Optimization of Software Estimation Models

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Abstract: In software engineering, estimations are frequently used to determine expected but yet unknown properties of software development processes or the developed systems, such as costs, time, number of developers, efforts, sizes, and complexities. Plenty of estimation models exist, but it is hard to compare and improve them as software technologies evolve quickly. We suggest an approach to estimation model design and automated optimization allowing for model comparison and improvement based on commonly collected data points. This way, the approach simplifies model optimization and selection. It contributes to a convergence of existing estimation models to meet contemporary software technology practices and provide a possibility for selecting the most appropriate ones.

1 INTRODUCTION

Several software engineering steps depend on estimations, e.g., estimating the complexity and the effort for implementing a particular task or estimating the time and costs of implementing a system. Consequently, there are plenty of methods and models suggested for these estimation tasks. We will introduce known methods and models for software estimation and continue with problem description and analysis to end the introduction presenting our research aims.

1.1 Known Methods and Models

There are plenty of known software estimations models, e.g., COCOMO, the first version known as COCOMO 81 and the second as COCOMO II (Boehm et al., 2009), COSYSMO (Valerdi, 2004), Evidence-based Scheduling (a refinement of typical agile estimating techniques using minimal measurement and total time accounting) (Spolsky, 2007), Function Point Analysis (Albrecht, 1979), Parametric Estimating (Jenson and Bartley, 1991), Planning Game (from Extreme Programming) (Beck and Andres, 2004), the ITK method (also known as the CETIN method), Proxy-based estimating (PROBE, from the Personal Software Process) (Humphrey, 1996), Program Evaluation and Review Technique (PERT) (Office, 1962), the Putnam model (also known as SLIM) (Putnam and Myers, 1991), the PRICE Systems (commercial parametric models that estimates the scope, cost, effort and schedule for software projects), SEESEM (Galorath and Evans, 2006) (parametric estimation of effort, schedule, cost, risk.) Minimum time and staffing concepts based on Brook's law (Brooks, 1995), the Use Case Points method (UCP) (Karner, 1993), Weighted Micro Function Points (WMFP), and Wideband Delphi¹.

1.2 Problem Description and Analysis

Modern software technology advances at a fast pace including development tools and environments, platforms, reusable components and libraries, languages, and processes. What is appropriate software technology today might be outdated in the near future. Therefore, also the estimation models need to change at the same speed because, otherwise, they become outdated and, hence, imprecise for modern software development.

Cloud technologies used in software development have the potential to provide us with access to previously unknown amounts of data, i.e., Big Data.

¹https://en.wikipedia.org/wiki/Cost estimation in software engineering
Converted into actionable knowledge, data has become an unprecedented economic value as witnessed by the success of companies like Google, Facebook, and Twitter. Big Data has also revolutionized software engineering research and its scientific methodologies leading a paradigm shift away from data-scarce, static, coarse-grained, and simple studies towards data-rich, dynamic, high resolution, and complex observations and simulations.

However, Big Data has not yet been fully exploited in estimation model building, validation, and exploitation. One reason might be that the relevant data is distributed among different stakeholders in an organization, e.g., relevant impact factors of development time are task complexity (estimated by the development team), team competence and availability (assessed by the personnel lead), and processes and environment maturity (evaluated by a CTO). Another reason might be that the relevant data becomes available at different points in time. For instance, the aforementioned factors of estimation are available before the project starts, while the actual ground truth only gets available afterward. Further to complicate matters, data points can be generated in different organizations and, hence, are hard to collect and to exploit commonly.

As a consequence, we face a lot of shortcomings in the currently suggested estimation models, despite all benefits of cloud and Big Data technologies. Estimation models are built from prior data collected years before and looking at software development practices of today; they are outdated. Consequently, they emphasize factors that are nowadays irrelevant and neglect factors of high contemporary importance. The model constants of these models are often outdated too and cannot be adapted to today’s technologies without significant effort. There is no automatic, continuous learning and improvement.

