Active Learning for SAT Solver Benchmarking

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Abstract. Benchmarking is one of the most important phases when developing algorithms. This also applies to solutions of the SAT (propositional satisfiability) problem. While the field of SAT solver benchmarking is well established, traditional benchmark selection approaches do not optimize for benchmark runtime. Benchmark selection chooses representative instances from a pool of instances such that they reliably discriminate SAT solvers based on their runtime. In this paper, we present a dynamic benchmark selection approach based on active learning. Our approach predicts the rank of a new solver among its competitors with minimum running time and maximum rank prediction accuracy. We evaluated this approach using the Anniversary Track dataset from the 2022 SAT Competition. Our selection approach can predict the rank of a new solver after about 10% of the time it would take to run the solver on all instances of this dataset, with a prediction accuracy of about 92 %. In this paper, we also discuss the importance of instance families in the selection process. Overall, our tool provides a reliable way for SAT solver engineers to efficiently determine the performance of a new SAT solver.

Keywords: Propositional satisfiability · Benchmarking · Active learning

1 Introduction

One of the main phases of algorithm engineering is benchmarking. This is also true for the propositional satisfiability problem (SAT), the archetypal \mathcal{NP} -complete problem. While state-of-the-art solvers embark upon more and more problem instances, SAT solver evaluation is steadily getting more costly. Competitive SAT solvers are the result of extensive experimentation and a variety of ideas and considerations [8,2]. While the field of SAT solver benchmarking is well established, traditional benchmark selection approaches do not optimize for benchmark runtime. Instead, the primary goal of traditional approaches is to select a representative set of instances for ranking solvers with a scoring scheme [10,16]. SAT Competitions typically use the widely adopted PAR-2 score, i.e., the average runtime with a penalty of 2τ for timeouts with time limit τ [8].

In this paper, we present a novel benchmark selection approach based on active learning. Our approach is able to predict the rank of a new solver with high accuracy in only a fraction of the time needed to evaluate the full benchmark. The problem we solve is specified in Definition 1.

Definition 1 (New-Solver Problem). Given a pool of instances \mathcal{I} , solvers \mathcal{A} , and runtimes $r: \mathcal{A} \times \mathcal{I} \to [0, \tau]$ with time limit τ , maximize the confidence in predicting the rank of a new solver $\hat{a} \notin \mathcal{A}$ while minimizing the total benchmark runtime by incrementally selecting instances in \mathcal{I} to evaluate \hat{a} .

Note that our scenario assumes that we know the runtimes of all solvers, except the new one, on all instances. One could also imagine a collaborative filtering scenario, where runtimes are only partially known [23,25].

Our approach satisfies several desirable criteria for benchmarking: It outputs a ranking to compare the new solver against the existing set of solvers. For this ranking, we do not even need to predict exact solver runtimes, which is trickier. We optimize the *runtime* that our strategy needs to arrive at its conclusion. We use problem-instance and runtime features. Moreover, we select instances non-randomly and incrementally. In particular, we take runtime information from already done experiments into account when choosing the next. By doing so, we can control the properties of the benchmarking approach such as its required runtime. Rather than solely outputting a binary classification, i.e, the new solver is worse than an existing solver, we provide a scoring function that shows by which margin a solver is worse and how similar it is to existing solvers. Our approach is scalable in that it ranks a new solver \hat{a} among any number of known solvers A. In particular, we only subsample the benchmark once instead of comparing pairwise against each other solver [21]. Since a new solver idea is rarely best on the first try, it is desired to obtain a solver ranking fast. In this way, a new solver idea can be discarded if it performs poorly across the board or may be further tweaked if it shows promising results at least in some cases.

We evaluate our approach with the SAT Competition 2022 Anniversary Track benchmark [2], consisting of 5355 instances and full runtimes of 28 solvers. We perform cross-validation by treating each solver as new once and predicting these solvers' PAR-2 rank. On average, our predictions reach about 92 % accuracy with only about 10 % of the runtime that would be needed to evaluate these solvers on the complete set of instances.

All our source code¹ and and data² are available on GitHub.

2 Related Work

Benchmarking is not only of high interest in many fields but also an active research area on its own. Recent studies show that benchmark selection is a difficult endeavor for multiple reasons. Biased benchmarks can easily lead to fallacious interpretations [7]. Also, benchmark selection has many movable parts, e.g., performance measures, aggregation, and handling of missing values. Questionable research practices might modify these elements a-posteriori to fit expectations, leading to biased results [27]. In the following, we discuss related work from the areas of static benchmark selection, algorithm configuration, incremental benchmark selection, and active learning.

