# Auto-Sklearn 2.0: Hands-free AutoML via Meta-Learning

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#### Abstract

Automated Machine Learning (AutoML) supports practitioners and researchers with the tedious task of designing machine learning pipelines and has recently achieved substantial success. In this paper we introduce new AutoML approaches motivated by our winning submission to the second ChaLearn AutoML challenge. We develop *PoSH Auto-sklearn*, which enables AutoML systems to work well on large datasets under rigid time limits using a new, simple and meta-feature-free meta-learning technique and employ a successful bandit strategy for budget allocation. However, *PoSH Auto-sklearn* introduces even more ways of running AutoML and might make it harder for users to set it up correctly. Therefore, we also go one step further and study the design space of AutoML itself, proposing a solution towards truly hands-free AutoML. Together, these changes give rise to the next generation of our AutoML system, *Auto-sklearn 2.0*. We verify the improvements by these additions in a large experimental study on 39 AutoML benchmark datasets and conclude the paper by comparing to other popular AutoML frameworks and *Auto-sklearn 1.0*, reducing the relative error by up to a factor of four, and yielding a performance in 10 minutes that is substantially better than what *Auto-sklearn 1.0* achieves within an hour.

**Keywords:** Automated Machine Learning, Hyperparameter Optimization, Meta-Learning

### 1. Introduction

The recent substantial progress in machine learning (ML) has led to a growing demand for hands-free ML systems that can support developers and ML novices in efficiently creating new ML applications. Since different datasets require different ML pipelines, this demand has given rise to the area of automated machine learning (AutoML; Hutter et al. (2019)). Popular AutoML systems, such as Auto-WEKA (Thornton et al., 2013), hyperopt-sklearn (Komer et al., 2014), Auto-sklearn (Feurer et al., 2015a), TPOT (Olson et al., 2016a) and Auto-Keras (Jin et al., 2019) perform a combined optimization across different preprocessors, classifiers or regressors and their hyperparameter settings, thereby reducing the effort for users substantially.

To assess the current state of AutoML and, more importantly, to foster progress in AutoML, ChaLearn conducted a series of AutoML challenges (Guyon et al., 2019), which

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evaluated AutoML systems in a systematic way under rigid time and memory constraints. Concretely, in these challenges, the AutoML systems were required to deliver predictions in less than 20 minutes, which would allow for an efficient development of new ML applications on-the-fly and help to democratize ML.

We won both the first and second AutoML challenge with modified versions of Auto-sklearn. In this work, we describe in detail how we improved Auto-sklearn from the first version (Feurer et al., 2015a) to construct PoSH Auto-sklearn, which won the second competition and then describe how we improved PoSH Auto-sklearn further to yield our current approach for Auto-sklearn 2.0.

Particularly, while AutoML reliefs the user from making low-level design decisions (e.g. which model to use), AutoML itself opens a myriad of high-level design decisions, e.g. which model selection strategy to use (Guyon et al., 2010, 2015; Raschka, 2018). Whereas our submissions to the AutoML challenges were mostly hand-designed, in this work we go one step further by automating AutoML itself to fully unfold the potential of AutoML in practice.<sup>1</sup>

After detailing the AutoML problem we consider in Section 2, we present two main parts making the following contributions:

- Part I: Portfolio Successive Halving in PoSH Auto-sklearn. In this part (see Section 3), we introduce budget allocation strategies as a complementary design choice to model selection strategies holdout (HO) and cross-validation (CV) for AutoML systems, also using successive halving (SH) as an alternative to allocate more resources to promising ML pipelines. Furthermore, we introduce both the practical approach as well as the theory behind building better portfolios for the meta-learning component of Auto-sklearn. We show that this combination substantially improves performance, yielding stronger results in 10 minutes than Auto-sklearn 1.0 achieved in 60 minutes.
- Part II: Automating AutoML in Auto-sklearn 2.0. In this part (see Section 4), we propose a meta-learning technique based on algorithm selection to construct a model-based policy selector to automatically choose the best optimization policy  $\pi$  for an AutoML system for a given dataset, further robustifying AutoML itself. We dub the resulting system Auto-sklearn 2.0 and depict the evolution from Auto-sklearn 1.0 via PoSH Auto-sklearn to Auto-sklearn 2.0 in Figure 1.

In Section 5, we additionally use the AutoML benchmark (Gijsbers et al., 2019) to evaluate Auto-sklearn 2.0 against other popular AutoML systems and show improved performance under rigid time constraints. Section 6 then puts our work into the context of related works and Section 7 concludes the paper with open questions, limitations and future work.

#### 2. Problem Statement and Background

AutoML is a widely used term, so, here we first define the problem we consider in this work. Then, we discuss the building blocks of AutoML systems before we provide an overview of existing AutoML methods and systems, including *Auto-sklearn*.

<sup>1.</sup> The work presented in this paper is in part based on two earlier workshop papers introducing some of the presented ideas in preliminary form (Feurer et al., 2018; Feurer and Hutter, 2018).

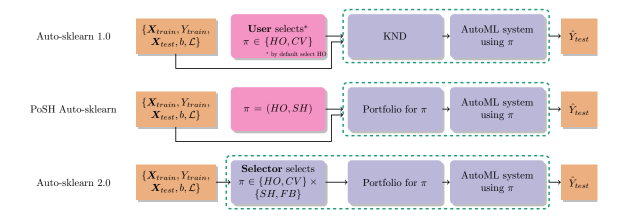


Figure 1: Schematic overview of Auto-sklearn 1.0, PoSH Auto-sklearn, and Auto-sklearn 2.0. Orange rectangular boxes refer to input and output data, while rounded purple boxes denote parts of the AutoML system (surrounded by a green dashed line). The pink, rounded box refers to human in the loop required for manual design decisions. The newer AutoML system simplify the usage of Auto-sklearn and reduce the required user input.