The models are incomplete, e.g., they estimate the size but not the effort, or estimate the effort based on the size but do not estimate the size itself. They are also incompatible with each other, i.e., they require similar, though not identical, factors. As an example, the development environments and product factors are regarded both to impact on the estimated system size and the development effort, which gives these factors too high weights overall.

Developing new estimation models is not a solution to the aforementioned shortcomings if the process of their development would not change. We claim that the estimation model development faces some inherent problems. New models cannot be tested without a significant cost in time and effort. The errors and, hence, the suitability of these models cannot be calculated automatically and in a universal way. Also, some estimation factors, e.g., parameters of a project, partially overlap, are dependent on each other, i.e., they correlate. If the correlation is strong, they are redundant and therefore, superfluous. To date, this can only be recognized and optimized with a manual effort, if done at all.

To train a new model, more data points have to be collected first. Given the fact that there are often not enough data points available or they do not exist in a uniform structure because no uniform process exists for collecting the data we face a so-called “cold start problem”. This leads to a lot of effort for data collection and model estimation before gaining any value. However, even imprecise objective estimation models have a value compared to and complementing subjective assessments.

### 1.3 Research Aims

This research contributes with an approach to test and improve estimation models mapping, e.g., cost drivers to costs. The approach shall be agnostic with respect to the specific domain and help to test and improve models for mapping any indicator of an outcome to the actual outcome.

Additionally, the following requirements shall be met: The approach is data-driven and improves with newly available data points. In such an approach, new model ideas should be easily trained and tested with old data points. New models should coexist and compete with previously defined models. The approach supports the calculation and comparison of the accuracy of all competing models based on all data points and adjusts the parameters of each model to minimize potential errors. The approach should also be understandable by human experts.

The goal of this work is not to develop one or more concrete, accurate estimation models. Because of the cold start problem and the fact that nowadays technologies change faster, simultaneously makes the continuous learning of the coefficients more important, our approach should help improve models and increase their accuracy over time. Moreover, it should not matter what a model looks like, i.e., what class of functions it exploits nor its domain. Instead, the approach shall be universal. If sufficient initial data points are available, arbitrary models can be tested, trained, iteratively improved, and eventually used.

The following research questions guided our work:

**RQ1** How can we define a continuously improving approach where new models are easy to implement, validate, and adjust?
RQ2 How can data points be collected while detecting correlations and dependencies between inputs while avoiding unnecessary inputs?

RQ3 Can we automate the approach?

In short, instead of suggesting the one and only model for estimations we aim at establishing an effective and efficient process to collect data points and develop different competing estimation models based on these data points that are useful at any point in time and that converge to more and more accurate tools.

We rely on a design science approach (Simon, 1996) and formulate the goal criteria as well as the design of our approach in Section 3. We evaluate an implementation of our approach in Section 4 against these goal criteria and discuss how well these are fulfilled in Section 4.5. In Section 5, we present our concluding remarks and our thoughts on future work.

2 THE COCOMO FAMILY OF SOFTWARE ESTIMATION MODELS

While our approach is agnostic to the actual estimation problem and the model to estimate, we still want to introduce a concrete model as an example for better relating to the issues and challenges.

The Constructive Cost Model (COCOMO) (Boehm et al., 2009) is arguably still the most widely used software implementation effort estimation model. It can also be applied to estimating the re-implementing effort. It was developed in 1981 and last updated with “new” calibration data—only 161 data points—in 2000. It aims at estimating the effort $E$ in person months depending on the estimated project size $N$ in lines of code and on different project and team parameters. Different model input parameters—so-called effort multipliers ($em$) and scale factors ($sf$)—have an impact on the estimated effort $E$, such as Development Flexibility, Team Cohesion, and Software Cost Drivers, which are then subdivided into Product, Personnel, Platform, and Project Cost Drivers.

The model is weighted with model constants $a$ and $b$ that need to be calibrated using data points from real projects. These are the formulas behind COCOMO:

$$
E = a \times (N/1000)^f \times \prod_{i=1}^{17} em_i
$$

$$
e = b \times 0.01 \times \sum_{i=1}^{5} sf_i
$$

Similar to COCOMO, the Constructive Systems Engineering Cost Model (COSYSMO) uses the size of a system and adapts it according to the corresponding parameters. It was developed in 2002 and was calibrated with approximately 50 projects. It estimates person months as a function of the four size parameters: number of system requirements, number of system interfaces, number of algorithms, and number of operational scenarios (Valerdi, 2004).

