more flexible and therefore, if treated appropriately, produce models of higher predictive performance.

4.7.5 Using ABFC for solving classification problems
The ABFC methods can also be used for solving binary classification problems where the output variable y can take value of only either 0 or 1. This can be done, for example, by constructing basis functions for logistic regression (also called maximum entropy classifier) models. Logistic regression (Hastie et al., 2003; Witten & Frank, 2005) represents log odds of y being equal to 1 as a linear model:

$$ \ln \left( \frac{P}{1-P} \right) = F(x) = \sum_{i=1}^{k} a_i f_i(x), $$

(16)
where $P$ is the predicted probability of $y$ being equal to 1. It is equivalent to the following representation of $P$:

$$P = \frac{1}{1 + \exp(-F(x))}. \quad (17)$$

The parameters $a$ of the model are usually estimated by minimizing the deviance:

$$-2 \sum_{j=1}^{n} (y_j \ln F(x_j) + (1 - y_j) \ln(1 - F(x_j))) \rightarrow \min. \quad (18)$$

Since there is no closed form solution to this minimization, the standard approach to solving it is to use iterative algorithms such as Iteratively Re-weighted Least-Squares (Hastie et al., 2003; Witten & Frank, 2005). Note that, in order to evaluate a model using AICC, the first term of (13) is replaced by the deviance.

F-ABFC and EF-ABFC for classification problems are implemented in the VariClass software tool freely available for non-commercial research and educational purposes at http://www.cs.rtu.lv/jekabsons/.

5. Related work

There exist also other polynomial regression modelling methods which use wide, potentially infinite, dictionaries of basis functions. In (Sutton & Matheus, 1991) an algorithm is proposed which starts model building with a first-degree model, with all the input variables already included in the model, and iteratively creates a user-predefined number of products of the already included basis functions thereby creating new basis functions. In (Orosz & Anderson, 1994) a modification of the algorithm is proposed where the initial model has none of the input variables included, however there was no empirical success and it was concluded that in practical applications the algorithms have three major disadvantages: inability to construct all the necessary basis functions, inability to discard unnecessary basis functions, and high sensitivity to noise and to number of samples in data.

More recently a different method was developed which can be seen also as a special case of the ABFC approach – Constrained Induction of Polynomial Equations for Regression, CIPER (Todorovski et al., 2004). CIPER was initially developed in the context of differential equation discovery, inductive databases, and constraint-based data mining. CIPER uses two state-transition operators and a Beam Search strategy. The first state-transition operator adds a new linear basis function while the second increases a single exponent of a single basis function. In (Jekabsons & Lavendels, 2008a), CIPER was empirically compared to F-ABFC and it was concluded that CIPER suffers from the nesting effect (Pudil et al., 1994) and has a tendency of getting stuck in local minima too early in the search. This is because CIPER is not able to preserve the structure of any of included basis functions (its second operator increases an exponent in an existing basis function but does not take into consideration the possibility that both versions of the basis function may be required) as well as because it is not able to simplify a model – decrease unnecessarily high exponents or discard unnecessary basis functions. In F-ABFC these issues are solved using Operator2 and the simplification operators.
Some similar ideas of constructing new features as combinations of original input variables are applied also in different other approaches. For example, in (Ritthoff et al., 2002) a feature construction method is proposed in which a genetic algorithm constructs linear and nonlinear combinations of original input variables further used as inputs for Support Vector Machines. In (Bloedorn & Michalski, 1998), on the other hand, the feature construction idea is used for data-driven expansion of the input space for induction of decision rules and decision trees.

6. Experiments

This section presents the results of comparisons of the proposed ABFC methods to the methods of subset selection and to a number of other well known state-of-the-art regression modelling methods using a series of synthetic and real-world regression data sets. The goal is to gain some understanding of the properties of F-ABFC and EF-ABFC and to evaluate their performance in both accuracy and speed. All the experiments were performed on a Pentium IV 2.4GHz machine with 1.5GB RAM.

