Learning and Predicting the Performance of Configurable Software Systems

Seminar - Advanced Software Engineering: Non-Functional Aspects in Software Engineering

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Abstract. Today's programs are mostly highly configurable and customizable. Popular applications like Apache or MySQL can have hundreds of configurable parameters. Whilst providing flexibility to a customer, this also brings some problems: having so many options can heavily influence the performance of a software system. And the more options there are, the harder it gets to predict the behavior of a software system. This paper will show that a brute force approach to this problem does not work as a general solution. Further it is going to explain and compare four different prediction methods that were developed by N. Siegmund et al. The comparison of those methods shows, that there is no 'best' approach to this problem, but rather a broad selection of well working methods exists.

1 Introduction

As modern programs grow larger and more powerful they also provide many configuration opportunities to their customers. In some cases the number of parameters can be even greater than 500. Examples for this can be found in Fig. 1. With this large amounts of configuration options stakeholders or customers can be satisfied easier since they can tailor a program to their specific requirements. But with this large amount of options comes a bigger problem: "Unpredictability". Looking at an example of Apache Storm (Fig. 2a) shows that the performance of two configurations of a program can differ significantly. Fig. 2b shows that solely changing a single parameter can increase the response time of Apache Storm by up to 100%. Without using prediction methods such results are only visible after executing and measuring multiple, if not all, configurations of a system. Or in other words: when looking only at a single configuration, one cannot conclude whether that configuration is any good for the current requirements in.

This is where prediction comes into play. By learning about the performance of some configurations of the system it tries to generate a function that can give an expected performance for a not measured configuration. This can be used to solve the just mentioned problem of finding a near optimal solution.

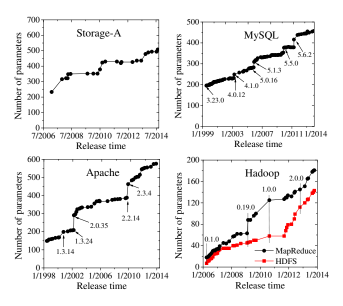


Fig. 1: Number of parameters of different popular programs [15].

Furthermore, performance prediction can be used to find default configurations. These should be configurations that fulfils most requirements to an acceptable level. The most straightforward approach to this problem would be a brute-force solution. In this case that would mean measuring each and every single valid configuration. As we will see later in Section 3 this approach is in general not feasible since the amount of valid configurations scales exponentially with the number of parameters. For that reason other approaches had to be found and especially the efficient sampling of a configuration space turned out to be a problem [10].

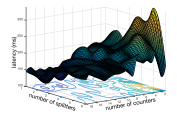
This paper will focus on showing different approaches and strategies to predicting the performance of a configurable software system. It will mainly discuss approaches developed by Norbert Siegmund et al. [2, 6, 10, 12]. They will be explained and compared. More specifically, this paper will have a look at four different approaches besides *brute-force*.

The first discussed technique is Automated Feature Interaction Detection (AFID) [12]. The goal of this approach is to assign a performance influence value to each feature and feature interaction. This is done by observing and measuring the behavior of certain configurations. The other 4 approaches make use of a CART Tree as their learning choice but differ in the way they choose their sample.

Variability Aware Performance Prediction (VAPP) [2] uses random sampling to pick which configurations to compile and measure.



(a) Configurations of Apache Storm sorted by measured throughput.



Only by tweaking 2 options out of 200 in Apache Storm - observed ~100% change in

(b) Possible influences of only two options on the latency of Apache Storm.

Fig. 2: Measurements done for Apache Strom which shows that configurations can have a significant influence on the performance of a software system.

WHAT [6], tires a more mathematical way to find groups of similar configurations without actually measuring them. For this distance based clustering/sampling is used.

The last two sampling approaches are proposed in the same paper by Sarkar et al. [10].

Progressive and Projective Sampling are quite similar, since they both take advantage of the fact, that the general formula behind a learning curve is known. With this knowledge they generate a part of the actual curve and fit a function to it. Based on this function an optimal size for the actual sample set can be calculated. Both methods also take the cost of measurements (resources and accuracy) into consideration.

All these approaches can reach accuracies of over 94% on average in the conducted tests of their corresponding papers. This makes them good enough to be relevant for the topic of this paper. Further their results can be compared straightforwardly since they are all tested on the same set of 6 software systems: Berkeley DB C, Berkeley DB Java, Apache, SQLite, LLVM, x264.

2 Definitions

Before the actual approaches are discussed it is important to pinpoint the definitions of terms which are used in this paper.

This paper often uses the terms "parameters", (configuration) "options" or "features". These terms are all equivalent and describe ways to adjust and optimize functional and non-functional properties of a software system [6]. One can divide into different types of options. Binary options usually have a value of 0 or 1 and describe the activation of a feature. Non-binary options support a wider

range of values. For example this could be a setting for the stack-size allowed for a program. Non-numeric options support the input of text. Those could be paths or other addresses. The set of all configuration options is denoted as \mathcal{O} .