With COCOMO and COSYSMO software engineers can estimate certain cost driving factors (independent variables), for example, lines of code, and then apply the model to estimate time and costs (dependent variables). Since function classes for mapping independent to dependent variables are known, regression of the function parameters can be done and redone. However, currently suggested parameters are originally trained on very few data points.

Based on the same data points, there is no possibility to create, test, and optimize parameters of new models, since the relevant independent variables and the function class of the model are static. The models are outdated and do not regard today’s practices in software developments, for example, new development processes like the agile approach. The projects on which COCOMO is based are all realized with the waterfall concept. It seems that COCOMO overestimates the effort since some cost drivers that had a high impact back in the time when they been defined are less important now. They are less influential nowadays due to technological progress, e.g., larger scale projects are easier to manage today than in 2000. In short, COCOMO and COSYSMO do not have any function type calibration capabilities and do not consider the addition of new parameters. At best, these models could provide an upper bound of the potentially achievable accuracy.

A recent study on effort estimation found that if a project gathers COCOMO-style data, none of the newer estimation models did any better than COCOMO (Menzies et al., 2017). They conclude that if such data were available, it should be used together with COCOMO to perform predictions.

The upcoming COCOMO III was first announced in 2015 (Clark, 2015) and should consider the scope of modern software. Especially, new development paradigms, new software domains like Software as a Service, mobile, and embedded as well as big data should be addressed. It should improve the realism and contain new and updated cost drivers. Unfortunately, the announcement has been out for a long time, and it is not known when or if at all, COCOMO III will ever be released.
It is not easy to keep old models updated or suggest new models, prove them superior, and abandon the previous ones.

3 ESTIMATION MODEL DEVELOPMENT

3.1 From Informal Requirements to Measurable Goal Criteria

The goal of this work is to help users develop models that improve over time. Model developers can thus add models. Along with new data points being added continuously, the accuracy of the model should improve. It should not matter what a model looks like, i.e., what class of functions it exploits or what domain it has. Instead, the approach should be universal; if a sufficient number of initial data points are available, arbitrary models can be tested, trained, iteratively improved, and eventually used. We define the following Goal Criteria (GC) for our solution refining the requirements posed in Section 1.3:

GC 1: The models should be able to make predictions based on learning from available data. Therefore, a suitable approach would allow collecting data points in central data storage and a consistent format. Data should be cleaned up, and it should be possible to deal with missing data values.

GC 2: The accuracy of the predictions should improve continuously through model optimization.

GC 3: Possible model improvements should be suggested automatically, e.g., if some parameters in a model become superfluous.

GC 4: It should be possible to determine if a prediction is trustworthy, e.g., by determining if it is based on outlier data.

GC 5: The models should be possible to inspect and be understandable by human domain experts. The influence of specific inputs on the result should be recognized.

GC 6: The approach should be universally applicable to any models and domains.

Further note that machine learning is often used to create models that can make predictions based on input data. However, these models are generally “black boxes” that are difficult to understand and learn from. Hence, while machine learning would fulfill some of the goal criteria, it would violate GC 3 and 5. Machine learning models also often exclude principle function classes and may result in over-trained or over-fitted models, especially when only a few data points are available. We instead rely on “white box” models. Designed parametric models that are explicitly defined based on expert domain knowledge, e.g., within software project management. While these might perform worse compared to machine learning models for some goal criteria, such as 2, they do fulfill all goal criteria.

3.2 Mathematical Foundations

A parametric model is provided as a function, e.g., $Y = f(x) = a \times X + b$, where $X$ is an input parameter, $a$ and $b$ are parameters, and $Y$ is an estimate. A data point consists of tuples of input variables and their corresponding, ideally correct estimates, e.g., $(x, y)$ with $x$ being an $X$ value and $y$ being a $Y$ value. The objective of the model optimization is to determine values for the parameters, so that they provide the best possible accuracy for the observed data, i.e., to minimize the mean squared error, $(y - (a \times x + b))^2$, for all data points $(x, y)$.