In all the experiments, predictive performance of a model is measured either using a completely independent test data set or using Cross-Validation. In any case the performance of a model is measured in terms of Relative Root Mean Squared Error:

$$RRMSE = 100\% \times \frac{\sqrt{\sum_{j=1}^{n} (y_j - F(x_j))^2}}{\sqrt{\sum_{j=1}^{n} (y_j - \bar{y})^2}} ,$$

where \( n \) is the number of samples in the test data set, \( F(x) \) is the predicted value corresponding to the value of \( y_j \) and \( \bar{y} \) is the mean of all the \( y \) values in the test set. While RMSE (Root Mean Square Error) represents model’s deviation from the data, the SD (Standard Deviation) captures how irregular the problem is. The lower the value of RRMSE, the more accurate is the model. The final RRMSE values stated are the values averaged over all evaluations.

All the employed regression modelling methods, except Regression Trees, Model Trees, Support Vector Machines, and Multi-Layer Perceptrons, are implemented in VariReg software tool version 0.9.21 freely available for non-commercial research and educational purposes at http://www.cs.rtu.lv/jekabsons/.

6.1 Synthetic data sets

To compare the performance of the proposed ABFC methods against subset selection (as well as against “full” polynomials with no subset selection) in different conditions of signal-to-noise ratio (SNR) and training data size, here two test functions are used – Synth1 (4 input variables) and Synth2 (10 input variables):

$$y_{\text{Synth1}} = \exp(2x_1 \sin(\pi x_4)) + \sin(x_2 x_3) ,$$

$$y_{\text{Synth2}} = (x_1^2 + (x_2 + x_3) (x_4 + x_5) (1 + x_6 x_7) + 0x_8 + 0x_9 + 0x_{10} .$$

For Synth1 the values of \( x \) are uniformly distributed in the interval [-0.25, 0.25]. For Synth2 they are uniformly distributed in the interval [0, 1]. For each test function three training set...
sizes (25 samples, 50 samples, and 100 samples) and three signal-to-noise ratios (no noise, SNR = 4, and SNR = 2) are used – a total of nine cases for each function. For each case a series of 20 training data sets are generated (randomly sampled in the domain of \( x \)) so that in each case for each regression modelling method the model building task is performed 20 times. For each test function a single test data set is generated containing 5000 samples randomly sampled in the domain of \( x \). The test data sets do not contain noise.

The heuristic search algorithms used for the subset selection are the SFS and the SFFS (the same algorithm adaptation of which is used in the ABFC methods). The algorithms are used together with the AICC criterion (also the same which is used in the ABFC methods). Note that the “recursion depth” hyperparameter of F-ABFC is set equal to 2 for Synth1 and equal to 1 (no recursion) for Synth2.

As for the full polynomials (FP) and the subset selection methods the desirable degree \( p \) is not known beforehand, the modelling results of these methods are stated in two forms: 1) average performance of models of a fixed \( p \); 2) average performance when a range of values for \( p \) are tried and the model of the lowest RRMSE value is picked. However, note that this second type of procedure for FP/SFS/SFFS is rather optimistic (in the sense of both predictive performance and speed) as for correct and fair evaluations there would be an additional validation data set or a Cross-Validation loop required.