A configuration can be defined in multiple different ways. Kaltenecker et al. [6] define a configuration as a function $c: \mathcal{O} \to \{0,1\}$. It assigns a 1 to each element of \mathcal{O} that is selected and a 0 to those which are not used. Guo et al. [2] and Nair et al. [9] have a similar approach. But instead of using a function to describe c they use a vector or an n-tuple over $\mathbb{Z}_2^+(=\{0,1\})$. Each position of those enumerations is associated with exactly one feature. As in Kaltenecker et al. [6] a 1 indicates an activation of a feature and a 0 means that the feature is not used. These definitions obviously describes binary options only, but can be expanded to support non-binary options by using the co-domain of \mathbb{N}_0 instead of $\{0,1\}$.

The *configuration space* describes all valid configurations of a system. It is denoted as C.

A *sample* is the subset of a *configuration space* that contains all configurations that are complied and measured during the process of sampling and learning. It may also contain the measured performance scores for each configuration.

To describe the quality of an approach an *accuracy* metric is often used. The *accuracy* is defined as 1-fault rate. And in turn the fault rate (or error rate) is defined as

$$fault rate = \frac{|actual - predicted|}{actual}$$
 (1)

This definition can be found in [9] and [12].

3 On the Applicability of the Brute Force Approach

As mentioned previously: brute force measuring is not feasible in most cases [12]. But what is the actual reason behind this? Let's have a look at an example first:

In one paper, Siegmund et al. [12] measured all valid configurations of multiple programs to analyse the accuracy of AFID. Berkley DB (C) was one of those programs. It is a database management program for embedded systems. It has 19 features and 2560 valid configuration. In the end, it took approximately 426 hours (= 17.75 days) to measure all these configurations. This value calculates to a time of about 10 minutes per configuration measurement. Considering that Berkeley DB (C) is a comparably small program, this is already a significant amount of time but still a time span that might be acceptable be used to get the perfect results of a $brute\ force$ approach.

This changes once one takes a look at larger programs. Modern applications like Apache or MySQL can have hundreds of configuration parameters [15]. Another example was displayed by Siegmund [11]: SQLite has 77 features that can produce $3\cdot 10^{77}$ valid configurations. Unfortunately, there is no evidence how the latter number was calculated. Yet, it can be assumed that at least the scale of this number in regard to the number of features is correct. This will be explained in

the next paragraph. Coming back to the example: Assuming measuring (compiling + profiling) one configuration would take 5 minutes, a brute-force approach would take $2.5 \cdot 10^{76}$ hours. Obviously, this is not an acceptable duration.

To find out why brute-force does not scale well, a generic look at how many configurations per program are needed to be measured helps. This set of all valid configurations can be written down as a Feature Model. An example for this is shown in Fig. 3. This feature model describes the software system called "DatabaseSystem". Feature Models can be annotated with further logical expressions to also contain cross-feature constraints. Such can also be seen in the just mentioned

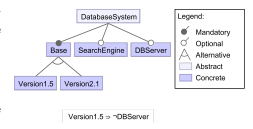


Fig. 3: An example of a feature model.

example. Each Feature Model can also be written down as a logical expression. The atomic variables of this expression are the options of the software system. In this context, a configuration is written down as an assignment of each variable of the expression. The tree in Fig. 3 is equivalent to the function

$$V(C) = \text{Base} \land (\text{Version1.5} \oplus \text{Version2.1}) \land (\text{Version1.5} \Rightarrow \neg \text{DBServer}).$$

For V(C) = 1, a configuration is considered *valid*. Otherwise, it is not accepted. An example configuration could look like this:

$$\begin{aligned} \textbf{DEFAULT} &= \{ \text{Base} = 1, \text{Version} 1.5 = 0, \text{Version} 2.1 = 1, \\ &\quad \text{DBServer} = 0, \text{SearchEngine} = 0 \} \end{aligned}$$

Note that these logical expressions can also be applied to non-binary options. The feature "Base" of the example could also be expressed as a tertiary value:

$$Base = \begin{cases} 0, Not \text{ selected} \\ 1, Version 1.5 \text{ selected} \\ 2, Version 2.1 \text{ selected} \end{cases}$$

V(C) would look like this, if Base is tertiary:

$$V(C) = (\text{Base} \Leftrightarrow 1 \vee \text{Base} \Leftrightarrow 2) \wedge ((\text{Base} \Leftrightarrow 1) \Rightarrow \neg \text{DBServer}).$$

In this context the most interesting property of this formula is the scaling of the number of valid configurations in relation to the number of options. Without loss of generality, one can assume that a system only has binary options. This can be done since all other types of options would just increase the overall number of options. However, already looking at binary options gives a satisfying result. Since naturally the total number of possible assignments for a logical expression is exponentially large, the number of valid configurations also lies in

the exponential space of $\mathcal{O}(2^{\#options})$. In other words: a configuration space that would have to measured for a brute force approach would be exponentially large. This exponential scaling is also the reason why brute-force does not scale well and more sophisticated methods are need for efficient predictions.