The main step is to train the parameters of a human-developed model. Note, that since the model formulas are not always linear, linear regression generally cannot be used. Non-parametric regression is also not applicable either since the goal is not to find the parametric model. It would result in a model that is massively over-trained, especially in the light of the aforementioned cold start problem. It would also be too complex to be understandable.

Optimization algorithms can calculate the minimum or maximum of a massively over-determined system of equations and, therefore, optimization algorithms for zero-point calculations should be used. Various groups of optimization algorithms exist for this purpose. We classify them based on which derivative they need: class A requires the second derivative and class B only the first derivative. For example, Gradient descent requires the first derivative and is thus of class B while the Broyden-Fletcher-Goldfarb-Shanno algorithm requires the first and second derivatives and is of class A.

In practice, derivatives of the model function do not always exist, because not all model functions can be differentiated at every argument vector. If no derivations are known, only optimization algorithms for non-differentiable functions can be used to determine constants. We, therefore, define a class C where no derivative is used, e.g., the Downhill Simplex algorithm.

In our implementation, we currently consider the
following five optimization algorithms: Broyden-Fletcher-Goldfarb-Shanno (A) (Fletcher, 1987), Quasi-Newton (A) (Huang, 2017), Gradient descent (B), Downhill simplex (C) (Nelder and Mead, 1965), and Trust-region (C) (Sorensen, 1982). Note that our system is extensible with any number of such algorithms.

### 3.3 Optimizing Model Parameters and Assessing Models

Our approach contains the following steps. Data points are continuously collected, and domain experts develop and add models. Based on the data points, optimization algorithms are used to determine the parameters in the models that best fit the collected data points. To determine the performance of the various models, the error is calculated and based on this error, we determine whether the model can be used for predictions or not. Finally, it is suggested how the models can be improved.

The model function is entered manually in our approach. Then an algorithmic differentiation is used to try to determine the derivatives automatically. If this is successful, the corresponding optimization algorithms of class A and B are available for finding the constants. If not, the approach asks for manual entering of the first and second derivatives of a model. If the model developer does not want to or cannot provide them only optimization algorithms of class C can be used.

All optimization algorithms that can be used for a suggested model, depending on the available derivations, belong to the “pool of available algorithms” and are used for subsequent optimizations of the corresponding model. For example, a model for which the second derivative is defined, the pool consists of optimization algorithms from classes A, B, and C. Each model function is optimized for all applicable data points using all optimization algorithm from the pool.

The set of applicable data points consists of all data points that have corresponding values for all input variables occurring in the model. If a model function requires an input \( x \), and this is not contained in a data point, this data point is not used for optimization. If, on the other hand, input \( x = N/A \), i.e., it was requested but not provided, standard procedures from statistics are used to deal with missing data and the data point is used. In the simplest case, the missing data are replaced with the mean value, no matter what kind values were missing. Other statistical standard methods like regression imputation, substitution by positional dimensions, or maximum likelihood methods can also be used.

Assume an existing model \( M \) is manually improved and saved as \( M' \). If \( M' \) contains at most the input variables of its predecessor \( M \) then existing data points could be used to train the new model as well. This is because these data points include all input variables needed to train \( M \) and are, hence, applicable also for \( M' \). This makes the two models comparable directly. Conversely, if \( M' \) adds a new input variable, it is critical to apply the standard procedures for dealing with missing data in \( M \) and to compare \( M \) and \( M' \) based on existing data points.

If the error functions are convex then it is easy to find the global minimum. If they are not convex, it is potentially difficult to find the global minimum. It is, however, enough to find several local minima and select the best of these. In addition to the application of various optimization methods, various measures should be taken to increase the probability of finding different local minima to choose the best from. These measures include learning parameters to make larger or smaller jumps, the addition of random values to weights, and the use of Momentum Terms to add to past weight adjustments, 0.25, 0.125, etc. All this remains transparent for the model developer.