| Method | RRMSE (sn) | Time (s) | RRMSE (sn) | Time (s) | RRMSE (sn) | Time (s) |
|--------|------------|----------|------------|----------|------------|----------|
| FP, \( p \in [1, 4] \) | 9.29 (1.88) | - | 6.74 (0.55) | - | 0.78 (0.21) | - |
| SFS, \( p = 2 \) | 7.17 (1.03) | < 0.1 | 6.38 (0.58) | < 0.1 | 5.63 (0.28) | < 0.1 |
| SFS, \( p = 6 \) | 0.77 (2.24) | 0.3 | 0.06 (0.02) | 1.4 | 0.04 (1e-2) | 8.0 |
| SFS, \( p = 10 \) | 3.19 (7.41) | 0.8 | 0.03 (0.04) | 16.1 | 2e-3 (7e-3) | 104.3 |
| SFS, \( p \in [1, 10] \) | 0.77 (2.24) | 4.5 | 0.03 (0.03) | 34.1 | 2e-3 (7e-3) | 226.1 |
| SFFS, \( p \in [1, 10] \) | 0.77 (2.24) | 4.5 | 0.03 (0.03) | 34.1 | 2e-4 (2e-4) | 236.9 |
| F-ABFC | 0.11 (0.15) | 0.1 | 0.01 (0.02) | 2.0 | 3e-7 (5e-7) | 43.8 |
| EF-ABFC | 0.27 (0.31) | 0.8 | 0.02 (0.02) | 11.5 | 1e-4 (4e-4) | 250.6 |

| Method | RRMSE (sn) | Time (s) | RRMSE (sn) | Time (s) | RRMSE (sn) | Time (s) |
|--------|------------|----------|------------|----------|------------|----------|
| FP, \( p \in [1, 4] \) | 42.71 (16.31) | - | 18.36 (2.53) | - | 12.12 (1.59) | - |
| SFS, \( p = 2 \) | 24.24 (10.87) | < 0.1 | 15.62 (2.99) | < 0.1 | 10.64 (2.38) | < 0.1 |
| SFS, \( p = 6 \) | 59.37 (28.09) | 0.1 | 41.37 (11.76) | 0.3 | 25.60 (9.49) | 0.8 |
| SFS, \( p = 10 \) | 112.02 (149.28) | 0.7 | 74.10 (40.38) | 3.4 | 39.23 (10.92) | 7.9 |
| SFS, \( p \in [1, 10] \) | 24.24 (10.87) | 1.7 | 15.62 (2.99) | 8.7 | 10.64 (2.38) | 19.5 |
| SFFS, \( p \in [1, 10] \) | 24.24 (10.87) | 1.8 | 15.62 (2.99) | 7.6 | 10.64 (2.38) | 21.0 |
| F-ABFC | 39.05 (17.97) | < 0.1 | 33.13 (15.64) | 0.1 | 22.64 (10.73) | 0.3 |
| EF-ABFC | 20.24 (6.76) | 0.3 | 13.65 (3.82) | 0.8 | 9.08 (3.16) | 2.1 |

| Method | RRMSE (sn) | Time (s) | RRMSE (sn) | Time (s) | RRMSE (sn) | Time (s) |
|--------|------------|----------|------------|----------|------------|----------|
| FP, \( p \in [1, 4] \) | 79.40 (37.25) | - | 35.97 (8.35) | - | 21.79 (4.29) | - |
| SFS, \( p = 2 \) | 36.35 (12.40) | < 0.1 | 26.51 (9.87) | < 0.1 | 18.55 (4.35) | < 0.1 |
| SFS, \( p = 6 \) | 88.32 (32.57) | 0.1 | 70.34 (24.81) | 0.3 | 47.11 (16.95) | 1.0 |
| SFS, \( p = 10 \) | 209.98 (213.00) | 0.8 | 99.26 (40.07) | 2.8 | 78.04 (31.78) | 8.1 |
| SFS, \( p \in [1, 10] \) | 36.35 (12.40) | 1.7 | 26.51 (9.87) | 5.9 | 18.55 (4.35) | 18.7 |
| SFFS, \( p \in [1, 10] \) | 36.35 (12.40) | 1.7 | 26.64 (10.08) | 6.3 | 18.55 (4.35) | 19.5 |
| F-ABFC | 58.43 (19.72) | < 0.1 | 72.44 (62.43) | < 0.1 | 39.93 (19.91) | 0.2 |
| EF-ABFC | 35.23 (11.04) | 0.3 | 24.94 (6.18) | 0.7 | 17.67 (4.45) | 1.8 |

Table 1. The results of the performed experiments for function Synth1
The results of the performed experiments are summarized in Table 1 and Table 2 in terms of mean RRMSE value, with its standard deviation reported in parenthesis, and elapsed time. Note that, due to the space constraints, for fixed degrees only the results of $p \in \{2, 6, 10\}$ (for Synth1), $p \in \{2, 5\}$ (for Synth2) are given. Detailed results are available at http://www.cs.rtu.lv/~jekabsons/.

