Faster Discovery of Faster System Configurations with Spectral Learning

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Abstract Despite the huge spread and economical importance of configurable software systems, there is unsatisfactory support in utilizing the full potential of these systems with respect to finding performance-optimal configurations. Prior work on predicting the performance of software configurations suffered from either (a) requiring far too many sample configurations or (b) large variances in their predictions. Both these problems can be avoided using the WHAT spectral learner. WHAT’s innovation is the use of the spectrum (eigenvalues) of the distance matrix between the configurations of a configurable software system, to perform dimensionality reduction. Within that reduced configuration space, many closely associated configurations can be studied by executing only a few sample configurations. For the subject systems studied here, a few dozen samples yield accurate and stable predictors—less than 10% prediction error, with a standard deviation of less than 2%. When compared to the state of the art, WHAT (a) requires 2 to 10 times fewer samples to achieve similar prediction accuracies, and (b) its predictions are more stable (i.e., have lower standard deviation). Furthermore, we demonstrate that predictive models generated by WHAT can be used by optimizers to discover system configurations that closely approach the optimal performance.

Keywords Performance Prediction · Spectral Learning · Decision Trees · Search-Based Software Engineering · Sampling.

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1 Introduction

Most software systems today are configurable. Despite the undeniable benefits of configurability, large configuration spaces challenge developers, maintainers, and users. In the face of hundreds of configuration options, it is difficult to keep track of the effects of individual configuration options and their mutual interactions. So, predicting the performance of individual system configurations or determining the optimal configuration is often more guess work than engineering. In their recent paper, Xu et al. documented the difficulties developers face with understanding the configuration spaces of their systems [40]. As a result, developers tend to ignore over 5/6ths of the configuration options, which leaves considerable optimization potential untapped and induces major economic cost [40].

Addressing the challenge of performance prediction and optimization in the face of large configuration spaces, researchers have developed a number of approaches that rely on sampling and machine learning [15, 29, 35]. While gaining some ground, state-of-the-art approaches face two problems: (a) they require far too many sample configurations for learning or (b) they are prone to large variances in their predictions. For example, prior work on predicting performance scores using regression trees had to compile and execute hundreds to thousands of specific system configurations [15]. A more balanced approach by Siegmund et al. is able to learn predictors for configurable systems [35] with low mean errors, but with large variances of prediction accuracy (e.g., in half of the results, the performance predictions for the Apache Web server were up to 50% wrong). Guo et al. [15] also proposed an incremental method to build a predictor model, which uses incremental random samples with steps equal to the number of configuration options (features) of the system. This approach also suffered from unstable predictions (e.g., predictions had a mean error of up to 22%, with a standard deviation of up 46%). Finally, Sarkar et al. [29] proposed a projective-learning approach (using fewer measurements than Guo et al. and Siegmund et al.) to quickly compute the number of sample configurations for learning a stable predictor. However, as we will discuss, after making that prediction, the total number of samples required for learning the predictor is comparatively high (up to hundreds of samples).

The problems of large sample sets and large variances in prediction can be avoided using the WHAT spectral learner, which is our main contribution. WHAT’s innovation is the use of the spectrum (eigenvalues) of the distance matrix between the configurations of a configurable system, to perform dimensionality reduction. Within that reduced configuration space, many closely associated configurations can be studied by measuring only a few samples. In a number of experiments, we compared WHAT against the state-of-the-art approaches of Siegmund et al. [35], Guo et al. [15], and Sarkar et al. [29] by means of six real-world configurable systems: Berkeley DB, the Apache Web server, SQLite, the LLVM compiler, and the x264 video encoder. We found that WHAT performs as well or better than prior approaches, while requiring far fewer samples (just a few dozen). This is significant and most surprising, since some of the systems explored here have up to millions of possible configurations.
Overall, we make the following contributions:

- We present a novel sampling and learning approach for predicting the performance of software configurations in the face of large configuration spaces. The approach is based on a spectral learner that uses an approximation to the first principal component of the configuration space to recursively cluster it, relying only on a few points as representatives of each cluster.
- We demonstrate the practicality and generality of our approach by conducting experiments on six real-world configurable software systems (see Figure 1). The results show that our approach is more accurate (lower mean error) and more stable (lower standard deviation) than state-of-the-art approaches. A key finding is the utility of the principal component of a configuration space to find informative samples from a large configuration space.

All materials required for reproducing this work are available at https://goo.gl/689Dve.

2 Background & Related Work

A configurable software system has a set $X$ of Boolean configuration options, also referred to as features or independent variables in our setting. We denote the number of features of system $S$ as $n$. The configuration space of $S$ can be represented by a Boolean space $\mathbb{Z}_2^n$, which is denoted by $F$. All valid configurations of $S$ belong to a set $V$, which is represented by vectors $C_i$ (with $1 \leq i \leq |V|$) in $\mathbb{Z}_2^n$. Each element of a configuration represents a feature, which can either be True or False, based on whether the feature is selected or not. Each valid instance of a vector (i.e., a configuration) has a corresponding performance score associated to it.

The literature offers two approaches to performance prediction of software configurations: a maximal sampling and a minimal sampling approach: With maximal sampling, we compile all possible configurations and record the associated performance scores. Maximal sampling can be impractically slow. For example, the performance data used in our experiments required 26 days of CPU time for measuring (and much longer, if we also count the time required for compiling the code prior to execution). Other researchers have commented that, in real world scenarios, the cost of acquiring the optimal configuration is overly expensive and time consuming [39].

If collecting performance scores of all configurations is impractical, minimal sampling can be used to intelligently select and execute just enough configurations (i.e., samples) to build a predictive model. For example, Zhang et al. [42] approximate the configuration space as a Fourier series, after which they can derive an expression showing how many configurations must be studied to build predictive models with a given error. While a theoretically satisfying result, that approach still needs thousands to hundreds of thousands of executions of sample configurations.