 $^{^{1}}$ temporary, an onymized version for review: ${\tt xxx}$

² temporary, anonymized version for review: xxx

Feature	Hoos [16]	SMAC [17]	Matricon [21]	Our approach
Ranking	✓	Х	✓	✓
Runtime Minimization	X	✓	✓	✓
Incremental/Non-Random	X	×	✓	✓
Scoring	✓	×	X	✓
Scalability	✓	✓	×	✓

Table 1: Support for desired criteria from Section 1. The table compares a static benchmark-selection approach [16], an algorithm configuration system [17], an existing active-learning approach [21], as well as our approach.

Static Benchmark Selection. Benchmark selection is an important issue for competitions, e.g., the SAT Competition. In such competitions, the organizers define the rules for composing the corresponding benchmarks. Balint et al. provide an overview of benchmark-selection criteria in different solver competitions [1]. Manthey and Möhle find that competition benchmarks might contain redundant instances, and propose a feature-based approach to remove redundancy [20]. Misir presents a feature-based approach to reduce benchmarks by matrix factorization and clustering [24].

Hoos et al. discuss which properties are most desirable when selecting SAT benchmark instances [16]. The selection strategy presented is static, i.e., it does not depend on particular solvers to distinguish. Selection criteria are instance variety to avoid over-fitting, adapted instance hardness (not too easy but also not too hard), and avoiding duplicate instances. To filter too similar instances, they use a distance-based approach with the SATzilla features [35,36]. Their approach ranks solvers, is feature-based, and scalable in the sense that the evaluation time scales linearly with more instances and solvers. Also, it allows scoring and thereby comparing solvers. However, the approach does not optimize for benchmark runtime and selects instances randomly, apart from constraints on the instance hardness and feature distance.

Algorithm Configuration. Further related work can be found within the field of algorithm configuration [15,32], e.g., the configuration system SMAC [17]. There, the goal is to tune SAT solvers for a given sub-domain of problem instances. Although this task is different from our goal, e.g., we do not need to navigate configuration space, there are similarities to our approach as well. For example, SMAC also employs an iterative, model-based selection procedure, though for configurations rather than instances. SMAC's configuration sampling has also proven to be better than random sampling in all tested scenarios and has even proven to be significantly better in most cases. While this approach considers the configuration's runtime, is feature-based, and scales by having a fixed time budget, it does not satisfy the other three criteria listed in Table 1. First, an algorithm configurator cannot be used to rank a new solver since it only looks at promising solvers/configurations rather than the overall average performance.

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Second, while using a model-based selection strategy to sample configurations, instance sampling is done *randomly*, i.e., without building a model over instances. And third, only few configurations within their approach are *scored* since algorithm configuration seeks to find solemnly the best performing configuration.

Incremental Benchmark Selection. Matricon et al. present an incremental benchmark selection approach [21]. Their per-set efficient algorithm selection problem (PSEAS) is similar to our New-Solver Problem (cf. Definition 1). Given a pair of SAT solvers, they iteratively select a subset of instances until the desired confidence level is reached to decide which of the two solvers is better. The selection of instances is dependent on the choice of the solvers to distinguish. They calculate a scoring metric for all unselected instances, run the experiment with the highest score, and update the confidence. Most of the proposed strategies in [21] are instance-based, i.e., no model of the solver runtimes is built. Regarding our desired criteria of benchmark selection (Table 1), the aforementioned approach ranks solvers, optimizes for runtime, is feature-based, and uses incremental non-random sampling. However, the approach only compares solvers binarily rather than providing a scoring. Thus, it is not clear how similar two given solvers are or on which instances they behave similarly. Moreover, a major shortcoming is the lacking scalability with the number of solvers. Comparing only pairs of solvers, evaluating a new solver requires sampling several benchmarks. In contrast, our approach allows comparing a new solver against a set of existing solvers with one benchmark.

Active Learning (AL). The posed New-Solver Problem has stark similarities to the well-studied field of active learning AL within recommender systems, especially the new-user problem [29]. On the one hand, we want to maximize the utility an instance provides to our model and, on the other hand, minimize the cost (CPU time) that is associated with its acquisition. In contrast to traditional passive machine-learning methods with given instance labels, active learning allows for selecting instances for which to acquire labels. AL algorithms can be categorized into synthesis-based [5,9,33] and pool-based approaches [12,14,19]. While synthesis-based methods generate instances for labeling, pool-based methods rely on a fixed set of unlabeled instances from which to sample.

Recent synthesis-based methods within the field of SAT solving show how to generate problem instances with desired properties. This goal is, however, orthogonal to ours [5,9]. While those approaches want to generate problem instances on which a solver is good or bad, we want to predict whether a solver is good or bad on an existing benchmark. Volpato and Guangyan use pool-based AL to learn an instance-specific algorithm selector [34]. Rather than benchmarking a solver's overall performance, the goal is to recommend the best solver out of a set of solvers for each SAT instance.