Figure 6 and Figure 7 visualizes the performance changes of the methods for different training set sizes and SNRs.

Table 2. The results of the performed experiments for function Synth2

| No noise | $n = 25$ | $n = 50$ | $n = 100$ |
|----------|---------|---------|---------|
| **Method** | RRMSE | Time (s) | RRMSE | Time (s) | RRMSE | Time (s) |
| FP, $p \in [1, 2]$ | 45.40 (4.86) | - | 38.73 (2.06) | - | 13.07 (1.37) | - |
| SFS, $p = 2$ | - | 38.17 (13.99) | <0.1 | - | 19.13 (3.20) | 0.2 |
| SFS, $p = 5$ | 80.25 (23.80) | 12.8 | 29.13 (12.73) | 53.6 | - | 6.3 |
| SFS, $p \in [1, 5]$ | 38.17 (13.99) | 14.7 | 19.13 (3.20) | 57.1 | - | 6.3 |
| SFFS, $p \in [1, 5]$ | 37.40 (12.36) | 15.9 | 20.20 (4.03) | 82.5 | - | 6.3 |
| F-ABFC | 52.86 (11.46) | <0.1 | 13.14 (6.96) | 0.7 | 1.59 (1.58) | 16.5 |
| EF-ABFC | 56.39 (16.40) | 0.3 | 12.92 (3.08) | 4.2 | 0.95 (0.46) | 98.7 |

| SNR = 4 |
|----------|
| **Method** | RRMSE | Time (s) | RRMSE | Time (s) | RRMSE | Time (s) |
| FP, $p \in [1, 2]$ | 51.78 (8.53) | - | 41.31 (3.25) | - | 37.38 (1.51) | - |
| SFS, $p = 2$ | - | 58.85 (15.18) | <0.1 | - | 35.44 (7.05) | 0.1 |
| SFS, $p = 5$ | 180.68 (68.02) | 10.7 | 82.78 (23.99) | 66.0 | - | 298.8 |
| SFS, $p \in [1, 5]$ | 58.85 (15.18) | 13.1 | 35.44 (7.05) | 77.8 | - | 298.8 |
| SFFS, $p \in [1, 5]$ | 61.01 (17.19) | 14.2 | 35.88 (8.69) | 78.2 | - | 298.8 |
| F-ABFC | 79.07 (35.39) | <0.1 | 45.59 (9.28) | 0.1 | 28.89 (7.25) | 0.5 |
| EF-ABFC | 62.79 (11.47) | 0.3 | 35.77 (3.73) | 1.3 | 20.37 (3.51) | 6.3 |

| SNR = 2 |
|----------|
| **Method** | RRMSE | Time (s) | RRMSE | Time (s) | RRMSE | Time (s) |
| FP, $p \in [1, 2]$ | 61.23 (9.17) | - | 46.61 (4.73) | - | 40.81 (3.12) | - |
| SFS, $p = 2$ | - | 73.81 (17.63) | <0.1 | - | 51.69 (8.15) | 0.1 |
| SFS, $p = 5$ | 180.68 (68.02) | 11.0 | 135.99 (37.13) | 47.6 | - | 208.9 |
| SFS, $p \in [1, 5]$ | 73.81 (17.63) | 13.3 | 47.92 (6.96) | 57.4 | - | 253.0 |
| SFFS, $p \in [1, 5]$ | 76.43 (11.56) | 14.3 | 50.41 (9.44) | 64.3 | - | 369.3 |
| F-ABFC | 82.42 (18.79) | <0.1 | 68.08 (15.40) | 0.1 | 49.61 (13.73) | 0.3 |
| EF-ABFC | 70.84 (9.52) | 0.2 | 51.11 (8.79) | 0.8 | 34.54 (5.93) | 3.9 |