#### 2.1 Problem Statement

Let  $P(\mathcal{D})$  be a distribution of datasets from which we can sample an individual dataset's distribution  $P_d = P_d(\mathbf{x}, y)$ . The AutoML problem we consider is to generate a trained pipeline  $\mathcal{M}_{\lambda} : \mathbf{x} \mapsto y$ , hyperparameterized by  $\lambda \in \Lambda$  that automatically produces predictions for samples from the distribution  $P_d$  minimizing the expected generalization error:<sup>2</sup>

$$GE(\mathcal{M}_{\lambda}) = \mathbb{E}_{(\mathbf{x},y) \sim P_d} [\mathcal{L}(\mathcal{M}_{\lambda}(\mathbf{x}), y)].$$
 (1)

Since a dataset can only be observed through a set of n independent observations  $\mathcal{D}_d = \{(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_n, y_n)\} \sim P_d$ , we can only empirically approximate the generalization error on sample data:

$$\widehat{GE}(\mathcal{M}_{\lambda}, \mathcal{D}_d) = \frac{1}{n} \sum_{i=1}^n \mathcal{L}(\mathcal{M}_{\lambda}(\mathbf{x}_i), y_i).$$
 (2)

In practice we have access to two disjoint, finite samples which we from now on denote  $\mathcal{D}_{\text{train}}$  and  $\mathcal{D}_{\text{test}}$  ( $\mathcal{D}_{d,\text{train}}$  and  $\mathcal{D}_{d,\text{test}}$  in case we reference a specific dataset drawn from  $P_d$ ). For searching the best ML pipeline, we only have access to  $\mathcal{D}_{\text{train}}$ , however, in the end performance is estimated on  $\mathcal{D}_{\text{test}}$ .

AutoML systems use this to automatically search for the best  $\mathcal{M}_{\lambda^*}$ :

$$\mathcal{M}_{\lambda^*} \in \operatorname*{argmin} \widehat{GE}(\mathcal{M}_{\lambda}, \mathcal{D}_{\text{train}})$$

$$(3)$$

<sup>2.</sup> Our notation follows Vapnik (1991).

and estimate GE, e.g., by a K-fold cross-validation:

$$\widehat{GE}_{\text{CV}}(\mathcal{M}_{\lambda}, \mathcal{D}_{\text{train}}) = \frac{1}{K} \sum_{k=1}^{K} \widehat{GE}(\mathcal{M}_{\lambda}^{\mathcal{D}_{\text{train}}^{(\text{train},k)}}, \mathcal{D}_{\text{train}}^{(\text{val},k)})$$
(4)

where  $\mathcal{M}_{\boldsymbol{\lambda}}^{\mathcal{D}_{\mathrm{train}}^{(\mathrm{train},k)}}$  denotes that  $\mathcal{M}_{\boldsymbol{\lambda}}$  was trained on the training split of k-th fold  $\mathcal{D}_{\mathrm{train}}^{(\mathrm{train},k)} \subset \mathcal{D}_{\mathrm{train}}$ , and it is then evaluated on the validation split of the k-th fold  $\mathcal{D}_{\mathrm{train}}^{(\mathrm{val},k)} = \mathcal{D}_{\mathrm{train}} \setminus \mathcal{D}_{\mathrm{train}}^{(\mathrm{train},k)}$ . Assuming that, via  $\boldsymbol{\lambda}$ , an AutoML system can select both, an algorithm and its hyperparameter settings, this definition using  $\widehat{GE}_{\mathrm{CV}}$  is equivalent to the definition of the CASH (Combined Algorithm Selection and Hyperparameter optimization) problem (Thornton et al., 2013; Feurer et al., 2015a).

### 2.1.1 Time-bounded AutoML

In practice, users are not only interested to obtain an optimal pipeline  $\mathcal{M}_{\lambda^*}$  eventually, but have constraints on how much time and compute resources they are willing to invest. We denote the time it takes to evaluate  $\widehat{GE}(\mathcal{M}_{\lambda}, \mathcal{D}_{\text{train}})$  as  $t_{\lambda}$  and the overall optimization budget by T. Our goal is to find

$$\mathcal{M}_{\lambda^*} \in \underset{\lambda \in \Lambda}{\operatorname{argmin}} \widehat{GE}(\mathcal{M}_{\lambda}, \mathcal{D}_{\text{train}}) \text{ s.t.} \left(\sum t_{\lambda_i}\right) < T$$
 (5)

where the sum is over all evaluated pipelines  $\lambda_i$ , explicitly honouring the optimization budget T.

#### 2.1.2 Generalization of AutoML

Ultimately, an AutoML system  $\mathcal{A}: \mathcal{D} \mapsto \mathcal{M}^{\mathcal{D}}_{\lambda}$  should not only perform well on a single dataset but on the entire distribution over datasets  $P(\mathcal{D})$ . Therefore, the meta-problem of AutoML can be formalized as minimizing the generalization error over this distribution of datasets:

$$GE(\mathcal{A}) = \mathbb{E}_{\mathcal{D}_d \sim P(\mathcal{D})} \left[ \widehat{GE}(\mathcal{A}(\mathcal{D}_d), \mathcal{D}_d) \right],$$
 (6)

which in turn can again only be approximated by a finite set of meta-train datasets  $\mathbf{D}_{\text{meta}}$  (each with a finite set of observations):

$$\widehat{GE}(\mathcal{A}, \mathbf{D}_{\text{meta}}) = \frac{1}{|\mathbf{D}_{\text{meta}}|} \sum_{d=1}^{|\mathbf{D}_{\text{meta}}|} \widehat{GE}(\mathcal{A}(\mathcal{D}_d), \mathcal{D}_d).$$
 (7)

Having set up the problem statement, we can use this to further formalize our goals. Instead of using a single, fixed AutoML system  $\mathcal{A}$ , we will introduce optimization policies  $\pi$ , a combination of hyperparameters of the AutoML system and specific components to be used in a run, which can be used to configure an AutoML system for specific use cases. We then denote such a configured AutoML system as  $\mathcal{A}_{\pi}$ . We will first construct  $\pi$  manually in Section 3, introduce a novel system for designing  $\pi$  from data in Section 4

<sup>3.</sup> Alternatively, one could use holdout to estimate GE with  $\widehat{GE}_{HO}(\mathcal{M}_{\lambda}, \mathcal{D}_{train}) = \widehat{GE}(\mathcal{M}_{\lambda}^{\mathcal{D}_{train}^{train}}, \mathcal{D}_{train}^{val})$ .

and then extend this to a (learned) mapping  $\Xi:\mathcal{D}\to\pi$  which automatically suggests an optimization policy for a new dataset using algorithm selection. This problem setup can also be used to introduce generalizations of the algorithm selection problem such as algorithm configuration (Birattari et al., 2002; Hutter et al., 2009; Kleinberg et al., 2017), per-instance algorithm configuration (Xu et al., 2010; Malitsky et al., 2012) and dynamic algorithm configuration (Biedenkapp et al., 2020) on a meta-level; but we leave these for future work. Also, instead of selecting between multiple policies of a single AutoML system the presented method can be applied to choose between different AutoML without adjustments. However, our goal is to improve single AutoML systems to make them easier to use instead of maximizing performance by invocing many AutoML systems, thereby increasing the complexity.