3 Active Learning for SAT Solver Benchmarking

The outline of our benchmarking framework is depicted in Algorithm 1. Given a set of known solvers \mathcal{A} , instances \mathcal{I} and runtimes r, we first create a prediction model \mathcal{M} for the new solver \hat{a} (Line 1). The prediction model is used to repeatedly select an instance (Line 4) for benchmarking \hat{a} (Line 5). The acquired result is subsequently used to update the prediction model \mathcal{M} (Line 6). When the stopping criterion is met (Line 3), we quit the benchmarking loop and predict the final score of \hat{a} (Line 8). Algorithm 1 returns the predicted score of \hat{a} as well as the acquired runtime measurements (Line 9).

Section 3.1 specifies the details of the underlying prediction model and describes how we may derive a solver ranking from that model. We discuss possible criteria for sampling instances in Section 3.2. Section 3.3 concludes with possible stopping conditions.

Since the potential runtime of experiments is by magnitudes larger than the model's update time, we only consider incrementing our benchmark by one instance at a time rather than using batches, which is also proposed in current active-learning advances [31,33].

Algorithm 1: Incremental Benchmarking Framework

```
Input: Solvers \mathcal{A}, Instances \mathcal{I}, Runtimes r: \mathcal{A} \times \mathcal{I} \to [0, \tau], Solver \hat{a}
    Output: Predicted Score of \hat{a}, Measured Runtimes \mathcal{R}
1 \mathcal{M} \leftarrow \text{initModel}(\mathcal{A}, \mathcal{I}, r, \hat{a})
2 \mathcal{R} \leftarrow \emptyset
                                                                                                             // cf. Section 3.3
3 while not stop(\mathcal{M}) do
                                                                                                             // cf. Section 3.2
          e \leftarrow \text{selectNextInstance}(\mathcal{M})
          t \leftarrow \text{runExperiment}(\hat{a}, e)
                                                                                     // Runs \hat{a} on e with time-out \tau
          updateModel(\mathcal{M}, (e, t))
                                                                                                            // cf. Section 3.1
      \mathcal{R} \leftarrow \mathcal{R} \cup \{(e,t)\}
8 s_{\hat{a}} \leftarrow \operatorname{predictScore}(\mathcal{M})
                                                                                                             // cf. Section 3.1
9 return (s_{\hat{a}}, \mathcal{R})
```

3.1 Solver Model

Let benchmark instances \mathcal{I} , SAT solvers \mathcal{A} , and runtimes $r: \mathcal{A} \times \mathcal{I} \to [0, \tau]$ be given. We denote the new solver to be ranked by $\hat{a} \notin A$ and define $\hat{A} := A \cup \{\hat{a}\}$ to denote all solvers including the new solver. Model \mathcal{M} provides a prediction function $f: \hat{\mathcal{A}} \times \mathcal{I} \to \mathbb{R}$ that powers the decisions within our framework.

To be applicable to a new solver \hat{a} , whose runtimes are unknown, we extend score by replacing $\gamma(a,e)$ with the model's predictions $f: \hat{\mathcal{A}} \times \mathcal{I} \to \{1,\ldots,k\}$.

Runtime Transformation. We transform the runtimes into discrete runtime labels on a per-instance basis. For each instance $e \in \mathcal{I}$, we use a single-link hierarchical clustering algorithm to assign the runtimes in $\{r(a,e) \mid a \in A\}$ to one of the k clusters C_1, \ldots, C_k such that the fastest runtimes for instance e are in cluster C_1 and the slowest are in cluster C_k . For instances with many timeouts, we have to manually fix the clustering results such that all timeouts τ are in cluster C_k . The runtime transformation function $\gamma_k : \mathcal{A} \times \mathcal{I} \to \{1, \ldots, k\}$ is then specified as follows:

$$\gamma_k(a,e) = j \iff r(a,e) \in C_j$$

Given an instance $e \in \mathcal{I}$, a solver $a \in A$ belongs to the $\gamma_k(a, e)$ -fastest solvers on instance e. In preliminary experiments, we achieved higher accuracy predicting such discrete runtime labels than predicting raw runtimes. The mean squared error of raw runtime regression was within the same magnitude as the values to be predicted. Research on portfolio solvers has also shown that directly predicting runtimes without any transformation does not work well in practice [4,26].

To determine solver ranks, we use the transformed runtimes $\gamma_k(a, e)$ in the adapted scoring function $s_k : A \to [1, 2 \cdot k]$ as follows:

$$s_k(a) := \frac{1}{|\mathcal{I}|} \sum_{e \in \mathcal{I}} \gamma_k'(a, e) \qquad \gamma_k'(a, e) := \begin{cases} 2 \cdot \gamma_k(a, e) & \text{if } \gamma_k(a, e) = k \\ \gamma_k(a, e) & \text{otherwise} \end{cases}$$
(1)

I.e., we apply a PAR-2 scoring, which is commonly used in SAT competitions [8], on the discrete labels. The scoring function s_k induces a ranking among solvers.