All optimization procedures contained in the pool of available algorithms are applied several times with different settings, and each suggests parameters. Then it is checked which of these produces the best result, i.e., the smallest errors.

In the first step, the parameters for each given model are calculated with each implemented optimization algorithm and all data as training data. These parameters are potentially over-optimized. For error calculation, we calculate the mean squared error \( mse \), i.e., the distance between \( Y_{calc} \) and \( Y_{real} \) is calculated, squared, and averaged, cf. Equation 1.

\[
\text{mse} = \frac{1}{n} \sum_{i=1}^{n} (Y_{calc}^i - Y_{real}^i)^2.
\] (1)

In the second step, the principles of cross-validation are used. Iteratively, the data points are randomly divided into training and test data sets. Then, the parameters are calculated from the training data, and the corresponding error is calculated from the test data according to equation 1. For each iteration, the calculated errors may differ. Therefore, the errors are averaged out. This is done until the change in the averaged error is smaller than \( \epsilon \) constant or after a fixed number of iterations. The latter is an emergency stop function guaranteeing termination that is used whenever the parameters found are not stable.

The following data is stored for each run: the model, optimization algorithm, the algorithm setting, the parameters, the mean squared error for the train-
ing session, the averaged mean squared error for the testing data sets, and a list of the data points used.

3.4 Improving Models

The calculated models might have a large number of input variables, which may be very time-consuming to retrieve and to enter. It is possible that 30 out of 50 input variables in a model function can explain 95% of the information gain. A human expert can then improve the model so that in the future, only the relevant variables need to be assessed and entered. For example, if two variables are linearly highly dependent on each other, then one of the two can be omitted. Our approach supports this manual model improvement.

Principal Component Analysis (PCA) forms the basis for calculating the information gain of input variables. After each parameter optimization, it can give hints on model improvements. Therefore, Eigenvalues and Eigenvectors are calculated. The Eigenvalues indicate how significant the respective Eigenvectors are for the explanatory power. The Eigenvalues corresponding to a component divided by the sum of all Eigenvalues gives the information gain of the respective component. First, the principal component is calculated. After that, it is calculated how good the principle component replacement is. Finally, the input variables are ranked based on their explanatory power.

\[
a_1 \times X_1^2 + a_2 \times X_2 + a_3
\]  

(2)

Based on the indication for the explanatory power of an input variable, the model developer is usually interested in how the model behaves if input variables or even complete terms are omitted. To avoid creating a new model for each such constellation, individual terms and parameters can be “switched off” semi-automatically. For example, the parametric model given in Equation 2 can be entered with optional parts. Three different alternatives are shown in the Equations 3a–3c.

\[
a_1 \times [X_1^2] + [a_2 \times X_2] + a_3
\]  

(3a)

\[
a_1 \times X_1^2 + [a_2 \times X_2 + a_3]
\]  

(3b)

\[
[a_1 \times X_1^2] + [a_2 \times X_2 + a_3]
\]  

(3c)

Brackets \([\ ]\) are used to indicate optional parts. The terms between an opening and a closing square bracket are included in the model. However, a model developer can optionally skip them during the parameter optimization steps. The errors of the different options of the model formula can always be compared with each other. It would also be feasible to work with neutral and inverse elements, but this is not that flexible, since the expression of the inverse element is often complicated, and the optimization time grows because more variables need to be evaluated.

3.5 Using Models

With a trained parameterized model, it is in principle always possible to make predictions. Depending on their accuracy, they can be a complement to or a substitute for human based, subjective estimations. To do that, a data point for which the estimate is not known can be entered. Then a predicted value is calculated, always using the best constants of the best applicable model.

Using the results of a trained model only works reliably if the designed parametric model is suitable in the first place and if the data points collected are suitable. Suitability of the model means that it reflects the actual relation between input and estimation variables and only disregards the actual parameter value. Suitability of the data points means that they are representative samples of the real-world development processes. Then, over time, the errors tend to decrease with new data points.