Another set of approaches are the four "additive" minimal sampling methods of Siegmund et al. [35]. Their first method, called feature-wise sampling (FW), is their

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1 In this paper, we concentrate on Boolean options, as they make up the majority of all options; see Siegmund et al., for how to incorporate numeric options [34].
basic method. To explain FW, we note that, from a configurable software system, it is theoretically possible to enumerate many or all of the valid configurations. Since each configuration \( C_i \) is a vector of \( n \) Booleans, it is possible to use this information to isolate examples of how much each feature individually contributes to the total run time:

1. Find a pair of configurations \( C_1 \) and \( C_2 \), where \( C_2 \) uses exactly the same features as \( C_1 \), plus one extra feature \( f_i \).
2. Set the run time \( \Pi(f_i) \) for feature \( f_i \) to be the difference in the performance scores between \( C_2 \) and \( C_1 \).
3. The run time for a new configuration \( C_i \) (with \( 1 \leq i \leq |V| \)) that has not been sampled before is then the sum of the run time of its features, as determined before:

\[
\Pi(C_i) = \sum_{f_j \in C_i} \Pi(f_j)
\]

When many pairs, such as \( C_1, C_2 \), satisfy the criteria of point 1, Siegmund et al. used the pair that covers the smallest number of features. Their minimal sampling method, FW, compiles and executes only these smallest \( C_1 \) and \( C_2 \) configurations. Siegmund et al. also offers three extensions to the basic method, which are based on sampling not just the smallest pairs, but also additional configurations covering certain kinds of interactions between features. All the following minimal sampling policies compile and execute valid configurations selected via one of three heuristics:

**PW (pair-wise):** For each pair of features, try to find a configuration that contains the pair and has a minimal number of features selected.

**HO (higher-order):** Select extra configurations, in which three features, \( f_1, f_2, f_3 \), are selected if two of the following pair-wise interactions exist: \((f_1, f_2)\) and \((f_2, f_3)\) and \((f_1, f_3)\).

**HS (hot-spot):** Select extra configurations that contain features that are frequently interacting with other features.

Guo et al. \[15\] proposed a progressive random sampling approach, which samples the configuration space in steps of the number of features of the software system in question. They used the sampled configurations to train a regression tree, which is then used to predict the performance scores of other system configurations. The termination criterion of this approach is based on a heuristic, similar to the PW heuristics of Siegmund et al.

Sarkar et al. \[29\] proposed a cost model for predicting the effort (or cost) required to generate an accurate predictive model. The user can use this model to decide whether to go ahead and build the predictive model. This method randomly samples configurations and uses a heuristic based on feature frequencies as termination criterion. The samples are then used to train a regression tree; the accuracy of the model is measured by using a test set (where the size of the training set is equal to size of the test set). One of four projective functions (e.g., exponential) is selected based on how

\[\text{Though, in practice, this can be very difficult. For example, in models like the Linux Kernel such an enumeration is practically impossible.}\]

\[\text{[29]}\]
correlated they are to accuracy measures. The projective function is used to approximate the accuracy-measure curve, and the elbow point of the curve is then used as the optimal sample size. Once the optimal size is known, Sarkar et al. uses the approach of Guo et al. to build the actual prediction model.

The advantage of these previous approaches is that, unlike the results of Zhang et al., they require only dozens to hundreds of samples. Also, like our approach, they do not require to enumerate all configurations, which is important for highly configurable software systems. That said, as shown by our experiments (see Section 4), these approaches produce estimates with larger mean errors and partially larger variances than our approach. While sometimes the approach by Sarkar et al. results in models with (slightly) lower mean error rates, it still requires a considerably larger number of samples (up to hundreds), while WHAT requires only few dozen.

3 Approach

3.1 Spectral Learning

The minimal sampling method we propose here is based on a spectral-learning algorithm that explores the spectrum (eigenvalues) of the distance matrix between configurations in the configuration space. In theory, such spectral learners are an appropriate method to handle noisy, redundant, and tightly inter-connected variables, for the following reasons: When data sets have many irrelevancies or closely associated data parameters $d$, then only a few eigenvectors $e, e \ll d$ are required to characterize the data. In this reduced space:

- Multiple inter-connected variables $i, j, k \subseteq d$ can be represented by a single eigenvector;
- Noisy variables from $d$ are ignored, because they do not contribute to the signal in the data;
- Variables become (approximately) parallel lines in $e$ space. For redundancies $i, j \in d$, we can ignore $j$ since effects that change over $j$ also change in the same way over $i$;

That is, in theory, samples of configurations drawn via an eigenspace sampling method would not get confused by noisy, redundant, or tightly inter-connected variables. Accordingly, we expect predictions built from that sample to have lower mean errors and lower variances on that error.

Spectral methods have been used before for a variety of data mining applications [20]. Algorithms, such as PDDP [4], use spectral methods, such as principle component analysis (PCA), to recursively divide data into smaller regions. Software-analytics researchers use spectral methods (again, PCA) as a pre-processor prior to data mining to reduce noise in software-related data sets [37]. However, to the best of our knowledge, spectral methods have not been used before as a basis of a minimal sampling method.

WHAT is somewhat different from other spectral learners explored in, for instance, image processing applications [31]. Work on image processing does not aim
at defining a minimal sampling policy to predict performance scores. Also, a standard spectral method requires an $O(N^2)$ matrix multiplication to compute the components of PCA \[18\]. Worse, in the case of hierarchical division methods, such as PDDP, the polynomial-time inference must be repeated at every level of the hierarchy. Competitive results can be achieved using an $O(2N)$ analysis that we have developed previously \[24\], which is based on a heuristic proposed by Faloutsos and Lin \[19\] (which Platt has shown computes a Nyström approximation to the first component of PCA \[27\]).