# 3. Part I: Portfolio Successive Halving in PoSH Auto-sklearn

In this section we introduce our winning solution for the second AutoML competition (Guyon et al., 2019), PoSH Auto-sklearn, short for **PO**rtfolio Successive Halving. We first describe our use of portfolios to warmstart an AutoML system and then motivate the use of the successive halving bandit strategy. Next, we describe practical considerations for building PoSH Auto-sklearn. We end this first part of our main contributions by an experimental evaluation demonstrating the performance of PoSH Auto-sklearn.

# 3.1 Portfolio Building

Finding the optimal solution to the time-bounded optimization problem from Eq. (5) requires searching a large space of possible ML pipelines as efficiently as possible. BO is a strong approach for this, but in its vanilla version starts from scratch for every new problem. A better solution is to warmstart BO with ML pipelines that are expected to work well, as done in the k-nearest dataset (KND) approach of *Auto-sklearn 1.0* (Reif et al. (2012); Feurer et al. (2015b,a), see also the related work in Section 6.4.1). However, we found this solution to introduce new problems:

- 1. It is time consuming since it requires to compute meta-features describing the characteristics of datasets, where good meta-features are often quite expensive to compute.
- 2. It adds complexity to the system as the computation of the meta-features must also be done with a time and memory limit.
- 3. Many meta-features are not defined with respect to categorical features and missing values, making them hard to apply for most datasets.
- 4. It is not immediately clear which meta-features work best for which problem.
- 5. In the KND approach, there is no mechanism to guarantee that we do not execute redundant ML pipelines.

We indeed suffered from these issues in the first AutoML challenge, failing on one track due to running over time for the meta-feature generation, although we had already removed landmarking meta-features due to their potentially high runtime. Therefore, here we propose a meta-feature-free approach which does not warmstart with a set of configurations specific to a new dataset, but which uses a static portfolio – a set of complementary configurations that covers as many diverse datasets as possible and minimizes the risk of failure when facing a new task.

So, instead of evaluating configurations chosen *online* by the KND method, we construct a portfolio consisting of high-performing and complementary ML pipelines to perform well on as many datasets as possible *offline*. Then for a dataset at hand all pipelines in this portfolio are simply evaluated one after the other and if time left afterwards, we continue with pipelines suggested by BO warmstarted with the evaluated portfolio pipelines. The portfolio can thus be seen as an optimized initial design for the BO method.

In the following, we describe our offline procedure how to construct such a portfolio.

#### 3.1.1 Approach

We first describe how we construct a portfolio given a finite set of candidate pipelines  $\mathcal{C} = \{\lambda_1, ..., \lambda_l\}$  and later describe how we generate this set from our infinitely large configuration space. Additionally, we assume that there exists a set of datasets  $\mathbf{D}_{\text{meta}} = \{\mathcal{D}_1, ..., \mathcal{D}_{|\mathbf{D}_{\text{meta}}|}\}$  and we wish to build a portfolio  $\mathcal{P}$  consisting of a subset of the pipelines in  $\mathcal{C}$  that performs well on  $\mathbf{D}_{\text{meta}}$ . We outline the process to build such a portfolio in Algorithm 1. First, we initialize our portfolio  $\mathcal{P}$  to the empty set (Line 2). Then, we repeat the following procedure until  $|\mathcal{P}|$  reaches a pre-defined limit: From a set of candidate ML pipelines  $\mathcal{C}$ , we greedily add a candidate  $\lambda^* \in \mathcal{C}$  to  $\mathcal{P}$  that reduces the estimated generalization error over all meta-datasets most (Line 4), and then remove  $\lambda^*$  from  $\mathcal{C}$  (Line 5).

The estimated generalization error of a portfolio  $\mathcal{P}$  on a single dataset  $\mathcal{D}$  is the performance of the best pipeline  $\lambda \in \mathcal{P}$  on  $\mathcal{D}$  according to the model selection and budget allocation strategy. This can be described via a function  $S(\cdot,\cdot,\cdot)$ , which takes as input a function to compute the estimated generalization error (e.g., as defined in Equation 4), a set of machine learning pipelines to train, and a dataset. It then returns the pipeline with the lowest estimated generalization error as

$$\mathcal{M}_{\boldsymbol{\lambda}^*}^{\mathcal{D}} = S(\widehat{GE}, \mathcal{P}, \mathcal{D}) \in \underset{\mathcal{M}_{\boldsymbol{\lambda}}^{\mathcal{D}} \in \mathcal{P}}{\operatorname{argmin}} \widehat{GE}(\mathcal{M}_{\boldsymbol{\lambda}}^{\mathcal{D}}, \mathcal{D}). \tag{8}$$

In case argmin is not unique we return the model that was evaluated first. The estimated generalization error of  $\mathcal{P}$  across all meta-datasets  $\mathbf{D}_{\text{meta}} = \{\mathcal{D}_1, \dots, \mathcal{D}_{|\mathbf{D}_{\text{meta}}|}\}$  is then

$$\widehat{GE}_{S}(\mathcal{P}, \mathbf{D}_{\text{meta}}) = \sum_{d=1}^{|\mathbf{D}_{\text{meta}}|} \widehat{GE}\left(S\left(\widehat{GE}, \mathcal{P}, \mathcal{D}_{d}\right), \mathcal{D}_{d}^{\text{val}}\right), \tag{9}$$

Here, we give the equation for using holdout and in Appendix A we give the exact notation for cross-validation and successive halving.