However, we need to ensure that discretized labels still discriminate solvers and align with the actual PAR-2 ranking. We analyzed the ranking induced by s_3 in preliminary experiments with the SAT Competition 2022 Anniversary Track [2]. According to a Wilcoxon-signed-rank test with $\alpha=0.05,\,87.83\,\%$ of solver pairs have significantly different scores after discretization, only a slight drop compared to 89.95 % before discretization. Further, our ranking approach is capable of correctly deciding for almost all (about 97.45 %; $\sigma=3.68\,\%$) solver pairs which solver is faster. In particular, the Spearman correlation of s_3 and PAR-2 ranking is about 0.988, which is very close to the optimal value of 1 [6]. All these results show that discretized runtimes are suitable for our framework.

3.2 Instance Sampling

Selecting an instance based on the model is a core functionality of our framework (cf. Algorithm 1, Line 2). In this section, we introduce our sampling strategies, which make use of the model's label-prediction function f and are inspired by existing work within the realms of active learning [30]. We implement a model-uncertainty and an information-gain sampling strategy. These methods require the model's predictions to also include probabilities for the k possible runtime labels. Let $f': \hat{\mathcal{A}} \times \mathcal{I} \to [0,1]^k$ denote this modified prediction function.

Model Uncertainty Sampling. The model-uncertainty sampling strategy simply selects the instance that is closest to the model's decision boundary.

$$\underset{e \in \mathcal{I} \setminus \tilde{\mathcal{I}}}{\operatorname{arg\,min}} \left| \frac{1}{k} - \max_{n \in \{1, \dots, k\}} f'(\hat{a}, e)_n \right|$$

Information Gain Sampling. The information-gain sampling strategy selects the instance with the highest expected entropy reduction regarding the runtime labels of the instance. To be more specific, we select the instance e that maximizes IG(e), which is specified as follows:

$$IG(e) := H(e) - \sum_{n=1}^{k} f'(\hat{a}, e)_n \, \hat{H}_n(e)$$

Here, H(e) denotes the entropy of the runtime labels $\gamma(a,e)$ over all $a \in \mathcal{A}$ and H(e,n) denotes the entropy of these labels plus n as the runtime label for \hat{a} . The term $\hat{H}_n(e)$ is computed for every possible runtime label $n \in \{1,\ldots,k\}$.

3.3 Stopping Criteria

In this section, we present two dynamic stopping criteria, the Wilcoxon and the Model Uncertainty Stopping Criterion (cf. Algorithm 1, Line 1).

Wilcoxon Stopping Criterion. The Wilcoxon stopping criterion stops the active-learning process when we are certain enough that the predicted runtime labels of the new solver are sufficiently different from the labels of the existing solvers. We use the average p-value $W_{\hat{a}}$ of a Wilcoxon signed-rank test w(S, P) of the two runtime label distributions $S = \{\gamma(a, e) \mid e \in \mathcal{I}\}$ for an existing solver a and $P = \{f(\hat{a}, e) \mid e \in \mathcal{I}\}$ for the new solver \hat{a} :

$$W_{\hat{a}} := \frac{1}{|\mathcal{A}|} \sum_{a \in \mathcal{A}} w(S, P)$$

To improve the stability of this criterion, we use an exponential moving average to smooth out outliers and stop as soon as $W_{\text{exp}}^{(i)}$ drops below a certain threshold:

$$W_{\text{exp}}^{(0)} := 1$$

 $W_{\text{exp}}^{(i)} := \beta W_{\hat{a}} + (1 - \beta) W_{\text{exp}}^{(i-1)}$

Model Convergence Stopping Criterion. The model-convergence stopping criterion is less sophisticated in comparison. It stops the active-learning process if the ranking induced by the model's predictions (Equation 1) remains unchanged within the last l iterations. Note that the concrete values of $s_k(\hat{a})$ might still change. We are solemnly interested in the induced ranking in this case.

4 Experimental Design

Given all the previously presented instantiations for Algorithm 1, this section briefly outlines our experimental design, including our optimization goal, evaluation framework, used data sets, hyper-parameter choices, as well as implementation details.