4 Measuring and Predicting the Performance of Highly Configurable Systems

By learning about the performance difference of multiple configurations it is possible to predict a program's performance accurately. This means, that not all (possibly exponentially many) configurations need to be measured. Instead, a small sample size should be enough to predict a program's performance. Multiple ways that use different methods have been proposed over time [9]. This paper will take a look at

- Automated Feature Interaction Detection by Siegmund et al. [12],
- o an incremental/statistical learning approach by Guo et al. [2],
- Cost efficient sampling by Sarkar et al. [10],
- WHAT a spectral learning approach by Nair et al. [9],

4.1 General Approach

Guo et al. [2] defines two problems that prediction approaches should solve:

- 1. Predicting the performance of a not measured configurations.
- 2. Finding a function f that shows the correlation between the properties of measured configurations and their performance value and that makes each predicted performance $f(\mathbf{x})$ of a configuration \mathbf{x} as close as possible to its actual performance.

$$f: \mathcal{C} \to \mathbb{R}$$
 such that $\sum_{(\mathbf{x}, y) \in S} L(y, f(\mathbf{x}))$ is minimal (2)

S is a sample and L is a loss function to penalize errors in prediction. (\mathbf{x}, y) is a pair of a configuration \mathbf{x} and its measured performance value y. The function f is also called the *performance model* of the system.

There is a general pattern for the solution of those problems that comes apparent when looking at different prediction approaches. It can be divided into two steps as displayed in Fig. 4.

The first step is to sample the exponential configuration space. This means finding configurations from which can be learned about the system. This is done by using efficient sampling techniques like spectral sampling [6] or progressive sampling [10].

Once enough and meaningful configurations are found, the learning process starts. Usually the previously chosen configurations are measured, under the

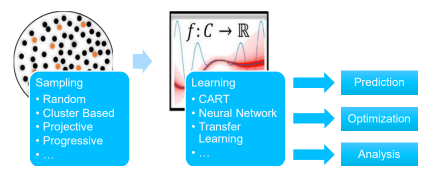


Fig. 4: General pattern of prediction approaches.

condition that this was not already done whilst sampling. The measurement results are fed to a learning process. A lot of different machine learning strategies can be applied [11]. The relative papers typically use CART's. Based on the found performance model continuing tasks like finding near optimal solutions or in-depth performance analysis can be done [11].

4.2 Automated Feature Interaction Detection

Automated feature interaction detection (AFID) is a measurement-based approach to predicting the performance of a highly configurable system. It was developed by Siegmund et al. [12]. The following section is also based on the proposing paper of AFID [12]. Under the usage of linear regression AFID tries to determine a performance influence value for each feature and feature interaction. A feature interaction is defined as an unexpected influence on the performance of a system when using a specific feature combination. Furthermore a goal of ?] was also to find explainable predictions. In the conducted experiments of Sarkar et al. [10] and Siegmund et al. [12] this method reaches average accuracies of 85% and 95% respectively.

The general process of AFID is displayed in Fig. 5. It can be divided into two different steps:

- 1. Finding interacting Features.
- 2. Measuring the performance influence of feature interactions.

For the first step some notation is needed. For simplicity AFID is defined for binary options only. The composition of performance influencing units (features or feature interactions) is denoted by a \cdot . In case two features are used simultaneously it is denoted by using a \times . This would also be another way to describe a configuration. So a program P that uses the two features a and b can be denoted as $P = a \times b$.

To get the performance of P, $\Pi(P)$ has to be calculated. The exact definition of the performance influence determining function Π can be found in the paper of Siegmund et al. [12]. For now, it is important to note that when

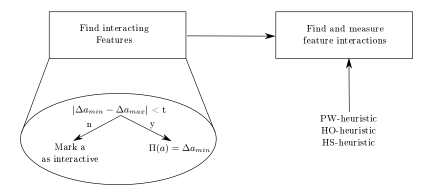


Fig. 5: General steps of *AFID*.

calculating $\Pi(P) = \Pi(a \times b)$ not only the performance influence of a and b are necessary, but also a#b has to be considered. The latter is the performance influence of the possible interaction between a and b. This means $\Pi(a \times b) = \Pi(a\#b) + \Pi(a) + \Pi(b)$. Higher order interactions are also possible: When using a program configuration $P_2 = a \times b \times c$ then $\Pi(P_2) = \Pi(a) + \Pi(b) + \Pi(c) + \Pi(a\#b) + \Pi(a\#c) + \Pi(b\#c) + \Pi(a\#b\#c)$. Some interactions do not exist or have an influence on the system, those will be ignored by the algorithm. Otherwise, a brute-force solution would be equally efficient.