Given a way how to construct the portfolio, we now detail how to construct the set of candidate pipelines  $\mathcal{C}$ . We limit ourselves to a finite set of portfolio candidates  $\mathcal{C}$  that we compute offline before performing Algorithm 1; we defer the use of other candidate generation mechanisms, for example the method of Hydra (Xu et al., 2010, 2011), to future work. To create our portfolio candidates, we first run BO on each meta-dataset in  $\mathbf{D}_{\text{meta}}$  and use the best found pipelines. Then, we evaluate each of these candidates on each meta-dataset to obtain a performance matrix which we use as a lookup table to construct

### Algorithm 1: Greedy Portfolio Building

```
    Input: Set of candidate ML pipelines C, D<sub>meta</sub> = {D<sub>1</sub>,...,D<sub>|D<sub>meta|</sub></sub>}, maximal portfolio size p, model selection strategy S
    P = ∅
    while |P| 
    λ* = argmin<sub>λ∈C</sub> GE<sub>S</sub>(P ∪ {λ}, D<sub>meta</sub>)
        // Ties are broken favoring the model trained first.
    P = P ∪ λ*, C = C \ {λ*}
    end while
    return Portfolio P
```

the portfolio. To build a portfolio across datasets, we need to take into account that the generalization errors for different datasets live on different scales (Bardenet et al., 2013). Thus, before taking averages, for each dataset we transform the generalization errors to the distance to the best observed performance scaled between zero and one, a metric named distance to minimum; which when averaged across all datasets is known as average distance to the minimum (ADTM) (Wistuba et al., 2015a, 2018). We compute the statistics for zero-one scaling by taking the results of all model selection and budget allocation strategies into account (i.e., we use the lowest observed test loss and the largest observed test loss for each meta-dataset).

For each meta-dataset  $\mathcal{D}_d \in \mathbf{D}_{\text{meta}}$  we have access to both  $\mathcal{D}_{d,\text{train}}$  and  $\mathcal{D}_{d,\text{test}}$ . In the case of holdout, we split the training set  $\mathcal{D}_{d,\text{train}}$  into two smaller disjoint sets  $\mathcal{D}_{d,\text{train}}^{\text{train}}$  and  $\mathcal{D}_{d,\text{train}}^{\text{val}}$ . We usually train models using  $\mathcal{D}_{d,\text{train}}^{\text{train}}$ , use  $\mathcal{D}_{d,\text{train}}^{\text{val}}$  to choose a ML pipeline  $\mathcal{M}_{\lambda}$  from the portfolio by means of the model selection strategy S, and judge the portfolio quality by the generalization loss of  $\mathcal{M}_{\lambda}$  on  $\mathcal{D}_{d,\text{train}}^{\text{val}}$  (instead of holdout we can of course also use cross-validation to compute the validation loss). However, if we instead choose the ML pipeline on the test set  $\mathcal{D}_{d,\text{test}}$ , Equation 9 becomes a monotone and submodular set function, which results in favorable guarantees for the greedy algorithm that we detail in Section 3.1.2. We follow this approach for the portfolio construction in the offline phase; we emphasize that for a new dataset, we of course do not require access to the test set  $\mathcal{D}_{\text{new,test}}$ .

#### 3.1.2 Theoretical Properties of the Greedy Algorithm

Besides the already mentioned practical advantages of the proposed greedy algorithm, this algorithm also enjoys a bounded worst-case error.

**Proposition 1** Minimizing the test loss of a portfolio  $\mathcal{P}$  on a set of datasets  $\mathcal{D}_1, \ldots, \mathcal{D}_{|\mathbf{D}_{meta}|}$ , when choosing an ML pipeline from  $\mathcal{P}$  for  $\mathcal{D}_d$  using holdout or cross-validation based on its performance on  $\mathcal{D}_{d,test}$ , is equivalent to the sensor placement problem for minimizing detection time (Krause et al., 2008).

We detail this equivalence in Appendix C.2. Thereby, we can apply existing results for the sensor placement problem to our problem. Using the test set of the meta-datasets  $\mathbf{D}_{\text{meta}}$  to construct a portfolio is perfectly fine as long as we do not use the meta-test datasets  $\mathbf{D}_{\text{test}}$ .

Corollary 1 The penalty function for all meta-datasets is submodular.

We can directly apply the proof from Krause et al. (2008) that the so-called penalty function (maximum estimated generalization error minus the observed estimated generalization error) is submodular and monotone to our problem setup. Since linear combinations of submodular functions are also submodular (Krause and Golovin, 2014), the penalty function is also submodular.

**Corollary 2** The problem of finding an optimal portfolio  $\mathcal{P}^*$  is NP-hard (Nemhauser et al., 1978; Krause et al., 2008).

**Corollary 3** The portfolio obtained by the greedy algorithm reduces the penalty (as defined in the explanation of Corollary 1) such that  $R(\mathcal{P}^*) \geq R(\mathcal{P}) \geq (1 - \frac{1}{e})R(\mathcal{P}^*)$ , where R is the expected penalty reduction across all datasets.

This means that there is at most an additional ADTM 37% of what the best possible portfolio  $\mathcal{P}^*$  of size  $|\mathcal{P}|$  would achieve (Nemhauser et al., 1978; Krause and Golovin, 2014). A generalization of this result given by Krause and Golovin (2014) also tighten the bound to 1% of what the optimal portfolio  $\mathcal{P}^*$  of size  $|\mathcal{P}^*|$  would achieve by extending the portfolio constructed by the greedy algorithm to size  $5|\mathcal{P}|$ . This means that we can find a close-to-optimal portfolio on the meta-train datasets  $\mathbf{D}_{\text{meta}}$ . Under the assumption that we apply the portfolio to datasets from the same distribution of datasets, we have a strong set of default ML pipelines.

We can also apply other strategies for the sensor set placement in our problem setting, such as mixed integer programming strategies, which can solve it optimally; however, these do not scale to portfolio sizes of a dozen ML pipelines (Krause et al., 2008; Pfisterer et al., 2018).

The same proposition (with the same proof) and corollaries apply if we select a ML pipeline based on an intermediate step in a learning curve or use cross-validation instead of holdout. We discuss using the validation set and other model selection and budget allocation strategies in Appendix C.3 and Appendix C.4.

# 3.2 Budget Allocation using Successive Halving

Two key components of any efficient AutoML system are its model selection and budget allocation strategies, which serve the following purposes: 1) approximate the generalization error of a single ML pipeline and 2) decide how many resources to allocate for each pipeline evaluation. The first point is typically tackled by using a holdout set or K-fold cross-validation (see Section 6.4.1). Here, we focus on the second point.

A key issue we identified during the last AutoML challenge was that training expensive configurations on the complete training set, combined with a low time budget, does not scale well to large datasets. At the same time, we noticed that our (then manual) strategy to run predefined pipelines on subsets of the data already yielded predictions good enough for ensemble building.