4.1 Optimization Goal

As already stated in the introductory section, this work addresses the New-Solver Problem (cf. Definition 1). We create a prediction model \mathcal{M} as described in Section 3.1 that provides us with a scoring function s_k . First and foremost, our goal is to provide the engineer of new SAT solvers with an accurate ranking. We define the ranking accuracy $O_{\text{acc}} \in [0,1]$ (higher is better) by the fraction of pairs (\hat{a},a) for all $a \in \mathcal{A}$ that are decided correctly by the given ranking (cf. Algorithm 2). For now, we exclude the equality of solvers since data shows that the PAR-2 scores of all our solvers are different. So, possible ranking decisions may be, for example, solver a is better than b or b is better than a. The ranking accuracy is affected by whether this decision is correctly made. Second, we also have to optimize for runtime. The fraction of runtime that the algorithm needs to arrive at its conclusion is denoted by $O_{\text{rt}} \in [0,1]$ (lower is better). This metric puts the runtime summed over the sampled instances in relation to the runtime summed over all instances in the dataset (cf. Algorithm 2). Overall, we want to find an approach that maximizes

$$O_{\delta} := \delta O_{\text{acc}} + (1 - \delta) \left(1 - O_{\text{rt}} \right) , \qquad (2)$$

whereby $\delta \in [0,1]$ allows for linear weighting between the two optimization goals $O_{\rm acc}$ and $O_{\rm rt}$. Plotting the approaches that maximize O_{δ} for all $\delta \in [0,1]$ on a $O_{\rm rt}$ - $O_{\rm acc}$ -diagram provides us with a Pareto front of the best approaches for different optimization-goal weightings.

4.2 Evaluation Framework

To evaluate a concrete instantiation of Algorithm 1 (a concrete choice for all the sub-routines), we perform cross-validation on our set of solvers. That means that each solver once plays the role of the new solver. Algorithm 2 shows this. Note that the *new* solver in each iteration is excluded from the set of solvers \mathcal{A} to avoid data leakage. Also, the runtime function r is restricted to $\mathcal{A} \times \mathcal{I}$.

4.3 Data

In our experiments, we work with the SAT Competition 2022 Anniversary Track instances [2]. The dataset consists of 5355 instances with respective runtime data. It has complete runtimes of 28 solvers. We use the GBD metadata database [18] to provide us with the aforementioned instance features, problem instances, and

Algorithm 2: Evaluation Framework

```
Input: Solvers \mathcal{A}, Instaces \mathcal{I}, Runtimes r: \mathcal{A} \times \mathcal{I} \to [0, \tau]
        {\tt Output}\colon {\tt Average}Rank Accuracy \bar{O}_{\tt acc}, Average Fraction of Runtime \bar{O}_{\tt rt}
  1~O \leftarrow \emptyset
 2 for \hat{a} \in \mathcal{A} do
                 \mathcal{A}' \leftarrow \mathcal{A} \setminus \{\hat{a}\}
  3
                  (s_{\hat{a}}, \mathcal{R}) \leftarrow \text{runALAlgorithm}(\mathcal{A}', \mathcal{I}, r, \hat{a})
                                                                                                                                                    // Refer to Algorithm 1
                  // Determine Rank Accuracy
                  O_{\rm acc} \leftarrow 0
  5
                 for a \in \mathcal{A} do
  6
                          \begin{array}{l} \text{if } \left( s_k(a) - s_{\hat{a}} \right) \cdot \left( \operatorname{par}_2(a) - \operatorname{par}_2(\hat{a}) \right) > 0 \text{ then} \\ \left[ \begin{array}{c} O_{\operatorname{acc}} \leftarrow O_{\operatorname{acc}} + \frac{1}{|\mathcal{A}|} \end{array} \right] \end{array}
  7
                 // Determine Runtime Fraction
                 r \leftarrow \sum_{e \in \mathcal{I}} r(\hat{a}, e)O_{\text{rt}} \leftarrow 0
 9
10
                  for e \in \mathcal{I} do
11
                      \left| \begin{array}{c} \text{if } \exists t, (e,t) \in \mathcal{R} \text{ then} \\ \left| \begin{array}{c} O_{\mathrm{rt}} \leftarrow O_{\mathrm{rt}} + \frac{t}{r} \end{array} \right. \end{array} \right| 
12
13
               O \leftarrow O \cup \{(O_{\mathrm{acc}}, O_{\mathrm{rt}})\}
15 (\bar{O}_{acc}, \bar{O}_{rt}) \leftarrow average(O)
16 return (\bar{O}_{\mathrm{acc}}, \bar{O}_{\mathrm{rt}})
```

solver runtimes. Thereby, all our approaches make use of the 56 base features that are maintained in the base database of GBD [18]. They are inspired by the SATzilla features [36]. Those features comprise general instance-size features and graph-representation features among others. All features are numeric and fortunately free of any missing values. We drop 10 out of 56 features because of zero variance. For hyper-parameter tuning, we randomly sample $10\,\%$ of the complete set of 5355 instances with stratification regarding the instance's family. All instance families that are too small, i.e., $10\,\%$ of them corresponds to less than one instance, are put into one meta-family for stratification. This smaller dataset allows for a more extensive exploration of hyper-parameter space.