Finding these interactions requires looking for all interacting features first. This is done in by intelligently measuring certain configurations. AFID defines a feature a as interacting when

$$a \text{ interacts} \Leftrightarrow \exists C, D \subseteq \mathcal{C} | C \neq D \land |\Delta a_C - \Delta a_D| \le t$$
 (3)

with

$$\Delta a_C = \Pi(C \times a) - \Pi(C)$$

= $\Pi(a \# C) + \Pi(a)$. (4)

t is a threshold depending on the given performance metric. Using these two equations it can be determined, whether a feature is interacting with other features using 4 measurements. These include $\Delta a_{min} = \Pi(a \times min(a)) - \Pi(min(a))$ and $\Delta a_{max} = \Pi(a \times max(a)) - \Pi(max(a))$. min(a) is a configuration, that contains the minimum possible features without using a. Simultaneously, max(a) is also a configuration that contains the maximum amount of possible features without a. Once these measurements are done Eq. (3) can be applied with $C = \Delta a_{min}$ and $D = \Delta a_{max}$. This is done for all features to find the interacting ones. If a feature f is not found to be interactive, its performance influence $\Pi(f)$ equals Δf_{min} .

Once all interacting features are found, the search for the actual interactions starts. In this second step three different heuristics are used to determine which interactions are searched for.

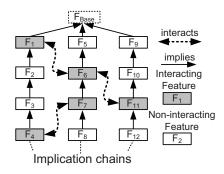


Fig. 6: Implication tree example found in [12].

Pair-Wise Heuristic Most groups of interacting features appear in the size (PW): of two [7, 12]. So it makes sense to look for pair interaction first.

Higher-Order Interactions Siegmund et al. [12] only look at higher order inter-Heuristic (HO): actions of the rank of three. Even higher ranks would take up too many measurement resources.

Hot-Spot Features Based on [1] and [14] Siegmund et al. [12] assume that Heuristic(HS): hot spot features exist. At last these specific type of interactions are findable too.

The three heuristics will be applied in the order of $PW \to HO \to HS$ and use the data provided by the previous ones. Using a SAT-Solver an implication-graph as seen in Fig. 6 is generated. Each implication-chain in this tree should have at least one interacting feature. When analysing the tree each chain is walked from the top down.

First the influence of every feature on another chain is measured (PW-heuristic). In the example of Fig. 6 the interactions would be measured in this order: F1#F6, F1#F7, F4#F6, F4#F7, F6#F11, F7#F11, F1#F11, F4#F11. Once an interaction impact exceeds a threshold it is recorded.

Secondly, the higher order interaction heuristic can be applied. Seconded order interactions can be relatively easily found by looking hat the results of the PW-Heuristik. Three features that interact pair-wise are likely to interact in a third order interaction. For example, looking at features a, b and c. If $\Delta a \# b_{C1}$ and $\Delta b \# c_{C2}$ have been recorded, $\{a\# b,b\# c,a\# c\}$ all have to be non-zero to find a third order interaction. Interactions with and order higher than three are not considered to prevent too many measurements.

Lastly, Hot-Spot features may be detected (HS-heuristic). This is done by counting the interactions per feature. If the number of interactions of a feature is above a certain threshold (e.g. the arithmetic mean) it is categorized as a Hot-Spot feature. Based on these hotspot features further third order interactions are explored. Again higher order interactions are not considered.

After applying the three heuristics all detected interacting features or feature

combinations are assigned a Δ to represent their performance influence on the program.

As mentioned, Siegmund et al. [12] tested AFID on six different software systems. Each program was tested under four approaches: Feature-Wise, Pair-Wise, Higher-Order, Hot-Spot (in this order). Each approach also used the data found by the previous one. Accordingly, the results get better the more heuristics are used as Table 1 shows. Using only the FW-heuristic means, that interactions are

Table 1: Results of average accurcy found by Siegmund et al. [12].

Approach	avg.	Accuracy
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FW	79.7%
PW	91%
HO	93.7%
HS	95.4%

not considered at all. However, the accuracy when using this heuristic is already at 80% on average. A significant improvement can be made by using the PW heuristic. It uses 8.5 times more measurements than the FW-heuristic on average, but improves the accuracy to 91%. Using the HO- or HS-heuristics improves the accuracy further by 2-4%. However, for Apache, using the HO over the PW heuristic even deteriorated the average result by 3.9% and doubled the standard deviation. As expected, using the HS-heuristic gives the best accuracy for all 6 tested applications. Siegmund et al. [12] also notes that analysing SQLite only needed about 0.1% of all possible configurations. This hints to the good scalability of AFID.

In conclusion AFID measures only a quadric amount of configurations to find the performance model of the software system. Depending on which heuristics are chosen, the accuracy and measurement costs can variate. The goal giving explainable predictions was also. Siegmund et al. [12] implemented this approach in their tool SPLConquerer¹. This tool was also used to produce all experiment results.

4.3 Classification and Regression Trees

The next 3 approaches all use a specific type of machine learning strategy to construct their predictors. This method is call Classification and Regression Tree's (CART). These trees are typically binary trees that divide the given data points into small enough groups so a direct local prediction can be done. These local prediction are then combined into a global predictor [2]. In the context of configurable software systems each point consists at least out of a configuration and an associated performance score. These data point are then fed into the algorithm seen in Listing 1.1.