**WHAT** receives $N$ (with $1 \leq |N| \leq |V|$) valid configurations ($C$), $N_1, N_2, \ldots$, as input and then:

1. Picks any point $N_i$ ($1 \leq i \leq |N|$) at random;
2. Finds the point $\text{West} \in N$ that is furthest away from $N_i$;
3. Finds the point $\text{East} \in N$ that is furthest from $\text{West}$.

The line joining $\text{East}$ and $\text{West}$ is our approximation for the first principal component. Using the distance calculation shown in Equation 2, we define $\delta$ to be the distance between $\text{East} (x)$ and $\text{West} (y)$. **WHAT** uses this distance ($\delta$) to divide all the configurations as follows: The value $x_i$ is the projection of $N_i$ on the line running from $\text{East}$ to $\text{West}$. We divide the examples based on the median value of the projection of $x_i$. Now, we have two clusters of data divided based on the projection values of $N_i$ on the line joining $\text{East}$ and $\text{West}$. This process is applied recursively on these clusters until a predefined stopping condition. In our study, the recursive splitting of the $N_i$’s stops when a sub-region contains less than $\sqrt{|N|}$ examples.

\[
dist(x,y) = \begin{cases} 
\sqrt{\sum_i (x_i - y_i)^2} & \text{if } x_i \text{ and } y_i \text{ is numeric} \\
0 & \text{if } x_i = y_i \\
1 & \text{otherwise}
\end{cases}
\]  

We explore this approach for three reasons:

– **It is very fast**: This process requires only $2|n|$ distance comparisons per level of recursion, which is far less than the $O(N^2)$ required by PCA \[8\] or other algorithms such as K-Means \[16\].

– **It is not domain-specific**: Unlike traditional PCA, our approach is general in that it does not assume that all the variables are numeric. As shown in Equation 2\[4\], we can approximate distances for both numeric and non-numeric data (e.g., Boolean).

– **It reduces the dimensionality problem**: This technique explores the underlying dimension (first principal component) without getting confused by noisy, related, and highly associated variables.

\[3\] The projection of $N_i$ can be calculated in the following way:

\[
a = \text{dist}(\text{East}, N_i); b = \text{dist}(\text{West}, N_i); x_i = \sqrt{\frac{a^2 + b^2 - 2ab\sin^2\theta}{2}}.
\]

\[4\] In our study, $\text{dist}$ accepts pair of configuration ($C$) and returns the distance between them. If $x_i$ and $y_i \in \mathbb{R}^n$, then the distance function would be same as the standard Euclidean distance.
3.2 Spectral Sampling

When the above clustering method terminates, our sampling policy (which we call $S_1$) is then applied:

- **Random sampling ($S_1$):** compile and execute one configuration, picked at random, from each leaf cluster;

We use this sampling policy, because (as we will show later) it performs better than:

- **East-West sampling ($S_2$):** compile and execute the East and West poles of the leaf clusters;
- **Exemplar sampling ($S_3$):** compile and execute all items in all leaves and return the one with lowest performance score.

Note that $S_3$ is *not a minimal* sampling policy (since it executes all configurations). We use it here as one baseline against which we can compare the other, more minimal, sampling policies. In the results that follow, we also compare our sampling methods against another baseline using information gathered after executing all configurations.

3.3 Regression-Tree Learning

After collecting the data using one of the sampling policies ($S_1$, $S_2$, or $S_3$), as described in Section 3.2, we use a CART regression-tree learner [5] to build a performance predictor. Regression-tree learners seek the attribute-range split that most increases our ability to make accurate predictions. CART explores splits that divide $N$ samples into two sets $A$ and $B$, where each set has a standard deviation on the target variable of $\sigma_1$ and $\sigma_2$. CART finds the “best” split defined as the split that minimizes

$$\frac{A}{N} \sigma_1 + \frac{B}{N} \sigma_2$$

Using this best split, CART divides the data recursively.

In summary, **WHAT** combines:

- The FASTMAP method of Faloutsos and Lin [10], which rather than $N^2$ comparisons only performs $2N$ where $N$ is the number of configurations in the configuration space;
- A spectral-learning algorithm initially inspired by Boley’s PDDP system [4], which we modify by replacing PCA with FASTMAP (called “WHERE” in prior work [24]);
- The sampling policy that explores the leaf clusters found by this recursive division;
- The CART regression-tree learner that converts the data from the samples collected by sampling policy into a run-time prediction model [5].

That is,

$$\text{WHERE} = \text{PDDP} - \text{PCA} + \text{FASTMAP}$$

$$\text{WHAT} = \text{WHERE} + \{S_1, S_2, S_3\} + \text{CART}$$

This unique combination of methods has not been previously explored in the software-engineering literature.
4 Experiments

4.1 Research Questions

We formulate our research questions in terms of the challenges of exploring large complex configuration spaces. As our approach explores the spectral space, our hypothesis is that only a small number of samples is required to explore the whole space. However, a prediction model built from a very small sample of the configuration space might be very inaccurate and unstable, that is, it may exhibit very large mean prediction errors and variances on the prediction error.

Also, if we learn models from small regions of the training data, it is possible that a learner will miss trends in the data between the sample points. Such trends are useful when building optimizers (i.e., systems that receives one configuration as input and propose an alternate configuration that has, for instance, a better performance). Such optimizers might need to evaluate hundreds to millions of alternate configurations. To speed up that process, optimizers can use a surrogate model\(^5\) that mimics the outputs of a system of interest, while being computationally cheap(er) to evaluate \(^{23}\). For example, when optimizing performance scores, we might ask a CART for a performance prediction (rather than compile and execute the corresponding configuration). Note that such surrogate-based reasoning critically depends on how well the surrogate can guide optimization.