This questions the common choice of assigning to all pipeline evaluations the same amount of resources, i.e. time, compute and data. As a principled alternative we used the successive halving bandit (SH, Karnin et al. (2013); Jamieson and Talwalkar (2016)), which assigns more budget to promising machine learning pipelines and can easily be combined with iterative algorithms.

### 3.2.1 Approach

AutoML systems evaluate each pipeline under the same resource limitations and on the same budget (e.g., number of iterations using iterative algorithms). To increase efficiency for cases with tight resource limitations, we suggest to allocate more resources to promising pipelines by using SH (Karnin et al., 2013; Jamieson and Talwalkar, 2016) to aggressively prune poor-performing pipelines.

Given a minimal and maximal budget per ML pipeline, SH starts by training a fixed number of ML pipelines for the smallest budget. Then, it iteratively selects  $\frac{1}{\eta}$  of the pipelines with lowest generalization error, multiplies their budget by  $\eta$ , and re-evaluates. This process is continued until only a single ML pipeline is left or the maximal budget is spent. While SH itself chooses new pipelines  $\mathcal{M}_{\lambda}$  to evaluate at random, we follow work combining SH with BO (Falkner et al., 2018). Specifically, we use BO to iteratively suggest new ML pipelines  $\mathcal{M}_{\lambda}$ , which we evaluate on the lowest budget until a fixed number of pipelines has been evaluated. Then, we run SH as described above. The implementation by Falkner et al. (2018) builds the model for BO on the highest available budget when there are n+1 finished pipelines where n is the number of hyperparameters to be optimized. As this is a mathematical requirement by their model, we can reduce this number and set it to  $\frac{n}{2}$ . Since we use only iterative ML models in our configuration space, we rely on the number of iterations as the budget.

SH potentially provides large speedups, but it could also too aggressively cut away good configurations that need a higher budget to perform best. Thus, we expect SH to work best for large datasets, for which there is not enough time to train many ML pipelines for the full budget, but for which training a ML pipeline on a small budget already yields a good indication of the generalization error.

We note that SH can be used in combination with both, holdout or cross-validation, and thus indeed adds another hyper-hyperparameter to the AutoML system, namely whether to use SH or always evaluate the algorithm on the full budget (FB). However, it also adds more flexibility to tackle a broader range of problems.

# 3.3 Practical Considerations and Challenge Results

In order to make best of the given budgets and the successive halving algorithm we had to do certain adjustments to obtain high performance.

First, we restricted the search space to contain only iterative algorithms and no more feature preprocessing. This simplifies the usage of SH as we only have to deal with a single type of fidelity, the number of iterations, while we would otherwise have to also consider dataset subsets as an alternative. This leaves us with extremely randomized trees (Geurts et al., 2006), random forests (Breimann, 2001), histogram-based gradient boosting (Friedman, 2001; Ke et al., 2017), a linear model fitted with a passive aggressive algorithm (Crammer et al., 2006) or stochastic gradient descent and a multi-layer perceptron. The exact configuration space can be found in Table 16 of the appendix.

<sup>4.</sup> Falkner et al. (2018) proposed using Hyperband (Li et al., 2018) together with BO, however, we use only SH as we expect it to work better in the extreme of having very little time, as it more aggressively reduces the budget per ML pipeline.

Second, because of using only iterative algorithms, we are able to store partially fitted models to disk to prevent having no predictions in case of time- and memouts. That is, after 2, 4, 8, . . . iterations we make predictions for the validation set and dump the model for later usage.

We give further details, such as the restricted search space, in Appendix B. For our submission to the second AutoML challenge, we implemented the following safeguards and tricks (Feurer et al., 2018), which we do not use in this paper since we instead focus on automatically designing a robust AutoML system:

- For the submission we also employed support vector machines using subsets of the dataset as lower fidelities. As none of the support vector machines was chosen for a final ensemble in the competition, we did not consider them any more for this paper, simplifying our methodology.
- We developed an additional library pruning method for ensemble selection. However, in preliminary experiments, we found that this at most provides an unsignificant boost for area under curve and not balanced accuracy, which we use in this work and thus did not follow that any further.
- To increase robustness against arbitrarily large datasets, we reduced all datasets to have at most 500 features using univariate feature selection. Similarly, we also reduced all datasets to have at most 45 000. We do not think these are good strategies in general and only implemented them as we had no information about the dimensionality of the datasets used in the challenge and to prevent running out of time and memory.
- In case the datasets had less than 1000 datapoints, we would have reverted from holdout to cross-validation. This fallback was however not triggered due to the datasets being larger.
- We manually added a linear regression fitted with stochastic gradient descent with its hyperparameters optimized for fast runtime as the first entry in the portfolio to maximize the chances of fitting a model within the given time.

Our submission, *PoSH Auto-sklearn*, was the overall winner of the second AutoML challenge. We give the results of the competition in Table 1 and refer to Feurer et al. (2018) and Guyon et al. (2019) for further details, especially for information on our competitors.

# 3.4 Experimental Setup

So far, AutoML systems were designed without any optimization budget or with a single, fixed optimization budget T in mind (see Equation 5).<sup>5</sup> Our system takes the optimization budget into account when constructing the portfolio. We will study two optimization budgets: a short, 10 minute optimization budget and a long, 60 minute optimization budget as in the original Auto-sklearn paper. To have a single metric for binary classification, multiclass classification and unbalanced datasets, we report the balanced error

<sup>5.</sup> The OBOE AutoML system (Yang et al., 2019) is a potential exception that takes the optimization budget into consideration, but the experiments by Yang et al. (2019) were only conducted for a single optimization budget, not demonstrating that the system adapts itself to multiple optimization budgets.