4.4 Hyper-parameters

Given Algorithm 1, there are several possible instantiations for the three phases, i.e., *selection*, *stopping*, and *ranking*. Also, there are different choices for the runtime-label prediction model. Note that we are not considering all previously listed approaches since a grid search of all combinations would be infeasible. Rather, we filter approaches based on preliminary experimental results (cf. Section 5.1) and do the main end-to-end experiments only with a subset. The end-to-end experiment configurations are given below.

Ranking. Regarding *ranking* (cf. Section 3.1), we experiment with the following approaches, including our used hyper-parameter values:

- Observed PAR-2 ranking of already sampled instances
- Predicted ranking induced by runtime-label predictions
 - History size: consider the latest 1, 10, 20, 30, or 40 predictions within a voting approach for stability. The latest x predictions vote on the winner.
 - Fallback threshold: if the difference of scores between two solvers drops below 0.01, 0.05, or 0.1, use the partially observed PAR-2 ranking as a tie-breaker. As discussed in Section 3.1, ranking decisions are all correct if the difference in scores between two solvers exceeds a certain value.

Selection. For *selection* (cf. Section 3.2), we experiment with the following methods, including our used hyper-parameter values:

- Random sampling
- Model-based uncertainty sampling
 - Fallback threshold: use random sampling for the first 0%, 5%, 10%, 15%, or 20% of instances to explore the instance space.
 - Runtime scaling: prefer instances with the greatest uncertainty per average (over all solvers) runtime (True or False).
- Model-based information-gain sampling
 - Fallback threshold: use random sampling for the first 0%, 5%, 10%, 15%, or 20% of instances to explore the instance space.

- Runtime scaling: prefer instances with the greatest information-gain per average runtime (True or False).
- Neighborhood-aware random sampling (preliminary experiments only); randomly chooses among the instances with the highest amount of non-sampled k neighbors
- Ranking-based sampling [3] (preliminary experiments only)
- Discrimination-based sampling [11] (preliminary experiments only)
- Variance-based sampling [21] (preliminary experiments only)

Stopping. For *stopping* decisions (cf. Section 3.3), we experiment with the following criteria, including our used hyper-parameter values:

- Fixed subset size of $10\,\%$ or $20\,\%$ of instances
- Ranking convergence criterion
 - Minimum amount: sample at least 2%, 8%, 10%, or 12% of instances before applying the criterion.
 - Convergence duration: stop if the predicted ranking stays the same for the last 1% or 2% of sampled instances.
- Wilcoxon criterion
 - Minimum amount: sample at least 2%, 8%, 10%, or 12% of instances before applying the criterion.
 - Average of p-values to drop below: 5%.
 - Exponential-moving average: incorporate previous significance values by using an EMA with $\beta=0.1$ or $\beta=0.7$.

Runtime-label prediction model. As runtime-label prediction model, we only make use of one fixed model since an exhaustive grid search would be infeasible (cf. Section 5.1). Our model of choice is an ensemble, stacking a quadratic-discriminant analysis³ onto a random forest model⁴ using a decision tree⁵ to weight its members. If not stated explicitly, all other model parameters default to the presets of $scikit-learn^6$ [28].

4.5 Implementation Details

For reproducibility, our source code, all experiments, and data are available on GitHub⁷. Our code is implemented in Python using *scikit-learn* [28] for making predictions and *gbd-tools* [18] for SAT-instance retrieval.

 $^{^3}$ With a regularization parameter of zero and a singular value threshold of zero

⁴ With *entropy* splitting criterion and *balanced* class weights

With Gini impurity splitting criterion; choosing the best split for each branching decision and maximum tree depth of 5 levels

⁶ https://scikit-learn.org/stable/index.html; Version 1.0.2

⁷ https://github.com/mathefuchs/al-for-SATsolver-benchmarking

5 Evaluation

Before evaluating our approach end-to-end as described in Section 4, we discuss the choice of the machine-learning predictor and the runtime-discretization method. Thereafter, we look at the results of our active-learning approach. Finally, we also analyze the importance of different instance families to our framework. Instance families comprise instances derived from the same application domain, e.g., planning, cryptography, etc., and are a useful tool for both analyzing solver performance and portfolios.

5.1 Runtime Prediction

An exhaustive grid search of all active-learning approaches outlined in Section 3 and different runtime-prediction models is infeasible. Therefore, we compare predictors first and select one runtime-prediction model for all further experiments. We evaluate predictors with the *Matthews Correlation Coefficient* (MCC) scores [13,22] of the classification task. MCC scores are great in dealing with class imbalances in contrast to conventional metrics like accuracy.

We perform cross-validation among all solvers: we assume that all other solver runtimes are known and provide $64\,\%$ of ground-truth runtime labels of the target solver for training, use $16\,\%$ as a validation set for hyper-parameter tuning of the runtime predictor, and $20\,\%$ as a test set (5-fold cross-validation). This is repeated for each solver once. Note that we perform cross-validation on two levels, i.e., among the solvers and the target solver runtimes. We compare the average performances on the test sets.