The model designer is responsible for assessing the suitability of the model. Regarding the suitability of a data point, statistical methods can support the model designer: Assuming that the input variable values are generally normally distributed and are within a scattered range around the expected value, values further than, e.g., two times the standard deviation outside the expectation could be regarded as outliers. If the variable values are not normally distributed, other criteria must be defined in order to distinguish between outliers and non-outliers. For run time efficiency, a standard data-mining algorithm should be used in practice, e.g., the Local Outlier Factor Algorithm (Breunig et al., 2000), which does not depend on the normal distribution.

4 EVALUATION

We experimentally investigate GC 1 and GC 2, whether the parametric models can be parameterized automatically based on data, become more and more accurate, and that the errors converge.

4.1 Implementation

Therefore, we implemented the approach as a web application with a simple CRUD-interface in Laravel
and PHP to enter data points and models. These were stored in a MySQL database and all optimizations were implemented using R, specifically the optim and nlm functions. To make the application portable, we packaged it as a Docker container.

4.2 Training Data and Models for Evaluation

We use synthetic data and artificial models to evaluate, since the focus is on evaluating whether the approach fulfills the goal criteria, not to create realistic and useful models per se. In the evaluations, we use three data generators: \( d_1(x) = A \times x + B \), \( d_2(x) = A \times x^3 + B \), and \( d_3(x) = A \times \sin x + B \). The data points in these data sets are generated by placing random seed constants into a model so that the real fuzziness is simulated, cf. Figure 1 for the first data set.

We also use three parametric estimation models: 

1. \( f_1(x) = a \times x + b \),
2. \( f_2(x) = a \times x^3 + b \), and
3. \( f_3(x) = a \times \sin x + b \).

The approach should fit \( a \) to \( A \) and \( a \) to \( A \) when optimizing the first (second, third) model with the first (second, third) data set.

4.3 Experiments and Evaluation Method

4.3.1 One Model and Several Data Sets

We first investigate a single model, \( f_1(x) \) and three different data sets generated from \( d_1, d_2, \) and \( d_3 \). We generate data randomly for each data set, optimize the parameters \( a, b \) of the model \( f_1(x) \). The model optimization should fit the model to minimize the error. As discussed previously, we compute two errors, \( e_1 \) that uses all data points as the training set and \( e_2 \) that uses a cross-validation approach and divides the data set into training and test sets. Both are calculated as the mean squared error. For each new data point randomly generated, we assess these errors.

4.3.2 Several Models and One Data Set

We also investigate how several models can compete in accuracy when fit to a single data set. Therefore, we use three models: \( f_1, f_2(x), \) and \( f_3 \) and a single data set generated from \( d_1 \). We use the same method to generate the data set and the same error calculations. We now investigate which of the three models fits best to the given data set.

4.3.3 Performance of the Optimization Algorithms

Finally, we also want to compare how well different optimization algorithms perform. We assess the \( R \) functions \( \text{optim}(M_{\text{optim}}) \) and \( \text{nlm}(M_{\text{nlm}}) \) by repeating the previous experiment with these two parameter optimization algorithms.

4.3.4 Error Calculation

Finally, we evaluated which of the two error calculation approaches \( e_1 \) or \( e_2 \) is best suited for evaluating a model with estimated parameters. Error \( e_1 \) assumes that all data is training data, while \( e_2 \) divides similar procedures into training and validation data using cross-validation.

4.4 Results

4.4.1 One Model and Several Data Sets

The first data set \( d_1 \) corresponds well to the model \( f_1 \). This is the ideal case. As expected, the errors converge to near zero, cf. Figure 2.

The second data set \( d_2 \) is based on a cubic function, for which a linear model is not a good fit. While the errors converge, they remain quite large, cf. Figure 3.

The third data set \( d_3 \) is based on a sine function. The errors converge in this case as well but, are again relatively high, compared to the first data set, cf. Figure 4. The reason for this is that a horizontal straight line can be drawn through the data points, against which optimization is performed. The data points are relatively close to each other and the horizontal straight line.

4.4.2 Several Models and One Data Set

Again, the picture is clear: Although all errors become constant at some point, only the error of the linear model formula \( f_1 \) matching the data points \( d_1 \) is small. The error is a measure of the accuracy of the model. Figure 5 clearly shows the difference between a model that fits very well to the data set \( d_1 \) and a model that fits badly to it.