Name	Rank	Dataset #1	Dataset #2	Dataset #3	Dataset #4	Dataset #5
PoSH Auto-sklearn	2.8	0.5533(3)	0.2839(4)	<b>0.3932</b> ( <b>1</b> )	<b>0.2635</b> ( <b>1</b> )	0.6766(5)
narnars0	3.8	0.5418(5)	0.2894(2)	0.3665(2)	0.2005(9)	<b>0.6922</b> ( <b>1</b> )
Malik	5.4	0.5085(7)	0.2297(7)	0.2670(6)	0.2413(5)	0.6853(2)
wlWangl	5.4	$\mathbf{0.5655(2)}$	<b>0.4851</b> ( <b>1</b> )	0.2829(5)	-0.0886(16)	0.6840(3)
thanhdng	5.4	0.5131(6)	0.2256(8)	0.2605(7)	0.2603(2)	0.6777(4)

Table 1: Results for the second AutoML challenge (Guyon et al., 2019). Name is the team name, Rank the final rank of the submission, followed by the individual results on the five datasets used in the competition. All performances are the normalized area under the ROC curve (Guyon et al., 2015) with the per-dataset rank in brackets.

rate (1 – balanced accuracy), following the 1<sup>st</sup> AutoML challenge (Guyon et al., 2019). As different datasets can live on different scales, we apply a linear transformation to obtain comparable values. Concretely, we obtain the minimal and maximal error obtained by executing *Auto-sklearn* with portfolios and using ensembles for each combination of model selection and budget allocation strategies per dataset, and rescale by subtracting the minimal error and dividing by the difference between the maximal and minimal error (ADTM, as introduced in Section 3.1.1).<sup>6</sup> With this transformation, we obtain a normalized error which can be interpreted as the regret of our method.

We also limit the time and memory for each ML pipeline evaluation. For the time limit we allow for at most 1/10 of the optimization budget, while for the memory we allow the pipeline 4GB before forcefully terminating the execution.

#### 3.4.1 Datasets

We require two disjoint sets of datasets for our setup: (i)  $\mathbf{D}_{\text{meta}}$ , on which we build portfolios and the model-based policy selector that we will introduce in Section 4, and (ii)  $\mathbf{D}_{\text{test}}$ , on which we evaluate our method. The distribution of both sets ideally spans a wide variety of problem domains and dataset characteristics. For  $\mathbf{D}_{\text{test}}$ , we rely on 39 datasets selected for the AutoML benchmark proposed by Gijsbers et al. (2019), which consists of datasets for comparing classifiers (Bischl et al., 2019) and datasets from the AutoML challenges (Guyon et al., 2019).

We collected the meta datasets  $\mathbf{D}_{\text{meta}}$  based on OpenML (Vanschoren et al., 2014) using the OpenML-Python API (Feurer et al., 2021). To obtain a representative set, we considered all datasets on OpenML with more than 500 and less than 1000000 samples with at least two attributes. Next, we dropped all datasets that are sparse, contain time attributes or string type attributes as  $\mathbf{D}_{\text{test}}$  does not contain any such datasets. Then, we dropped synthetic datasets and subsampled clusters of highly similar datasets. Finally, we manually checked for overlap with  $\mathbf{D}_{\text{test}}$  and ended up with a total of 208 training datasets and used them to train our method.

<sup>6.</sup> We would like to highlight that this is slightly different than in Section 3.1.1 where we did not have access to the ensemble performance, and also only normalized per model selection strategy.

We show the distribution of the datasets in Figure 2. Green points refer to  $\mathbf{D}_{\text{meta}}$  and orange crosses to  $\mathbf{D}_{\text{test}}$ . We can see that  $\mathbf{D}_{\text{meta}}$  spans the underlying distribution of  $\mathbf{D}_{\text{test}}$  quite well, but that there are several datasets which are outside of the  $\mathbf{D}_{\text{meta}}$  distribution, which are marked with a black cross and for which our AutoML system selected a backup strategy (see Section 4.1.1). We give the full list of datasets for  $\mathbf{D}_{\text{meta}}$  and  $\mathbf{D}_{\text{test}}$  in Appendix E.

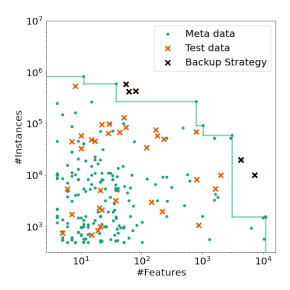


Figure 2: Distribution of meta and test datasets. We visualize each dataset w.r.t. its metafeatures and highlight the datasets that lie outside our meta distribution; for these, we apply a backup strategy.

For all datasets we use a single holdout test set of 33.33% which is defined by the corresponding OpenML task. The remaining 66.66% are the training data of our AutoML systems, which handle further splits for model selection themselves based on the chosen model selection strategy.

#### 3.4.2 Meta-data Generation

For each optimization budget we created four performance matrices, see Section 3.1.1 for details on performance matrices. Each matrix refers to one way of assessing the generalization error of a model: holdout, 3-fold CV, 5-fold CV or 10-fold CV. To obtain each matrix, we did the following. For each dataset  $\mathcal{D}$  in  $\mathbf{D}_{\text{meta}}$ , we used combined algorithm selection and hyperparameter optimization to find a customized ML pipeline. In practice, we ran SMAC (Hutter et al., 2011; Lindauer et al., 2017a) three times and picked the best resulting ML pipeline on the test split of  $\mathcal{D}$ . To ensure that SMAC finds a good configuration, we run it for ten times the *optimizationbudget* given by the user (see Equation 5). Then, we ran the cross-product of all ML pipelines and datasets to obtain the performance matrix.

We also stored intermediate results for the iterative algorithms so that we can build custom portfolios for SH, too.

#### 3.4.3 Other Experimental Details

We always report results averaged across 10 repetitions to account for randomness and report the mean and standard deviation over these repetitions. To check whether performance differences are significant, where possible, we ran the Wilcoxon signed rank test as a statistical hypothesis test with  $\alpha=0.05$  (Demšar, 2006). In addition, we plot the average rank as follows. For each dataset, we draw one run per method (out of 10 repetitions) and rank these draws according to performance, using the average rank in case of ties. We repeat this sampling 200 times to obtain the average rank on a dataset, before averaging these into the total average.

We conducted all experiments using ensemble selection, and we construct ensembles of size 50 with replacement. We give results without ensemble selection in the Appendix B.2.

All experiments were conducted on a compute cluster with machines equipped with 2 Intel Xeon Gold 6242 CPUs with 2.8GHz (32 cores) and 192 GB RAM, running Ubuntu 20.04.01. We provide scripts for reproducing all our experimental results at https://github.com/automl/ASKL2.0\_experiments and provide a clean integration of our methods into the official Auto-sklearn repository.

# 3.5 Experimental Results

In this subsection we now validate the improvements for *PoSH Auto-sklearn*. First, we will qualitatively show that different model selection strategies are required for different dataset sizes and time horizons. Then, we will compare the performance of using a portfolio to the previous KND approach and no warmstarting.