Table 2. The results of the performed experiments for function Synth2
The results in Table 1 indicate that for noise-free data the F-ABFC outperforms its much slower ensembled extension EF-ABFC while for noisy data it is vice versa. When the data contains noise, the F-ABFC here can be outperformed even by full polynomials which mostly give some of the worst performances. This suggests that for noisy data it is important to curb the flexibility of F-ABFC – to use the EF-ABFC even when the data is sparse.

![Graph showing performance of methods for different training set sizes and SNRs](image)

**Fig. 7.** Performance of the methods for function Synth2 for the different training set sizes and SNRs: (a) no noise; (b) SNR = 4 (solid lines) and SNR = 2 (dashed lines)

The results in Table 2 partially confirm those in Table 1 except that this time the EF-ABFC is always more accurate than F-ABFC which may be caused by the three irrelevant input variables (pure noise) in the data on which the Synth2 does not depend. Additionally, as now in the case of \( n = 25 \) the data are very sparse, for this case the ABFC methods are just too flexible – they largely overfit the data even when there is no additional noise.

Overall, the results for both Synth1 and Synth2 indicate the computational advantage of the ABFC methods in situations when the required regression model is more complex (of higher degree). And this advantage grows with the dimensionality of the problem.

For noisy data the best choice of \( p \) for SFS/SFFS almost always was 2. Then the speed of a single SFS/SFFS search can be outperformed only by F-ABFC. However, as the best \( p \) value is actually unknown and a number of values must be tried, F-ABFC as well as EF-ABFC is still faster than the subset selection.

Finally, it must also be noted that the overall results show evidence that for subset selection the choice of the search algorithm (either SFS or SFFS) was of no great importance. Therefore further in this study only the SFS algorithm for subset selection is considered.

### 6.2 Real-world machine learning data sets

The real-world machine learning regression data sets used are: autoMPG (7 input variables, 392 samples), AutoPrice (15 input variables, 159 samples), Bodyfat (14 input variables, 252 samples), Fishcatch (7 input variables, 158 samples), Housing (13 input variables, 506 samples), HousingNOX (13 input variables, 506 samples), MachineCPU (6 input variables, 209 samples), Pyrimidines (27 input variables, 73 samples), Servo (4 input variables, 167 samples), and Stock (9 input variables, 950 samples). The data sets are from UCI Machine Learning Repository (http://www.ics.uci.edu/~mlearn/MLRepository.html), Luis Torgo’s data sets repository (http://www.liaad.up.pt/~ltorgo/Regression/DataSets.html), and
Weka collection of data sets (http://www.cs.waikato.ac.nz/ml/weka/). They are chosen because of the relatively low number of samples, which is common in real-world practical applications, as well as because of mostly continuous input variables and no missing values. Here the performances of the different regression modelling methods are evaluated using 10-fold Cross-Validation. Note that prior to dividing the data into Cross-Validation folds, the order of the samples was randomized.