Therefore, to assess feasibility of our sampling policies, we must consider:

- Performance scores generated from our minimal sampling policy;
- The variance of the error rates when comparing predicted performance scores with actual ones;
- The optimization support offered by the performance predictor (i.e., can the model work in tandem with other off-the-shelf optimizers to generate useful solutions).

The above considerations lead to four research questions:

**RQ1:** Can WHAT generate good predictions after examining only a small number of configurations?

Here, by “good” we mean that the predictions made by models that were trained using sampling with WHAT are as accurate, or more accurate, as predictions generated from models supplied with more samples.

**RQ2:** Do less data used in building the predictions models cause larger variances in the predicted performance scores?

**RQ3:** Can “good” surrogate models (to be used in optimizers) be built from minimal samples?

Note that RQ2 and RQ3 are of particular concern with our approach, since our goal is to sample as little as possible from the configuration space.

**RQ4:** How good is WHAT compared to the state of the art of learning performance predictors from configurable software systems?

To answer RQ4, we will compare WHAT against approaches presented by Siegmund et al. \(^{35}\), Guo et al. \(^{15}\), and Sarkar et al. \(^{29}\).

\(^5\) Also known as response surface methods, meta models, or emulators.
Table 1 Subject systems used in the experiments.

| System               | #LOC  | #Features | #Configurations |
|----------------------|-------|-----------|-----------------|
| Berkeley DB C Edition (BDBC) | 219,811 | 18        | 2,560           |
| Berkeley DB Java Edition (BDBJ) | 42,596  | 32        | 400             |
| Apache               | 230,277 | 9         | 192             |
| SQLite               | 312,625 | 39        | 3,932,160       |
| LLVM                 | 47,549  | 11        | 1,024           |
| x264                 | 45,743  | 16        | 1,152           |

4.2 Subject Systems

The configurable systems we used in our experiments are described in Table 1. Note, with “predicting performance”, we mean predicting performance scores of the subject systems while executing test suites provided by the developers or the community, as described in Table 1. To compare the predictions of our and prior approaches with actual performance measures, we use data sets that have been obtained by measuring nearly all configurations. We say nearly all configurations, for the following reasoning: For all except one of our subject systems, the total number of valid configurations was tractable (192 to 2560). However, SQLite has 3,932,160 possible configurations, which is an impractically large number of configurations to test whether our predictions are accurate and stable. Hence, for SQLite, we use the 4500 samples for

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*http://openscience.us/repo/performance-predict/cpm.html*
testing prediction accuracy and stability, which we could collect in one day of CPU time. Taking this into account, we will pay particular attention to the variance of the SQLite results.

4.3 Experimental Rig

RQ1 and RQ2 require the construction and assessment of numerous runtime predictors from small samples of the data. The following rig implements that construction process.

For each configurable software system, we built a table of data, one row per valid configuration. We then ran all configurations of all software systems and recorded the performance scores (i.e., that are invoked by a benchmark). The exception is SQLite for which we measured only the configurations needed to detect interactions and additionally 100 random configurations. To this table, we added a column showing the performance score obtained from the actual measurements for each configuration.

Note that the following procedure ensures that we never test any prediction model on the data that we used to learn this model. Next, we repeated the following procedure 20 times (the figure of 20 repetitions was selected using the Central Limit Theorem): For each subject system in \{BDBC, BDBJ, Apache, SQLite, LLVM, x264\}

- Randomize the order of the rows in their table of data;
- For \(X\) in \{10, 20, 30, ... , 90\};
  - Let Train be the first \(X\) % of the data
  - Let Test be the rest of the data;
  - Pass Train to WHAT to select sample configurations;
  - Determine the performance scores associated with these configurations. This corresponds to a table lookup, but would entail compiling and executing a system configuration in a practical setting.
  - Using the Train data and their performance scores, build a performance predictor using CART.
  - Using the Test data, assess the accuracy of the predictor using the error measure of Equation 3 (see below).

The validity of the predictors built by CART is verified on testing data. For each test item, we determine how long it actually takes to run the corresponding system configuration and compare the actual measured performance to the prediction from CART. The resulting prediction error is then computed using:

\[
error = \frac{|predicted - actual|}{actual} \cdot 100
\]  

(Aside: It is reasonable to ask why this metrics and not some of the others proposed in the literature (e.g sum absolute residuals). In short, our results are stable across a range of different metrics. For e.g., the results of this paper have been repeated using sum of absolute residuals and, in those other results, we seen the same ranking of methods; see \[http://tiny.cc/sumAS\].

RQ2 requires testing the standard deviation of the prediction error rate. To support that test, we:
– Determine the $X$-th point in the above experiments, where all predictions stop improving (elbow point);
– Measure the standard deviation of the error at this point, across our 20 repeats.

As shown in Figure 1 all our results plateaued after studying $X = 40\%$ of the valid configurations. Hence to answer RQ2, we will compare all 20 predictions at $X = 40\%$.

RQ3 uses the learned regression tree as a surrogate model within an optimizer;
– Take $X = 40\%$ of the configurations;
– Apply WHAT to build a CART model using some minimal sample taken from that 40%;
– Use that CART model within some standard optimizer while searching for configurations with least runtime;
– Compare the faster configurations found in this manner with the fastest configuration known for that system.

This last item requires access to a ground truth of performance scores for a large number of configurations. For this experiment, we have access to that ground truth (since we have access to all system configurations, except for SQLite). Note that such a ground truth would not be needed when practitioners choose to use WHAT in their own work (it is only for our empirical investigation).