#### 3.5.1 Using different model selection strategies

Choosing the correct model selection and budget allocation strategy not only depends on the characteristics of the dataset at hand, but also on the given time-limit. While there exist general recommendations, we observed in practice that this is a crucial design decision that drastically impacts performance. To highlight this effect, in Figure 3 we show exemplary results comparing the balanced error rates of the best found ML pipeline found by searching our configuration space with BO using holdout, 3CV, 5CV and 10CV with SH and FB on different optimization budgets and datasets. We give details on the SH hyperparameters in Appendix D.3.

The top row shows results obtained using the same optimization budget of 10 minutes on two different datasets. While FB; 10CV is best on dataset sylvine (top left) the same strategy on median performs amongst the worst strategies on dataset adult (top right). Also, on sylvine, SH performs overall slightly worse in contrast to adult, where SH performs better on average. The bottom rows shows how the given time-limit impacts the performance on the dataset jungle\_chess\_2pcs\_raw\_endgame\_complete. Using a quite restrictive optimization budget of 10 minutes (bottom left), SH; 3CV, which aggressively cuts ML pipelines on lower budgets, performs best on average. With a higher optimization budget (bottom right), the

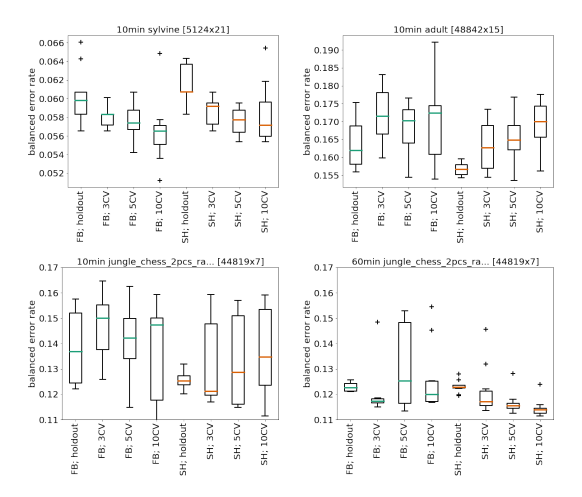


Figure 3: Final performance for BO using different model selection strategies averaged across 10 repetitions. Top row: Results for a optimization budget of 10 minutes on two different datasets. Bottom row: Results for a optimization budget of 10 and 60 minutes on the same dataset.

overall results improve and more strategies become competitive. We will demonstrate in Section 4.3 that using both, FB and SH, has advantages over using only one of them.

# 3.5.2 Using Portfolios instead of a k-nearest datasets approach

**Portfolio vs. KND.** We introduced the portfolio-based warmstarting to avoid computing meta-features for a new dataset. However, the portfolios work inherently differently. While KND aimed at using only well performing configurations, a portfolio is built such that there is a diverse set of configurations, starting with ones that perform well on average and then moving to more specialized ones, which also provides a different form of initial design for BO. Here, we study the performance of the learned portfolio and compare it against *Auto-sklearn 1.0*'s default meta-learning strategy using 25 configurations. Additionally, we also study how pure BO would perform. We give results in Table 2.

	1	0 minut	es	60 minutes			
	ВО	KND	Port	ВО	KND	Port	
holdout	5.98	5.29	3.70	3.84	3.98	3.08	
SH; holdout	5.15	4.82	4.11	3.77	3.55	3.19	
3CV	8.52	7.76	6.90	6.42	6.31	4.96	
SH; 3CV	7.82	7.67	6.16	6.08	5.91	5.17	
$5\mathrm{CV}$	9.48	9.45	7.93	6.64	6.47	5.05	
SH; 5CV	9.48	8.85	7.05	6.19	5.83	5.40	
10CV	16.10	<u>15.11</u>	12.42	10.82	10.44	9.68	
SH; 10CV	16.14	15.10	12.61	10.54	10.33	9.23	

Table 2: Averaged normalized balanced error rate. We report the aggregated performance across 10 repetitions and 39 datasets of our AutoML system using only Bayesian optimization (BO), or BO warmstarted with k-nearest-datasets (KND) or a greedy portfolio (Port). Per line, we boldface the best mean value (per model selection and budget allocation strategy and optimization budget, and underline results that are not statistically different according to a Wilcoxon-signed-rank Test  $(\alpha = 0.05)$ ).

For the new AutoML-hyperparameter  $|\mathcal{P}|$  we chose 32 to allow two full iterations of SH with our hyperparameter setting of SH. Unsurprisingly, warmstarting in general improves the performance on all optimization budgets and most model selection strategies, often by a large margin. The portfolios always improve over BO, while KND does so in all but one case. When comparing the portfolios to the KND approach, we find that the raw results are always favorable, and that for more than half of the settings the differences are also significant.

**PoSH Auto-sklearn vs Auto-sklearn 1.0.** Lastly, we can have a look at the performance of PoSH Auto-sklearn (SH; holdout and Port) compared to Auto-sklearn 1.0 (holdout and KND). For both time horizons there is a strong reduction in the loss (10min:  $5.29 \rightarrow 4.11$  and 60min:  $3.98 \rightarrow 3.19$ ), indicating that the proposed PoSH Auto-sklearn is indeed an improvement over the existing solution. In combination with the portfolio, the average results are inconclusive about whether our use of successive halving was the right choice or whether plain holdout would have been better. We also provide the raw numbers in Appendix B.3, but they are inconclusive, too.

### 4. Part II: Automating Design Decisions in AutoML

The goal of AutoML is to yield state-of-the-art performance without requiring the user to make low-level decisions, e.g., which model and hyperparameter configurations to apply. Using a portfolio and SH, PoSH Auto-sklearn is already an improvement over Auto-sklearn 1.0 in terms of efficiency and scalability. However, high-level design decisions, such as choosing between cross-validation and holdout or whether to use SH or not, remain. Thus, PoSH Auto-sklearn, and AutoML systems in general, suffer from a similar problem as they are trying to solve, as users have to manually set their arguments. Thus, we propose to ex-

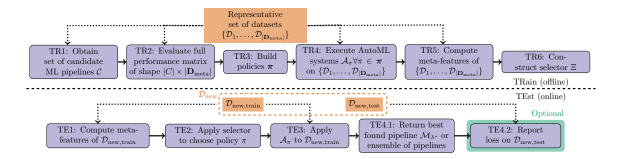


Figure 4: Schematic overview of the proposed *Auto-sklearn 2.0* system with the training phase (TR1-TR6) above and the test phase (TE1-TE4) below the dashed line. Rounded boxes refer to computational steps while rectangular boxes depict the input data to the AutoML system.

tend AutoML systems with a policy selector to automatically choose an optimization policy given a dataset (see Figure 1 in Section 1 for a schematic overview). In this second part, we discuss the resulting approach, which led to *Auto-sklearn 2.0* as the first implementation of it.