The node impurity found in the algorithm is typically calculated by the square mean error. To prevent under- or overfitting[3] the recursive splitting has to be stopped at the right time. This is possible by manual parameter tuning or using an empirical-determined automatic terminator. The size of the predictor

¹ https://github.com/se-passau/SPLConqueror

- 1. Start at the root node. Assign all configurations to it.
- 2. For each option, find the set of options X that minizes the sum of the node impurities in the two child nodes. Divide the configurations assigned to the current node based on X into two disjoint sets and assign those to the corresponding child nodes.
- 3. If a stopping criterion is reached, exit. Otherwise, apply step 2 and 3 to each child node inturn.

Listing 1.1: Pseudocode for generating a *CART*. Adopted from Loh [8] to fit software configurations.

variables set X is usually 1. This means, that we decide the branching based on whether a feature is selected or not (or based on its value). When using the CART found in Fig. 7a configuration $c=\{\ldots,\ x_3=1,\ \ldots,\ x_{14}=0,\ x_{15}=0,\ \ldots\}$ would be classified as S_{RRL} . The predicted performance of this class and therefore for c is $\ell_{S_{RRL}}=571$.

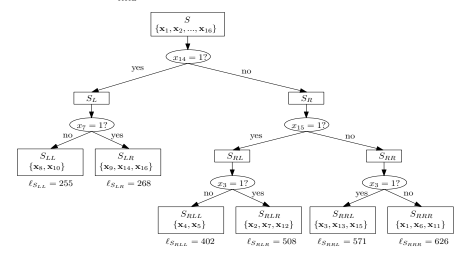


Fig. 7: Example performance model of X264 generated by CART based on the random sampling (N=16), using minimization of the sum of squared error loss [2].

4.4 Variability aware Performance Prediction

Variability aware Performance Prediction (VAPP) is a statistic based approach to performance prediction. With the help of random sampling and CARTs a simple yet effective predictor can be build. The following section is based upon

Guo et al. [2]. In their own tests Guo et al. [2] reached an average precision of 94% whilst using a sample as large as the ones *AFID* would be using under the PW-heuristic. Further, tests conducted by Nair et al. [9] with the same sample size showed an accuracy of 92.4%.

The basic idea of variability aware performance prediction is shown in Fig. 8. Two cycles can be found:

- \circ The first cycle is outside of the dashed box and describes the basic inputoutput behavior of a predictor. A user configures a new configuration \mathbf{x} for System A and asks the predictor (dashed box) for a prediction. It replies with a quantitative prediction for \mathbf{x} 's performance.
- o In the second cycle a actual prediction is generated based on decision rules which themselves are in turn created by simplifying a performance model (a CART). Random sampling is used to learn the performance model.

Like other approaches, the target of variability aware performance prediction is to get accurate predictions whilst only using a small sample for the creation of the performance model. Nonetheless, VAPP offers a free choice of the sample size. The configurations of the sample are chosen randomly out of C.

VAPP uses the tuple-based definition of a configuration. It further defines that each configuration $c_j \in \mathcal{C}$ has an actual performance value y_j . For formal correctness it is assumed that every option of a configuration actually influences the performance of the system. Otherwise, a CART could not be applied.

In the used CART each sub-tree is also called a segment S_i , where i determines the location of the sub-tree. This is also shown in Fig. 7.

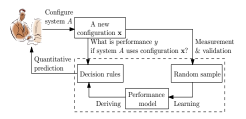


Fig. 8: Overview of the Approach of Variability aware Performance Prediction [2].

For the *local model* ℓ of the used CART Guo et al. [2] choose the arithmetic average:

$$\ell_{S_i} = \frac{1}{|S_i|} \sum_{y_j \in S_i} y_j \tag{5}$$

As a loss function to penalize the prediction errors (node impurity) the sum of squared error loss is selected.

$$\sum_{y_j \in S_i} L(y_j, \ell_{S_i}) = \sum_{y_j \in S_i} (y_j - \ell_{S_i})^2$$
 (6)

Therefore the best split for a segment S_i is found when

$$\sum_{y_j \in S_{iL}} L(y_j, \ell_{S_{iL}}) + \sum_{y_j \in S_{iR}} L(y_i, \ell_{S_{iR}})$$

is minimal.

Assuming there are q leafs in a tree than the predictor function $f(\mathbf{x})$ is defined as:

$$f(\mathbf{x}) = \sum_{i=1}^{q} \ell_{S_i} I(\mathbf{x} \in S_i)$$
 (7)

where I is an indicator function to indicate whether x belongs to a leaf S_i . For the example of Fig. 7, f(x) unwraps to:

$$\begin{split} f(x) &= 255 * I(x_{14} = 1, x_7 = 0) \\ &+ 268 * I(x_{14} = 1, x_7 = 1) \\ &+ 402 * I(x_{14} = 0, x_{15} = 1, x_3 = 0) \\ &+ 508 * I(x_{14} = 0, x_{15} = 1, x_3 = 1) \\ &+ 571 * I(x_{14} = 0, x_{15} = 0, x_3 = 1) \\ &+ 626 * I(x_{14} = 0, x_{15} = 0, x_3 = 0) \end{split}$$

Every possible configuration x is associated with a leaf of the tree. Therefore, f(x) can always be applied.