The goal of the performed experiments is to compare the proposed ABFC methods to the methods of subset selection and to other well known state-of-the-art regression modelling methods using a set of real-world regression data sets. The compared methods are the following: FP, SFS, F-ABFC, EF-ABFC, Multivariate Adaptive Regression Splines (MARS) (Friedman, 1993), M5’ Regression Trees (RT) (Witten & Frank, 2005), M5’ Model Trees (MT) (Witten & Frank, 2005), Support Vector Machines (SVM) (Vapnik, 1995; Smola & Scholkopf, 2004), and Multi-Layer Perceptrons (Witten & Frank, 2005). Note that for none of the methods any of the hyperparameters were manually tuned. MARS was that of piecewise-cubic type essentially without special limitation of the number of basis functions (i.e. the limit was 500) and with the smoothing parameter (the number of degrees of freedom associated with one basis function) either fixed to the default value of 3 or found using an additional 10-fold Cross-Validation from the range \([1, 5]\) with step size 0.5. SVM used Radial Basis Function kernel and improved Sequential Minimal Optimization algorithm (Shevade et al., 1999) for which the complexity parameter and the gamma parameter were found using grid search and Cross-Validation from the range \([10^{-1}, 10^{0}, 10^{1}, 10^{2}]\) for the complexity parameter and \([10^{-2}, 10^{-1}, 10^{0}, 10^{1}]\) for the gamma parameter. MLP had one hidden layer with the “best” number of neurons determined by 10-fold Cross-Validation from the range \([10, 20, 30, 40]\) and the weights were optimized using backpropagation. As implementations of RT, MT, SVM, and MLP the Weka software (Witten & Frank, 2005) was employed with its default parameters. Also note that for the ABFC methods the recursion of the state-transition operators was never used.

The results of the performed experiments are summarized in Table 3 in terms of mean RRMSE value, with the standard deviation reported in parenthesis, and elapsed time. Here the modelling results of SFS are stated in the same two forms as in Section 6.1 except that for the different data sets (different in size, in number of input variables, and in required model complexity) the values of \(p\) are tried in different intervals (named “\(p = \text{automatic}\)”) – the search for the best \(p\) is started with the first degree and \(p\) is increased as long as the RRMSE value improves. The results of FP are not stated, as due to matrix singularity in OLS for Pyrimidines data set the parameter values of FP models could not be calculated. Also note that, due to the space constraints, only the results averaged over all the data sets are given. Detailed results are available at http://www.cs.rtu.lv/jekabsons/.

From the results of the experiments it is concluded that in terms of predictive performance, the EF-ABFC outperformed all the other regression modelling methods involving polynomials as well as showed high competitiveness against the other “non-polynomial” methods. In terms of computational cost, both ABFC methods outperformed subset selection but were inferior to some of the “non-polynomial” methods, especially RT, MT, and MARS without CV.
Table 3. The average results of the performed experiments for the ten machine learning data sets

| Method     | RRMSE    | Time (s) |
|------------|----------|----------|
| SFS, $p = 1$ | 49.64 (12.05) | < 0.1 |
| SFS, $p = 2$ | 40.89 (15.11) | 4.8 |
| SFS, $p = 3$ | 37.83 (15.70) | 227.4 |
| SFS, $p = 4$ | 47.10 (29.17) | 2486.8 |
| SFS, $p =$ automatic | 34.83 (10.20) | 2207.5 |
| F-ABFC     | 39.61 (16.93) | 108.7 |
| EF-ABFC    | 31.24 (10.86) | 607.8 |
| RT         | 50.76 (9.85)  | 0.3  |
| MT         | 34.79 (12.35) | 0.4  |
| MARS       | 40.81 (17.29) | 3.2  |
| MARS + CV  | 39.87 (15.57) | 265.8 |
| SVM        | 31.87 (10.73) | 360.5 |
| MLP        | 41.50 (18.04) | 345.2 |

For the different data sets, the best found degree $p$ for SFS with “$p =$ automatic” varied in range [1, 6] (with the average value of 3.0), meaning that the maximal checked value of $p$ was 7 (though for only one of the data sets). However, the average degree of models constructed by F-ABFC and EF-ABFC was 6.4 and 7.2 correspondingly. If on average for SFS such large values of $p$ would be tried (instead of only 3.0), the SFS would take considerably more time (orders of magnitude) to complete.