For the sake of completeness, we explored a range of optimizers seen in the literature: DE [36], NSGA-II [6], and our own GALE [21, 43] system. Normally, it would be reasonable to ask why we used those three, and not the hundreds of other optimizers described in the literature [12, 17]. However, as shown below, all these optimizers in this domain exhibited very similar behavior (all found configurations close to the best case performance). Hence, the specific choice of optimizer is not a critical variable in our analysis.

5 Results

5.1 RQ1

Can WHAT generate good predictions after examining only a small number of configurations?

Figure 1 shows the mean errors of the predictors learned after taking $X\%$ of the configurations, then asking WHAT and some sampling method ($S_1$, $S_2$, and $S_3$) to (a) find what configurations to measure; then (b) asking CART to build a predictor using these measurements. The horizontal axis of the plots shows what $X\%$ of the configurations are studied; the vertical axis shows the mean relative error ($\mu$) from Equation 5. In this figure:

7 Just to clarify one frequently asked question about this work, we note that our rig “studies” 40% of the data. We do not mean that our predictive models require accessing the performance scores from the 40% of the data. Rather, by “study” we mean reflect on a sample of configurations to determine what minimal subset of that sample deserves to be compiled and executed.
Fig. 1 Errors of the predictions made by WHAT with four different sampling policies. Note that, on the y-axis, lower errors are better.

- The ×—× lines in Figure 1 show a baseline result where data from the performance scores of 100% of configurations were used by CART to build a runtime predictor.
The other lines show the results using the sampling methods defined in Section 3.2. Note that these sampling methods used runtime data only from a subset of 100% of the performance scores seen in configurations from 0 to X%.

In Figure 1, lower y-axis values are better since this means lower prediction errors. Overall, we find that:

- Some of the subject systems exhibit large variances in their error rate, below \(X = 40\%\) (e.g., BDBC and BDBJ).
- Above \(X = 40\%\), there is little effect on the overall change of the sampling methods.
- Mostly, \(S_3\) shows the highest overall error, so that it cannot be recommended.
- Always, the \(\times - \times\) baseline shows the lowest errors, which is to be expected since predictors built on the baseline have access to all data.
- We see a trend that the error of \(S_1\) and \(S_2\) are within 5% of the baseline results. Hence, we can recommend these two minimal sampling methods.

Figure 2 provides information about which of \(S_1\) or \(S_2\) we should recommend. This figure displays data taken from the \(X = 40\%\) point of Figure 1 and displays how many performance scores of configurations are needed by our sampling methods (while reflecting on the configurations seen in the range \(0 \leq X \leq 40\)). Note that:

- \(S_3\) needs up to thousands of performance scores points, so it cannot be recommended as minimal-sampling policy;
- \(S_2\) needs twice as much performance scores as \(S_1\) (\(S_2\) uses two samples per leaf cluster while \(S_1\) uses only one).
- \(S_1\) needs performance scores only for a few dozen (or less) configurations to generate the predictions with the lower errors seen in Figure 1.

Combining the results of Figure 1 and Figure 2, we conclude that:

\(S_1\) is our preferred spectral sampling method. Furthermore, the answer to RQ1 is “yes”, because applying WHAT, we can (a) generate runtime predictors using just a few dozens of sample performance scores; and (b) these predictions have error rates within 5% of the error rates seen if predictors are built from information about all performance scores.

5.2 RQ2

*Do less data used in building prediction models cause larger variances in the predicted values?*

Two competing effects can cause increased or decreased variances in performance predictions. In our study, we report standard deviation (\(\sigma\)) as a measure of variances in the performance predictions. The less we sample the configuration space, the less we constrain model generation in that space. Hence, one effect that can be expected is that models learned from too few samples exhibit large variances. But, a compensating effect can be introduced by sampling from the spectral space since that space contains fewer confusing or correlated variables than the raw configuration space.
Comparing evaluations of different sampling policies. We see that the number of configurations evaluated for $S_2$ is twice as high as $S_1$, as it selects 2 points from each cluster, whereas $S_1$ selects only 1 point.

Figure 3 shows that after some initial fluctuations, after seeing $X = 40\%$ of the configurations, the variances in prediction errors reduce to nearly zero, which is similar to the results in Figure 1.

Based on the results of Figure 3, we answer RQ2 with “no”: Selecting a small number of samples does not necessarily increase variance (at least to say, not in this domain).

5.3 RQ3

Can “good” surrogate models (to be used in optimizers) be built from minimal samples?
The results of answering RQ1 and RQ2 suggest to use WHAT (with $S_1$) to build runtime predictors from a small sample of data. RQ3 asks if that predictor can be used by an optimizer to infer what other configurations correspond to system configurations with fast performance scores. To answer this question, we ran a random set of 100 configurations, 20 times, and related that baseline to three optimizers (GALE [21], DE [36] and NSGA-II [6]) using their default parameters.
When these three optimizers mutated existing configurations to suggest new ones, these mutations were checked for validity. Any mutants that violated the system’s constraints (e.g., a feature excluding another feature) were rejected and the survivors were “evaluated” by asking the CART surrogate model. These evaluations either rejected the mutant or used it in generation $i + 1$, as the basis for a search for more, possibly better mutants.

Figure 4 shows the configurations found by the three optimizers projected onto the ground truth of the performance scores of nearly all configurations (see Section 4.2). Again note that, while we use that ground truth for the validation of these results, our optimizers used only a small part of that ground-truth data in their search for the fastest configurations (see the WHAT + S1 results of Figure 2).