For their Experiment Guo et al. [2] test the same software systems as Siegmund et al. [12] (Section 4.2). They also compared their prediction results with the results produced by AFID.

Since unlike AFID the size of a sample for VAPP can be chosen freely, some comparable sample sizes were chosen. Guo et al. [2] use 4 different sample sizes based on the size of the tested programs. For a program with N features they use samples the size of N, 2N, 3N and M. M is the amount of configurations measured by AFID using the PW-heuristic. It was found, that the prediction accuracy increases linear with the size of a sample. For a small sample with the size of N the prediction accuracy was at above 92% in 3 cases. However, for Berkeley DB (C) the prediction accuracy with an N sized samples was at 112.4% with a standard deviation of $\pm 354.6\%$. This shows that VAPP is not generally applicable for small samples. Using a sample size of M significantly improves the average prediction accuracy to a stable average of 93.8%.

4.5 Cost-Efficient Sampling

Just how its name suggests, cost-efficient sampling tires to minimize the cost-accuracy rate of a prediction approach. This is done by trying to find a (near) optimal sample size n^* for a given program.

Sarkar et al. [10] apply two different sampling techniques to reach this goal. Their general structure is displayed in Fig. 9.

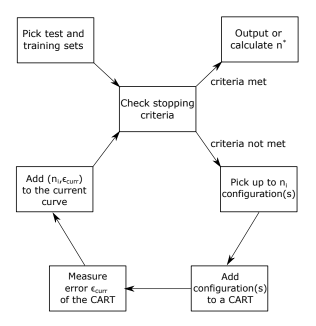


Fig. 9: General procedure of cost-efficient sampling to find an optimal sized n^* .

Firstly a test and a training set are picked from C. Then, in each iteration of this process, a performance model is build. The used sample is taken from the training set and is increased each iteration. This makes the respective performance model more accurate in each iteration. The error rate of each created performance model is recorded. Together with the size of the used sample a point in a graph is created. As iterations continue this graph resembles an approximation of the learning curve of the performance model of the system. A curve as displayed in Fig. 10 is created. This process continues until a stopping criterion is met. At this point n^* was already found or can be calculated based on the given curve and a cost-function. An optimal sample size should provide a good accuracy without being too large. Naturally n^* lies some-

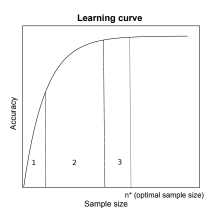


Fig. 10: A typical learning curve can be divided into three phases:

- 1. steep incline;
- 2. gradual incline;
- 3. plateau [10].

where at the beginning of the third phase.

Four different types of costs are considered by Sarkar et al. [10]:

$$TotalCost = Cost_{Measuerment(Training)} + Cost_{Measuerment(Testing)} + Cost_{ModelBuilding} + Cost_{PredictionError}$$

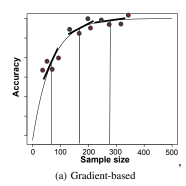
$$(8)$$

Based on this, the TotalCost for a sample the size of n is defined as:

$$TotalCost(n) = 2n + \epsilon_n \cdot |S| \cdot R \tag{9}$$

S is a score set that contains all configurations, whose performance value will be predicted with the current model. R is tuning parameter for the measuring cost of a training set.

The two sampling strategies based on this are progressive sampling and projective sampling. Both approach the search for n^* differently.



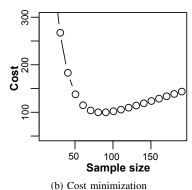


Fig. 11: Stopping criteria of progressive sampling [10].

Progressive sampling can be divided into two different strategy types. They differ in the way the next sample size n_{i+1} is calculated:

1. arithmetic: $n_i = n_0 + i \cdot a$ 2. geometric: $n_i = n_0 + a^i$

The constant parameter a determines the growth-rate of the samples. On one hand arithmetic progressive-sampling is more precise, but on the other hand we need to build more performance models in comparison to geometric progressive-sampling. For progressive sampling there are two types of stopping criteria:

- 1. Gradient-Based: Additional models around n_i will be build so the gradient around at n_i can be determined. Fig. 11a shows an example of this. Once the gradient or accuracy reaches a certain threshold the iterative process is stopped and n^* set as n_i .
- 2. Cost Minimization: For each n_i the current error-rate of the performance model is substituted into the cost function of Eq. (9). For well-behaved learning curves the cost function is convex as displayed in Fig. 11b. Once the cost-function increases for the first time, the iterative process is stopped and n^* set as n_{i-1} .

Projective sampling tires to find the actual function behind the learning curve of the performance model of the current system. Typical equations that describe a learning curve and the respective minimum of the cost-function can be found in Table 2. Unlike in progressive Sampling the size of the sample set is increased by only a small amount each iteration. The experiments conducted by Sarkar et al. [10] show that even an increment by the size of 1 is enough to get satisfying results. This time the iterative process is stopped once a certain accuracy is reached. Now an algorithm tires to fit the generated points onto the four functions shown in Table 2. The best fitted function is considered the learning curve of the current performance model. Once this function is found, it can be substituted into to cost-function Eq. (9) and the minimum of which can be calculated. These minimums are also shown in Table 2. Based on the found

Table 2: Functional representation of typical learning curves and thier optimal sample size based on the given cost function [10].