### 6.3 Real-world metamodelling data sets

In many different industrial applications, to cut down the computational cost of complex, high fidelity scientific and engineering simulations, regression models (in the context also referred to as metamodels or surrogate models) are constructed that mimic the behaviour of the simulation models as closely as possible while being computationally much cheaper to employ (Myers & Montgomery, 2002; Chen et al., 2006; Martin & Simpson, 2005; Kalnins et al., 2008b; Kalnins et al., 2008a; Kalnins et al., 2009a; Kalnins et al., 2009b). The process of design optimization involving metamodelling usually comprises three major steps which may be interleaved iteratively: 1) selection of samples (known as design of experiments); 2) construction of metamodel and estimation of its predictive performance; 3) employment of the metamodel in design optimization (i.e., finding the best values for input variables.
with which the studied system achieves the optimum response), design space exploration, what-if analysis, sensitivity analysis, and other routine tasks.

The metamodelling problem addressed here is modelling of the behaviour of “I-core” all-metal laser-welded sandwich panels under bending load for further design optimization and analysis in application as deck panels in a modularised watercraft concept (Kalnins et al., 2008a). The problem has six input variables and four output variables. The data are generated using finite element simulations and contains 500 samples distributed in the input space using sequential experimental design (Auzins, 2004).

Originally, metamodelling was associated with low-degree (usually quadratic) polynomial models. They have been well accepted in engineering practice, as they require only little data and are computationally very efficient. However, it is understood that they are loosing efficiency when highly nonlinear behaviour should be approximated.

In this section the compared regression modelling methods are the same as in the Section 6.2 with an addition of three methods which are rather popular in metamodelling literature: Locally-Weighted Polynomials (LWP) (Cleveland & Devlin, 1988; Kalnins et al., 2008b; Kalnins et al., 2005), Radial Basis Functions (RBF) (Gutmann, 2001), and Kriging (Martin & Simpson, 2005; Lophaven et al., 2002). Note again that for none of the methods any of the hyperparameters were manually tuned. LWP used the Gaussian weight function with the value of the bandwidth parameter found by Leave-One-Out Cross-Validation. Note that the LWP has a similar issue of degree \( p \) selection as FP and SFS, so here a number of different degrees are tried in the interval \([1, 4]\). RBF used the multi-quadric basis functions with the shape parameter fixed to 1. Kriging used first-degree polynomial as a trend function and employed the Gaussian correlation function. Note that the used source code for the Kriging technique was developed by (Lophaven et al., 2002). Also note that for the ABFC methods in the performed experiments the recursion of the state-transition operators was never used.

The results of the performed experiments are summarized in Table 4 in terms of mean RRMSE value, with its standard deviation reported in parenthesis, and elapsed time. Here the performances of the different regression modelling methods are evaluated using 5-fold Cross-Validation. The modelling results of FP and SFS are stated in the same two forms as in Section 6.1. Note that, due to the space constraints, only the results averaged over all the data sets are given. Detailed results (as well as the utilized data sets) are available at http://www.cs.rtu.lv/jekabsons/.

The results in Table 4 indicate that with the four metamodelling data sets (all of which are essentially noise-free) the ensembling of F-ABFC models was not necessary – the accuracy advantage of EF-ABFC is negligible while it is computationally about ten times slower than simple F-ABFC. However, both F-ABFC and EF-ABFC outperformed subset selection in terms of predictive performance as well as in terms of speed. In respect to the other methods the ABFC approach once again is highly competitive, especially the faster F-ABFC method.

With the metamodelling data sets, on average the best degree \( p \) for SFS was 6.3 while the average degree of models constructed by F-ABFC and EF-ABFC was 9.2 and 7.9 correspondingly. Similarly to the conclusions of the previous section, trying these larger values of \( p \) for SFS would take orders of magnitude more time to complete.
Note that in practice it turns out that the user all too often does model building in a “one-shot” manner, without consideration of different settings for a modelling method. With FP and SFS (as well as LWP) it could mean that almost any of the results stated in Table 4 (as well as in the other tables from previous sections) may be accepted as the final. Iterative and adaptive methods like those of ABFC, on the other hand, have the potential of relatively rapidly producing accurate models without the configuration burden.