ML classification models that we have tried include random forests, decision trees, quadratic-discriminant analysis, AdaBoost, logistic regression, k-nearest neighbor, multi-layer perceptrons, support vector machines, and naive Bayes. A stacking ensemble consisting of quadratic-discriminant analysis and random forests performed best. It produces an MCC score of 0.8882 for predicting three-class runtime labels. The results are significantly better than any other model that we have tried regarding a Wilcoxon signed-rank test with $\alpha=5\,\%$.

For runtime discretization, we use hierarchical clustering with k=3 and a log-single-link criterion for all further experiments. Each non-time-out runtime starts in a separate interval. We then gradually merge intervals whose single-link logarithmic distance is the smallest until the desired number of partitions is reached. As already mentioned in Section 3.1, this approach has good properties regarding the predictiveness of the PAR-2 score ranking and its discriminatory power. Also, experiments showed that our stacking ensemble performed particularly well for this kind of discretization method. Other clustering approaches that we have tried include hierarchical clustering with mean-, median- and complete-link criteria, as well as k-means and spectral clustering.

5.2 Active-Learning Results

Tuning our algorithm. Our end-to-end experiments follow the evaluation framework introduced in Section 4.2. Fig. 1 shows the performance of the ap-

proaches from Section 4.4 on $O_{\rm rt}$ - $O_{\rm acc}$ -diagrams for the smaller hyper-parameter-tuning dataset. Evaluating a particular configuration with Algorithm 2 returns a point $(O_{\rm rt},\,O_{\rm acc})$. The plotted lines represent the best performing configurations (convex hull) per top-level hyper-parameter choice, i.e., per ranking approach (Fig. 1a), selection approach (Fig. 1b), and stopping criterion (Fig. 1c).

Regarding ranking approaches (Fig. 1a), using the s_3 -induced ranking consistently outperforms the partially observed PAR-2 ranking for each possible value of the runtime-accuracy trade-off parameter δ . This is expected since selection decisions are not random. For example, we might sample more instances of one family if it benefits discrimination of solvers. While the partially observed PAR-2 score is skewed, the prediction model can account for this.

Regarding the selection approaches (Fig. 1b), the model-based uncertainty strategy performs best in most cases. However, the model-based information-gain sampling is beneficial if runtime is strongly favored (very small δ ; runtime fraction less than 5%). Random sampling performs quite well but is outperformed by the model-based uncertainty strategy in all cases, showing the benefit of actively selecting instances based on a prediction model.

Regarding the stopping criterion (Fig. 1c), the ranking convergence criterion performs the most consistently well. If accuracy is strongly favored (very high δ), the Wilcoxon stopping criterion performs better.

Figure 2 shows an interesting consequence of weighting our optimization goals: If we, on the one hand, desire to get a rough estimate of a solver's performance fast (low δ), approaches favor selecting many easy instances. In particular, the fraction of sampled instances is bigger than the fraction of runtime. By having many observations, it is easier to build a model. If we, on the other hand, desire to get a good estimate of a solver's performance in a moderate amount of time (high δ), approaches favor selecting fewer difficult instances. In particular, the fraction of instances is smaller than the fraction of runtime.

Full dataset evaluation. Having selected the most promising hyper-parameters, we run our end-to-end active-learning experiments on the complete Anniversary Track dataset (5355 instances). The best-performing approach for $\delta=0.85$ uses runtime-label prediction ranking, model-based uncertainty sampling, and a ranking-convergence stopping criterion. It can predict a new solver's PAR-2 ranking with about 92.33% accuracy $(O_{\rm acc})$, while only needing 10.35% of the time for running the solver on the full dataset $(O_{\rm rt})$. This trade-off provides a better way of prototyping and evaluating new SAT solvers rather than evaluating them on manually picked instance based on domain knowledge. The sampled instances account for about 5.24% of instances, indicating a preference for selecting difficult instances. The average Spearman correlation between the PAR-2 ranking and the predicted ranking is about 0.9572, indicating a high similarity of these two rankings. Because of that, the runtime-prediction model helps determine on what problem instances solvers perform similarly or are better/worse.