Name	Equation	Optimal Sample Size
Logarithmic	err(n) = a + b.log(n)	$n^* = -(\underline{R \cdot S \cdot b})/2$
Weiss and Tian	err(n) = a + bn/(n+1)	$n^* = \sqrt{(-R \cdot S \cdot b)/2}$
Power Law	$err(n) = an^b$	$n^* = \left(\frac{-2}{R \cdot S \cdot a \cdot b}\right)^{\frac{1}{b-1}}$ $n^* = \log_b \left(\frac{-2}{R \cdot S \cdot a \cdot \ln b}\right)$
Exponential	$err(n) = ab^n$	$n^* = \log_b \left(\frac{-2}{R \cdot S \cdot a \cdot \ln b} \right)$

minimum other techniques that do not have a fixed sample size like *VAPP* can be used. In the experiments conducted by Sarkar et al. [10] *projective sampling* was better than *progressive sampling* in every case. For each of the six tested software systems *projective sampling* had a better accuracy whilst having a lower cost than *progressive sampling*.

For both strategies it is important, that the initial sample generation/picking is successful. Therefore, the picking of representative configurations as n_i samples of each iteration is key. Sarkar et al. [10] use a heuristic based on feature frequencies to optimize the selection. As a general guideline each feature should be selected and deselected once at least. Looking at a set of configurations S the feature frequency of a feature i is defined as:

$$1 \le \sum_{j \in S} x_i(j) < |S| \tag{10}$$

Where $x_i \in \{0, 1\}$ describes, whether the feature i is selected in the configuration j or not. This feature frequency is recorded for whilst creating a new sample. In case a certain threshold is reached, the sample generation will be stopped.

The results of the experiments conducted by Sarkar et al. [10] show that projective sampling is superior to progressive sampling in all cases. It had a lower costs

and a higher accuracy when predicting configurations of the six tested programs. After the optimal sample size was found Sarkar et al. [10] used VAPP with an n^* sized sample to get the prediction results. The accuracy of this approach reached up to 99% and had an average of 94.5%.

4.6 WHAT

WHAT is a spectral learning approach developed by Nair et al. [9]. The following section is based on proposing paper of Nair et al. [9]. WHAT aims to find an accurate and stable performance model with fewer samples than the previous methods. To reach this goal it uses spectral and regression tree learning. The idea behind spectral learning is the mathematical concept of eigenvalues/vectors of a distance matrix between configurations. This has the advantage of automatic noise reduction. Nair et al. [9] explain, that when a data set has many irrelevancies or closely associated data parameters d, then only a few eigenvectors $e, e \ll d$ are required to characterize the data. To find these important eigenvalues/-vectors a simple clustering algorithm is applied to $\mathcal C$. The main advantage of this approach are a reduced sample size and a lower standard deviation compared to previously shown methods [9]. Nair et al. [9] divide WHAT into 3 parts.

1. Spectral Learning

This first step of spectral learning is used to cluster all valid configurations. As each configuration is an n-dimensional vector (or n-tuple) it can be placed in an n-dimensional space.

WHAT gets N different valid configurations as input and is picks a random configuration N_i and two configurations West and East. West is the configuration that is most different to N_i and East is the configuration most different to West. In mathematical terms the 'most different' means the largest euclidean distance between two configurations. After that, a straight through East and West is calculated and all configuration are dived into two clusters. This division is based on the median value of all calculated distances. This process is recursively repeated for each sub-cluster until they reach a threshold size. Nair et al. [9] use $\sqrt{|N|}$ as their termination value. Unlike other clustering algorithms like K-Means, spectral learning runs in linear time of $\mathcal{O}(2|n|)$.

2. Spectral Sampling

For the actual sampling different strategies can be applied. The most important is random sampling, where one configuration of each leaf cluster is randomly picked as a representative. This representative is then compiled and measured. There are also two other sampling strategies mentioned. East-West sampling complies and executes East and West of the current leaf-cluster. Whereas Exemplar sampling measures all configurations of a leaf-cluster, but only returns the lowest performance score as a representative. Both of these strategies get outperformed by the random sampling strategy.

3. Regression-Tree Learning

In this step a CART is build from the chosen samples. This time the best split is defined as reaching the minimum of $\frac{A}{N}\sigma_1 + \frac{B}{N}\sigma_2$, σ_1 and σ_2 are the standard deviations of the corresponding sub-trees. From this CART again decision rules can be derived.

Results of testing WHAT on the programs we introduced earlier show that it has an average precision of 93.4%. Furthermore, the standard deviation is comparably low, which was one of the original targets of WHAT [9].

5 Comparison

Table 3 displays the results of an experiment conducted by Nair et al. [9], to compare the different prediction methods. The table shows some characteristic properties of each approach.