7. Conclusion

This chapter introduced Adaptive Basis Function Construction – an adaptive sparse polynomial regression model building approach which can also be viewed as an alternative to the classical subset selection approach. In contrast to subset selection, the ABFC approach does not require putting restrictions on model’s degree, enables model building in polynomial time, and does not require repetition of the model building process. The basis functions required for the model are automatically adaptively constructed using heuristic search specifically for data at hand without using a restricted fixed finite user-predefined dictionary. The dictionary in the ABFC is infinite and polynomials of arbitrary complexity can be constructed.

In most of the performed empirical experiments, the ABFC methods outperformed subset selection in terms of predictive performance as well as in terms of the amount of required

| Method       | RRMSE    | Time (s) |
|--------------|----------|----------|
| FP, \( p = 1 \) | 49.85 (4.82) | -        |
| FP, \( p = 2 \) | 23.81 (3.10) | -        |
| FP, \( p = 3 \) | 12.81 (1.77) | -        |
| FP, \( p = 4 \) | 9.88 (1.46) | -        |
| FP, \( p \in [1, 4] \) | 9.17 (1.28) | -        |
| SFS, \( p = 1 \) | 49.75 (4.68) | < 0.1    |
| SFS, \( p = 2 \) | 23.42 (3.15) | 0.2      |
| SFS, \( p = 3 \) | 11.74 (1.84) | 4.2      |
| SFS, \( p = 4 \) | 7.31 (1.35) | 41.1     |
| SFS, \( p = 5 \) | 5.62 (1.14) | 220.3    |
| SFS, \( p = 6 \) | 5.03 (0.78) | 959.1    |
| SFS, \( p = 7 \) | 5.05 (1.12) | 1828.4   |
| SFS, \( p \in [1, 7] \) | 4.92 (0.75) | 3053.4   |
| F-ABFC       | 4.28 (0.55) | 71.9     |
| EF-ABFC      | 4.19 (0.55) | 715.4    |
| RT           | 60.18 (7.87) | 1.0      |
| MT           | 22.27 (4.97) | 4.7      |
| MARS         | 5.87 (0.96) | 0.9      |
| MARS + CV    | 5.31 (0.84) | 77.5     |
| SVM          | 13.14 (2.57) | 414.7    |
| MLP          | 8.47 (1.03) | 331.3    |
| LWP, \( p = 1 \) | 40.22 (4.12) | 2.8      |
| LWP, \( p = 2 \) | 20.23 (2.79) | 26.2     |
| LWP, \( p = 3 \) | 11.66 (1.68) | 210.6    |
| LWP, \( p = 4 \) | 9.76 (1.42) | 1576.7   |
| RBF          | 14.48 (3.42) | 1.9      |
| Kriging      | 7.40 (1.21) | 16.3     |

Table 4. The average results of the performed experiments for the four metamodelling data sets.
computational resources. Moreover, in respect to the other well-known state-of-the-art regression methods, the ABFC approach is also highly competitive. Additionally, the ABFC methods have advantages also in their simple application – the underlying algorithms have very small number of hyperparameters for the user to tune and result in simple explicit equations employable without specialized software.

Comparing the two specific methods F-ABFC and EF-ABFC, it is concluded that EF-ABFC has predictive performance advantage over F-ABFC when the data contains noise, be it in terms of signal-to-noise ratio or in terms of irrelevant input variables. On the other hand, F-ABFC is much faster than EF-ABFC and can produce more accurate models when the data is noise-free. Nevertheless, both methods may require the “recursion depth” hyperparameter to be set to a value higher than 1 when the data is of low dimensionality (e.g., \( d \leq 4 \)) and/or the existing structure in the data requires a very complex model.

As future work, some of the ideas described in Section 4.7 could be pursued.

Software (including open source) implementing the ABFC methods, as well as most of the other regression methods employed in this chapter, can be downloaded at the author’s webpage: http://www.cs.rtu.lv/jekabsons/.

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