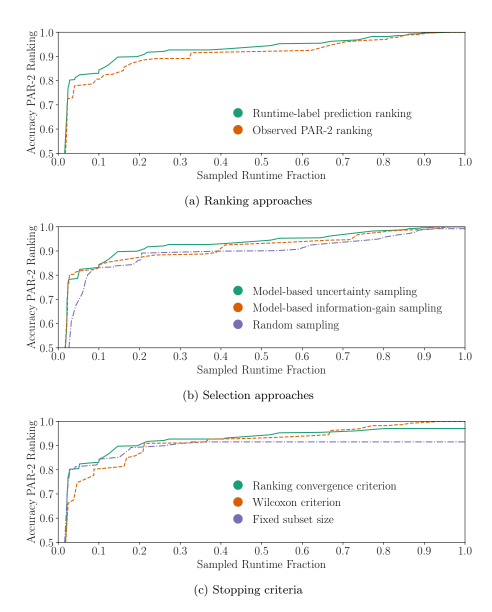


Fig. 1: $O_{\rm rt}$ - $O_{\rm acc}$ -diagrams comparing different hyper-parameter instantiations of our active-learning framework. The x-axis shows the ratio of total solver runtime on the sampled instances relative to all instances. The y-axis shows the ranking accuracy (cf. Section 4.1). Each line entails the front of Pareto-optimal configurations for the respective hyper-parameter instantiation.

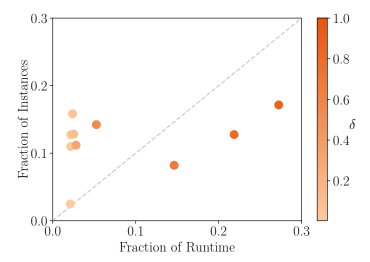


Fig. 2: Scatter plot comparing different instantiations of δ for our active-learning strategy on the hyper-parameter-tuning dataset. The x-axis shows the fraction of runtime $O_{\rm rt}$ of the sample, while the y-axis shows the fraction of instances sampled. The color indicates the weighting between different optimization goals $\delta \in [0, 1]$. The larger δ , the more we favor accuracy over runtime.

5.3 Instance-Family Importance

Selection decisions of our approach also reveal the importance of different instance families to our framework. Figure 3 shows the occurrence of instance families within the dataset and the benchmarks created by active learning. We use the best-performing configurations for all $\delta \in [0,1]$ and examine the selection decisions by the active-learning approach on the SAT Competition 2022 Anniversary Track dataset [2]. While most families appear with the same fraction in the dataset and the sampled benchmarks, there are a few outliers that need further discussion. Problem instances of the families fpga, quasigroup-completion, and planning are especially helpful to our framework in distinguishing solvers. Instances of these families are selected over-proportionally in comparison to the full dataset. In contrast to that, instances of the biggest family, i.e., hardwareverification, roughly appear with the same fraction in the dataset and the sampled benchmarks. Finally, instances of the family *cryptography* are less important in distinguishing solvers than their huge weight in the dataset suggests. A possible explanation is that these instances are very similar, such that a small fraction of them is sufficient to estimate a solver's performance on all of them.

6 Conclusion

In this work, we have discussed possible solutions to the *New-Solver Problem*: Given a new solver, we want to find its ranking amidst its competitors. Our

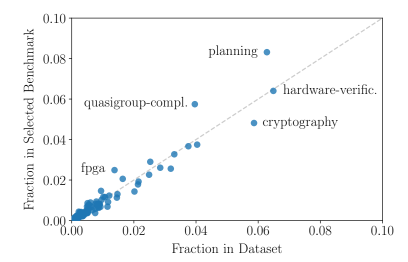


Fig. 3: Scatter plot showing the *importance* of different instance families to our framework. The x-axis shows the frequency of instance families in the full Anniversary Track dataset. The y-axis shows the average frequency of instance families in the samples selected by active learning. The dashed line represents families that are occur with the same frequency in both, dataset and samples.

approach provides accurate ranking predictions while needing significantly less runtime that a full evaluation on a given benchmark set. In particular, we can determine a new solver's PAR-2 ranking with about $92\,\%$ accuracy, while only needing $10\,\%$ of the full-evaluation time. We have found that our framework produces good results working with discrete runtime labels rather than regressing actual runtimes, albeit the former being more simple. We have evaluated several ranking algorithms, instance-selection approaches, and stopping criteria within our sequential active-learning process. We also took a brief look at which instance families are the most prevalent in selection decisions.

6.1 Future Work

Future work may compare further ranking algorithms, instance-selection approaches, and stopping criteria. Furthermore, it is possible to formulate runtime discretization as an optimization problem, directly selecting the most discriminatory discretization technique rather than doing an empirical comparison.

A major shortcoming of our current approach is the lack of parallelization, selecting instances one at a time. Benchmarking on a computing cluster with n cores benefits from having batches of n instances. However, bigger batch sizes n impede $active\ learning$. Also, it is unclear how to synchronize model update and instance selection without wasting too much runtime.

On a more general note, one can apply our evaluation framework to arbitrary \mathcal{NP} -complete problems as all discussed active-learning methods are problem-agnostic. Solely the runtime-prediction model might need some tweaking. Those problems share most of the relevant properties of SAT solving, i.e., there are established problem-instance features, a full benchmark run takes days, and solver development traditionally requires expert knowledge to hand-select instances.

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