The important information of Figure 4 is that all the optimized configurations fall within 1% of the fastest configuration according to the ground truth (see all the left-hand-side dots on each plot). Table 2 compares the performance of the optimizers used in this study. Note that the performances are nearly identical, which leads to the following conclusions:

Based on the results of figure 4 answer to RQ3 is “yes”: For optimizing performance scores, we can use surrogates built from few runtime samples. The choice of the optimizer does not critically effect this conclusion.

**Table 2** The table shows how the minimum performance scores as found by the learners GALE, NSGA-II, and DE, vary over 20 repeated runs. Mean values are denoted $\mu$ and IQR denotes the 25th–75th percentile. A low IQR suggests that the surrogate model build by WHAT is stable and can be utilized by off the shelf optimizers to find performance-optimal configurations.

| Searcher | GALE Mean | IQR | DE Mean | IQR | NSGAII Mean | IQR |
|----------|-----------|-----|---------|-----|-------------|-----|
| Apache   | 870       | 0   | 840     | 0   | 840         | 0   |
| BDBC     | 0.363     | 0.004 | 0.359 | 0.002 | 0.354       | 0.005 |
| BDBJ     | 3139      | 70   | 3139    | 70   | 3139        | 70   |
| LLVM     | 202       | 3.98 | 200     | 0   | 200         | 0   |
| SQLite   | 13.1      | 0.241 | 13.1   | 0   | 13.1        | 0.406 |
| X264     | 248       | 3.3  | 244     | 0.003 | 244         | 0.05 |

5.4 RQ4

*How good is WHAT compared to the state of the art of learning performance predictors from configurable software systems?*

We compare WHAT with the three state-of-the-art predictors proposed in the literature [35], [15], [29], as discussed in Section 2. Note that all approaches use
Fig. 4 Solutions found by GALE, NSGA-II, and DE (shown as points) laid against the ground truth (all known configuration performance scores). It can be observed that all the optimizers can find the configuration with lower performance scores.

regression-trees as predictors, except Siegmund’s approach, which uses a regression function derived using linear programming. The results were studied using non-parametric tests, which was also used by Arcuri and Briand at ICSE ’11 (25). For testing statistical significance, we used non-parametric bootstrap test 95% confidence [9] followed by an A12 test to check that any observed differences were not
| Rank | Approach | Mean MRE(µ) | STDev(σ) | #Evaluations |
|------|----------|-------------|----------|--------------|
| **Apache** | | | | |
| 1 | Sarkar | 7.49 | 0.82 | 55 |
| 1 | Guo(PW) | 10.51 | 6.85 | 29 |
| 1 | Siegmund | 10.34 | 11.68 | 29 |
| 1 | WHAT | 10.95 | 2.74 | 16 |
| 1 | Guo(2N) | 13.03 | 15.28 | 18 |
| **BDDBC** | | | | |
| 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 |
| 1 | WHAT | 4.75 | 4.46 | 16 |
| 2 | Sarkar | 5.67 | 6.97 | 48 |
| 2 | Siegmund | 6.98 | 7.13 | 57 |
| **LLVM** | | | | |
| 1 | Guo(PW) | 3.09 | 2.98 | 64 |
| 1 | WHAT | 3.32 | 1.05 | 32 |
| 1 | Sarkar | 3.72 | 0.45 | 62 |
| 1 | Guo(2N) | 4.99 | 5.05 | 22 |
| 2 | Siegmund | 8.50 | 8.28 | 43 |
| **SQLite** | | | | |
| 1 | Sarkar | 3.44 | 0.10 | 925 |
| 2 | WHAT | 5.60 | 0.57 | 64 |
| 3 | Guo(2N) | 8.57 | 7.30 | 78 |
| 3 | Guo(PW) | 8.94 | 6.24 | 566 |
| 4 | Siegmund | 12.83 | 17.0 | 566 |
| **x264** | | | | |
| 1 | Sarkar | 6.64 | 1.04 | 93 |
| 1 | WHAT | 6.93 | 1.67 | 32 |
| 1 | Guo(2N) | 7.18 | 7.07 | 32 |
| 1 | Guo(PW) | 7.72 | 2.33 | 81 |
| 2 | Siegmund | 31.87 | 21.24 | 81 |

**Fig. 5** Mean MRE(µ) seen in 20 repeats. Mean MRE is the prediction error as described in Equation 3 and STDev (σ) is the standard deviation of the MREs found during multiple repeats. Lines with a dot in the middle (e.g. •) show the mean as a round dot within the IQR (and if the IQR is very small, only a round dot will be visible). All the results are sorted by the mean values: a lower mean value of MRE is better than large mean value. The left-hand side column (rank) ranks the various techniques for e.g. when comparing various techniques for Apache, all the techniques have the same rank since their mean values are not statistically different. Rank is computer using Scott-Knott, bootstrap 95% confidence, and the A12 test.

Trivially small effects; i.e. given two lists X and Y, count how often there are larger numbers in the former list (and there are ties, add a half mark): \( a = \forall x \in X, y \in Y \#(x>y) + 0.5 \#(x=y) \) (as per Vargha [38], we say that a “small” effect has \( a < 0.6 \)). Lastly, to generate succinct reports, we use the Scott-Knott test to recursively divide our optimizers. This recursion used A12 and bootstrapping to group together subsets that are (a) not significantly different and are (b) not just a small effect different to
each other. This use of Scott-Knott is endorsed by Mittas and Angelis [25] and by Hassan et al. [13].

As seen in Figure 5, the FW heuristic of Siegmund et al. (i.e., the sampling approach using the fewest number of configurations) has the higher error rate and the highest standard deviation on that error rate (four out of six times). Hence, we cannot recommend this method or, if one wishes to use this method, we recommend using the other sampling heuristics (e.g., HO, HS) to make more accurate predictions (but at the cost of much more measurements). Moreover, the size of the standard deviation of this method causes further difficulties in estimating which configurations are those exhibiting a large prediction error.