4.4.3 Performance of the Optimization Algorithms

It shows that there is not a single best algorithm: while method \( M_{\text{optim}} \) is particularly suitable for \( f_2 \), cf. Figure 6), method \( M_{\text{nlm}} \) is more suitable for \( f_3 \), cf. Figure 7. \( M_{\text{optim}} \) and \( M_{\text{nlm}} \) are similarly well suited.
The first set of datapoints is based on \( a \times x + b \) and therefore corresponds very well to the trivial case \( f_1(x) = a \times x + b \). Data points can be collected while correlations between inputs can be detected by performing PCA. Since the models are designed by a human project developer, it makes sense to provide different optimization algorithms and to use them in parallel.

### 4.4.4 Error Calculation

The error calculation uses two approaches \( e_1 \) or \( e_2 \). Figures 2–4 show that their differences are marginal and converge to zero. However, \( e_1 \) is potentially overtrained and therefore less trustworthy than \( e_2 \).

### 4.5 Discussion

The evaluation focused on GC 1 and GC 2 and showed that the model parameters could be trained from data, are suitable to make predictions, and that the accuracy improves and converges with the number of training data points.

The remaining goal criteria are also fulfilled by the design and do not need to be evaluated experimentally. Since the models are designed by a human domain expert and only the parameters are optimized, it can be assumed that they are understandable and trusted. The parameters that the optimization algorithm determines can easily be validated against the data set; hence, GC 5 is fulfilled. Since outliers can be detected, the error is provided, and the model’s fit against the data sets can easily be inspected, we consider GC 4 fulfilled as well. The evaluation used artificial models and synthetic data, with a sample size of 500 for each data generator, not tied to the software estimation domain. While repeating the experiments with real models and data is a matter of future work, this evaluation method at least shows that the approach applies to models of different types and domains, so GC 6 is also fulfilled.

GC 3 is fulfilled by the use of PCA to determine how much explanatory power each parameter has, so the PCA can suggest that some data points are not needed for the model. It can also be used to determine if optional parts of the model should be included or not based on their explanatory power. This information can be provided to the model developer as suggestions.

We can now answer our research questions: Continuous model improvement can be realized, and new models can be easily tested and implemented, using the approach based on optimization algorithms (RQ1). Data points can be collected while correlations between inputs can be detected by performing...
Figure 5: Comparison of the models $f_1$, $f_2$, and $f_3$ on the data set generated from $d_1(x) = A \times x + B$.

Figure 7: Comparison of two optimization algorithms, $M_{\text{optim}}$ (lower) and $M_{\text{nlm}}$ (upper) for the model $f_3$ on the data set $d_1$.

Figure 6: Comparison of two optimization algorithms $M_{\text{optim}}$ (lower) and $M_{\text{nlm}}$ (upper) for the model $f_2$ on the data set $d_1$.

Figure 8: Comparison of two optimization algorithms, $M_{\text{optim}}$ and $M_{\text{nlm}}$ (overlapping) for the model $f_1$ on the data set $d_1$.

5 CONCLUDING REMARKS AND FUTURE WORK

We suggested an approach to estimation model design and automated optimization that allows for model comparison and improvement based on collected data sets. Our approach simplifies model optimization and selection. It contributes to a convergence of existing estimation models to meet contemporary software technology practices and to a possible selection of the most appropriate ones.

We evaluated the design and implementation of our approach and showed that it fulfilled all the goal criteria that we formulated. The model accuracy is improved through optimization, and the models are understandable by human domain experts. We also find that principal component analysis can determine the explanatory power of parameters and parts of models can suggest changes to the model developer.

We only evaluated our approach on synthetic data, so future work is to apply it to data from real-world software projects. We particularly want to use our approach to compare models that we design with established models such as COCOMO II to see how they compare. We are also interested to see how the approach will perform, both concerning accuracy and performance on larger data sets. Further yet, we relied on a small set of optimization algorithms, and we plan to incorporate more algorithms, e.g., genetic algorithms in the future.

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\(^2\)http://www.kks.se
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