Siegmund's approach of AFID was ranked last on 4 occasions. Its accuracy and standard deviation are the worst in most cases. It mostly ranks lower than VAPP whilst utilizing same sample size (Guo(PW)). WHAT is the oldest of the presented approaches and its low ranking can be seen as a demonstration on how prediction algorithms evolved over time.

Both versions of VAPP appear inconsistently with regard to their rank. Gou(PW) ranks lower than Gou(2N) in half of the six occasions. Generally the version which used a larger sample had a slightly better accuracy, but also suffered from a larger standard deviation. The results show that VAPP's predictions might not be consistent for not sufficiently large enough samples. This could be attributed to the complete randomness of configuration picking.

Sarkar's approach of using cost-efficient sampling in combination with CARTs ranked best in four out of the six cases. In the remaining two cases it still had an acceptable accuracy. But this consistent high accuracy comes with the cost of using larger samples then other methods. In the case of SQLite, the optimized sample size was 15 times larger than the sample of the next best approach WHAT.

Considering all tests WHAT had an average standard deviation of only 2.98%. Having such a low standard deviation was the main goal of WHAT. When comparing it to other methods, it consistently has a below average standard deviation and an above average accuracy whilst using a smaller sample. Since WHAT is also the most recent approach, this further confirms the progress that was made in recent years.

The paper ([9]) defines A and B as sets and N as a (natural) number. So it may be assumed that the formula actually should be $\frac{|A|}{N}\sigma_1 + \frac{|B|}{N}\sigma_2$. This does make sense, since this formula weights both standard deviations σ proportional to total number of current configurations N.

Table 3: Measureing results of Nair et al. [9] when comparing the methods of, AFID(Siegmund), VAPP(Gou), WHAT and projective sampling in combination with CARTs (Sarkar). The Rank column is computed using Scott-Knott, bootstrap 95% confidence, and A12 test [9].

Apache	Rank	Approach	Mean MRE(μ)	$STDev(\sigma)$		#Evaluations
1 Guo(PW) 10.51 6.85 — 29 1 Siegmund 10.34 11.68 — 16 1 Guo(2N) 13.03 15.28 — 18 BDBC 1 Sarkar 1.24 1.46 — 191 2 Siegmund 6.14 4.41 — 139 2 WHAT 6.57 7.40 — 64 2 Guo(PW) 10.16 10.6 — 139 3 Guo(2N) 49.90 52.25 — 36 BDBJ 1 Guo(2N) 2.29 3.26 — — 52 1 Guo(PW) 2.86 2.72 — 48 2 Sarkar 5.67 6.97 — 48 2 Sarkar 5.67 6.97 — 48 2 Siegmund 6.98 7.13 — 64 1 WHAT 3.32 1.05 — 32 1 <	Apache					
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6 Continuing, Related and Future Work

This paper covered different types of prediction approaches developed by N. Siegmund et al.

Many other techniques can be found that are related or used for performance prediction of configurable software systems. Some notable continuing techniques are:

T-Wise Sampling picks configurations that contain every combination of T different features to ensure a diverse sample [5].

Fourier Learning theoretically guarantees a accuracy level whilst using a minimal sample. It is based on the Fourier transform and was developed by Zhang et al. [16].

Transfer Learning uses prediction models that are generated based on simulations of the observed system. These models then get transferred to a predictor of the real system to improve its predictions [4].

As shown, a lot of different prediction approaches with satisfying accuracies are available. To further improve them Nair et al. [9] propose two different starting points:

- All presented approaches consider all features as equally important. But in some systems a certain option might has a higher priority than others. Hence, weighting techniques for features could improve prediction results.
- Currently, the sizes of samples are mostly picked manually. This can lead to
 problems with adaptability and scalability. A future field of research would be
 the integration of a dynamic progressive sampling techniques, as replacement
 for the typically used static sample size.

Furthermore, Siegmund et al. [13] note that finding valid configurations, for binary and non-binary options is still a problem with exponential complexity. Improvements on related algorithms would lead to a faster discovery of the *configuration space*, test and training sets.

Most approaches concentrate on binary and numeric options only. Siegmund et al. [13] show that both can be integrated smoothly, into a prediction approach. However, non-numeric options like paths are not mentioned much in the literature. For example, a database-path can arguably have a huge impact on the performance of a software system. The extension or development of techniques that support non-numeric option could improve predictions for more complex software systems that depend on those kinds of options.

7 Conclusion

There are many techniques for learning and predicting the performance of a configurable software system. This paper had a look at different approaches developed by N. Siegmund et al. It was quickly determined that, if enough time and resources are available or a program is sufficiently small, brute force can be applied to get perfect prediction results. But since brute force does not scale well, more sophisticated methods were developed. Most of those methods produce acceptable results with over 90% accuracy in many cases. But there is no best approach. Each technique has its own advantages and disadvantages that make it unique. AFID has well explainable results, VAPP is easy to implement, WHAT has a very low standard deviation and when using cost-efficient sampling extremely accurate predictions are possible. Which method a predictor should use is always a question of finding a balance between the possible size of the sample, the targeted accuracy and the applicability of the approach on the current system.

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