As to the approach of Guo et al (with PW), it does not stand out on any of our measurements. Its error results are within 1% of WHAT; its standard deviations are usually larger and it requires much more data than WHAT (Evaluations column of the figure 5).

In terms of the number of measure samples required to build a model, the right-hand column of Figure 5 shows that WHAT requires the fewest samples except for two cases: the approach of Guo et al. (with 2N) working on BDBC and LLVM. In both these cases, the mean error and standard deviation on the error estimate is larger than WHAT. Furthermore, in the case of BDBC, the error values are $\mu = 14\%$, $\sigma = 13\%$, which are much larger than WHAT’s error scores of $\mu = 6\%$, $\sigma = 5\%$.

Although the approach of Sarkar et al. produces an error rate that is sometimes less than the one of WHAT, it requires the highest number of measurements. Moreover, WHAT’s accuracy is close to Sarkar’s approach (1% to 2%) difference). Hence, we cannot recommend this approach, too.

Table 3 shows the number of evaluations used by each approaches. We see that most state-of-the-art approaches often require many more samples than WHAT. Using those fewest numbers of samples, WHAT has within 1% to 2% of the lowest standard deviation rates and within 1 to 2% of lowest error rates. The exception is Sarkar’s approach, which has 5% lower mean error rates (in BDBC, see the Mean MRE column of figure 5). However, as shown in right-hand side of Table 3, Sarkar’s approach needs nearly three times more measurements than WHAT.

To summarize, there are two cases in Figure 5 where WHAT performs worse than, at least, one other method:

– SQLite: The technique proposed by Sarkar et al. does better than WHAT (3.44 vs 5.6) but, as shown in the final column of Figure 5, it does so at the cost of $\frac{925}{64} \approx 15$ times more evaluations that WHAT. In this case, a pragmatic engineer could well prefer our solution over that of Sarkar et al. (since number of evaluations performed by Sarkar et al. more than an order of magnitude than WHAT).

– BDBC: Here again, WHAT is not doing the best but, compared to the number of evaluations required by all other solutions, it is not doing particularly bad.

Given the overall reduction of the error is small (5% difference between Sarkar and WHAT in mean error), the cost of tripling the data-collection cost is often not feasible in a practical context and might not justify the small additional benefit in accuracy.
Table 3 Comparison of the number of the samples required with the state of the art. The grey colored cells indicate the approach that requires the lowest number of samples. We notice that WHAT and Guo (2N) uses less data compared to the other approaches. The high fault rate of Guo (2N) accompanied with high variability in the predictions makes WHAT our preferred method.

| Samples | Siegmund | Guo (2N) | Guo (PW) | Sarkar | WHAT |
|---------|----------|----------|----------|--------|-------|
| Apache  | 29       | 181      | 29       | 55     | 16    |
| BDBC    | 139      | 36       | 139      | 191    | 64    |
| BDBJ    | 48       | 52       | 48       | 57     | 16    |
| LLVM    | 62       | 22       | 64       | 43     | 16    |
| SQLite  | 566      | 78       | 566      | 925    | 64    |
| X264    | 81       | 32       | 81       | 93     | 32    |

Based on the results of figure 5, we answer RQ4 with “yes”, since WHAT yields predictions that are similar to or more accurate than prior work, while requiring fewer samples.

6 Why does it work?

In this section, we present an in-depth analysis to understand why our sampling technique (based on a spectral learner) achieves such low mean fault rates while being stable (low variance). We hypothesize that the configuration space of the system configuration lie on a low dimensional manifold.

6.1 History

Menzies et. al [24] demonstrated how to exploit the underlying dimension to cluster data to find local homogeneous data regions in an otherwise heterogeneous data space. The authors used an algorithm called WHERE (see section 3.3), which recurses on two dimensions synthesized in linear time using a technique called FASTMAP [11]. The use of underlying dimension has been endorsed by various other researchers [1, 2, 7, 41]. There are numerous other methods in the literature, which are used to learn the underlying dimensionality of the data set such as Principal Component Analysis (PCA) [19], Spectral Learning [32] and Random Projection [3]. These algorithms use different techniques to identify the underlying, independent/orthogonal dimensions to cluster the data points and differ with respect to the computational complexity and accuracy. We use WHERE since it computationally efficient $O(2N)$, while still being accurate.

* WHERE is an approximation of the first principal component
6.2 Testing Technique

Given our hypothesis the configuration space lies in a lower dimensional hyperplane — it is imperative to demonstrate that the intrinsic dimensionality of the configuration space is less than the actual dimension. To formalize this notion, we borrow the concept of correlation dimension from the domain of physics [14]. The correlation dimension of a dataset with \( k \) items is found by computing the number of items found at distance within radius \( r \) (where \( r \) is the Euclidean distance between two configurations) while varying \( r \). This is then normalized by the number of connections between \( k \) items to find the expected number of neighbors at distance \( r \). This can be written as:

\[
C(r) = \frac{2}{k(k-1)} \sum_{i=1}^{n} \sum_{j=i+1}^{n} I(||x_i - x_j|| < r)
\]

where \( I(x < y) = \begin{cases} 1, & \text{if } x < y \\ 0, & \text{otherwise} \end{cases} \)

Given the dataset with \( k \) items and range of distances \([r_0, r_{max}]\), we estimate the intrinsic dimensionality as the mean slope between \( \ln(C(r)) \) and \( \ln(r) \).

6.3 Evaluation

On the configuration space of our subject systems, we observe that the intrinsic dimensionality of the software system is much lower than the actual dimension. Figure 6 presents the intrinsic dimensionality along with the actual dimensions of the software systems. If we take a look at the intrinsic dimensionality and compare it with the actual dimensionality, then it becomes apparent that the configuration space lies on a lower dimensional hyperplane. For example, SQLite has 39 configuration options, but the intrinsic dimensionality of the space is just 1.61 (this is a fractal dimension). At the heart of WHAT is WHERE (a spectral clusterer), which uses the approximation of the first principal component to divide the configuration space and hence can take advantage of the low intrinsic dimensionality.

As a summary, our observations indicate that the intrinsic dimension of the configuration space is much lower that its actual dimension. Hence, clustering based on the intrinsic dimensions rather than the actual dimension would be more effective. In other words, configurations with similar performance values lie closer to the intrinsic hyperplane, when compared to the actual dimensions, and may be the reason as to why WHAT achieves empirically good results.

7 Reliability and Validity

Reliability refers to the consistency of the results obtained from the research. For example, how well independent researchers could reproduce the study? To increase external reliability, we took care to either clearly define our algorithms or use implementations from the public domain (SciKitLearn) [26]. Also, all the data used in this
Fig. 6 The actual dimensions are shown on the x-axis and intrinsic dimensionality is shown on the y-axis. The points are annotated with the names of the corresponding software system. The intrinsic dimensionality of the systems are much lower than the actual dimensionality (number of columns in the dataset).

work are available on-line in the PROMISE code repository and all our algorithms are on-line at github.com/ai-se/where.

Validity refers to the extent to which a piece of research actually investigates what the researcher purports to investigate (13). Internal validity checks if the differences found in the treatments can be ascribed to the treatments under study.

http://openscience.us/repo/performance-predict/cpm.html
One threat to internal validity of our experiments is the choice of training and testing data sets discussed in Figure 1. Recall that, while all our learners used the same testing data set, our untuned learners were only given access to training data.

Another threat to internal validity is instrumentation. The very low $\mu$ and $\sigma$ error values reported in this study are so small that it is reasonable to ask whether they are due to some instrumentation quirk, rather than due to using a clever sample strategy:

- Our low $\mu$ values are consistent with prior work [29];
- As to our low $\sigma$ values, we note that, when the error values are so close to 0%, the standard deviation of the error is “squeezed” between zero and those errors. Hence, we would expect that experimental rigs that generate error values on the order of 5% and Equation 3 should have $\sigma$ values of $0 \leq \sigma \leq 5$ (e.g., like those seen in our introduction).

Regarding SQLite, we cannot measure all possible configurations in reasonable time. Hence, we sampled only 100 configurations to compare prediction and actual performance values. We are aware that this evaluation leaves room for outliers. Also, we are aware that measurement bias can cause false interpretations [24]. Since we aim at predicting performance for a special workload, we do not have to vary benchmarks.

We aimed at increasing the external validity by choosing software systems from different domains with different configuration mechanisms and implemented with different programming languages. Furthermore, our subject systems are deployed and used in the real world. Nevertheless, assuming the evaluations to be automatically transferable to all configurable software systems is not fair. To further strengthen external validity, we ran the model (generated by WHAT + S1) against other optimizers, such as NSGA-II and differential evolution [36]. That is, we validated whether the learned models are not only applicable for GALE style of perturbation. In Table 2, we see that the models developed are valid for all optimizers, as all optimizers are able to find the near optimal solutions.

8 Related Work

In 2000, Shi and Maik [31] claimed the term “spectral clustering” as a reference to their normalized cuts image segmentation algorithm that partitions data through a spectral (eigenvalue) analysis of the Laplacian representation of the similarity graph between instances in the data.

In 2003, Kamvar et al. [20] generalized that definition saying that “spectral learners” were any data-mining algorithm that first replaced the raw dimensions with those inferred from the spectrum (eigenvalues) of the affinity (a.k.a. distance) matrix of the data, optionally adjusted via some normalization technique.

Our clustering based on first principal component splits the data on an approximation to an eigenvector, found at each recursive level of the data (as described in §3.1). Hence, this method is a “spectral clusterer” in the general Kamvar sense. Note that, for our data, we have not found that Kamvar’s normalization matrices are needed.

Regarding sampling, there are a wide range of methods know as experimental designs or designs of experiments [38]. They usually rely on fractional factorial designs as in the combinatorial testing community [22].
Furthermore, there is a recent approach that learns performance-influence models for configurable software systems [34]. While this approach can handle even numeric features, it has similar sampling techniques for the Boolean features as reported in their earlier work [35]. Since we already compared to that earlier work and do not consider numeric features, we did not compare our work to performance-influence models.

9 Conclusions

Configurable software systems today are widely used in practice, but they impose challenges regarding finding performance-optimal configurations. State-of-the-art approaches require too many measurements or are prone to large variances in their performance predictions. To overcome these limitations, we have proposed a fast spectral learner, called WHAT, along with three new sampling techniques. The key idea of WHAT is to explore the configuration space with eigenvalues of the features used in a configuration to determine exactly those configurations for measurement that reveal key performance characteristics. This way, we can study many closely associated configurations with only a few measurements.

We evaluated our approach on six real-world configurable software systems borrowed from the literature. Our approach achieves similar to lower error rates, while being stable when compared to the state of the art. In particular, with the exception of Berkeley DB, our approach is more accurate than the state-of-the-art approaches by Siegmund et al. [35] and Guo et al. [15]. Furthermore, we achieve a similar prediction accuracy and stability as the approach by Sarkar et al. [29], while requiring a far smaller number of configurations to be measured. We also demonstrated that our approach can be used to build cheap and stable surrogate prediction models, which can be used by off-the-shelf optimizers to find the performance-optimal configuration. We use the correlation dimension to demonstrate how the high dimensional configuration space of our subject systems has a low intrinsic dimensionality, which might be the reason why WHAT performs so well on these datasets.

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