### 4.1 Automated Policy Selection

Specifically, we consider the case, where an AutoML system can be run with different optimization policies  $\pi \in \Pi$  and study how to further automate AutoML using algorithm selection on this meta-meta level. In practice, we extend the formulation introduced in Eq. 7 to not use an AutoML system  $\mathcal{A}_{\pi}$  with a fixed policy  $\pi$ , but to contain a policy selector  $\Xi : \mathcal{D} \to \pi$ :

$$\widehat{GE}(\mathcal{A}, \Xi, \mathbf{D}_{\text{meta}}) = \frac{1}{|\mathbf{D}_{\text{meta}}|} \sum_{d=1}^{|\mathbf{D}_{\text{meta}}|} \widehat{GE}(\mathcal{A}_{\Xi(\mathcal{D}_d)}(\mathcal{D}_d), \mathcal{D}_d).$$
(10)

In the remainder of this section, we describe how to construct such a policy selector.

# 4.1.1 Approach

Optimization strategies in AutoML itself are often heavily hyperparameterized. In our case, we deem the model selection strategy and budget allocation strategy (see Sections 6.4.1 and 3.2) as important choices the user has to make when using an AutoML system to obtain high performance. These decisions depend on both the given dataset and the available resources. As there is also an interaction between the two strategies and the optimal portfolio  $\mathcal{P}$ , we consider here that the optimization policy  $\pi$  is parameterized by a combination of (i) model selection strategy, (ii) budget allocation strategy and (iii) a portfolio constructed for the choice of the two strategies. In our case these are eight different policies ({3-fold CV, 5-fold CV, 10-fold CV, holdout} × {SH, FB}).

We introduce a new layer on top of AutoML systems that automatically selects a policy  $\pi$  for a new dataset. We show an overview of this system in Figure 4 which consists of a training (TR1–TR6) and a testing stage (TE1–TE4). In brief, in training steps TR1–

TR3, we obtain a performance matrix of size  $|\mathcal{C}| \times |\mathbf{D}_{meta}|$ , where  $\mathcal{C}$  is a set of candidate ML pipelines, and  $|\mathbf{D}_{meta}|$  is the number of representative meta-datasets. Having collected datasets  $\mathbf{D}_{meta}$  (we describe in Section 3.4.1 how we did this for our experiments), we obtain the candidate ML pipelines (TR1) by running Auto-sklearn on each dataset and build the performance matrix as described in Section 3.1.1 by evaluating each of these candidate pipelines on each dataset (TR2) as described in Section 3.4.2. This matrix is used to build a portfolio as described in Section 3.1 for each combination of model selection and budget allocation strategy in training step TR3. These combinations of portfolio, model selection strategy and budget allocation strategy are our policies. We then execute the full AutoML system for each such policy in step TR4 to obtain a realistic performance estimate. In step TR5 we compute meta-features and use them together with the performance estimate from TR4 in step TR6 to train a model-based policy selector  $\Xi$ , which will be used in the online test phase.

In order to not overestimate the performance of  $\pi$  on a dataset  $\mathcal{D}_d$ , dataset  $\mathcal{D}_d$  must not be part of the meta-data for constructing the portfolio. To overcome this issue, we perform an inner 5-fold cross-validation and build each  $\pi$  on four fifths of the meta-datasets and evaluate it on the left-out fifth of meta-datasets. For the final AutoML system we then use a portfolio built on all meta-datasets.

For a new dataset  $\mathcal{D}_{\text{new}} \in \mathbf{D}_{\text{test}}$ , we first compute meta-features describing  $\mathcal{D}_{\text{new}}$  (TE1) and use the model-based policy selector from step TR7 to automatically select an appropriate policy for  $\mathcal{D}_{\text{new}}$  based on the meta-features (TE2). This will relieve users from making this decision on their own. Given an optimization policy  $\pi$ , we then apply the AutoML system  $\mathcal{A}_{\pi}$  to  $\mathcal{D}_{\text{new}}$  (TE3). Finally, we return the best found pipeline  $\mathcal{M}_{\lambda*}$  based on the training set of  $\mathcal{D}_{\text{new}}$  (TE4.1). Optionally, we can then compute the loss of  $\mathcal{M}_{\lambda*}$  on the test set of  $\mathcal{D}_{\text{new}}$  (TE4.2); we emphasize that this would be the only time we ever access the test set of  $\mathcal{D}_{\text{new}}$ .

In the following, we describe two ways to construct a policy selector and introduce an additional backup strategy to make it robust towards failures.

Constructing the single best policy A straight-forward way to construct a selector relies on the assumption that the meta-datasets  $\mathbf{D}_{meta}$  are homogeneous and that a new dataset is similar to these. In such a case we can use per-set algorithm selection (Kerschke et al., 2019), which aims to find the best algorithm for a set of problem instances; in our case it aims to find the best combination of model selection and budget allocation for a given set of meta-datasets  $\mathbf{D}_{meta}$ . This single best policy is then the automated replacement for our manual selection of SH and holdout in PoSH Auto-sklearn. While this seems to be a trivial baseline, it actually requires the same amount of compute power as the more elaborate strategy we introduce next.

Constructing the per-dataset Policy Selector Instead of using a fixed, learned policy, we now propose to adapt the policy to the dataset at hand. To construct the meta selection model (TR6), we follow the policy selector design of HydraMIP (Xu et al., 2011): for each pair of AutoML policies, we fit a random forest to predict whether policy  $\pi_A$  outperforms policy  $\pi_B$  given the current dataset's meta-features. Since the misclassification loss depends on the difference of the losses of the two policies (i.e. the ADTM when choosing the wrong policy), we weight each meta-observation by their loss difference. To make errors comparable

hyperparameter	type	values
Min. number of samples to create a further split	int	[3, 20]
Min. number of samples to create a new leaf	int	[2, 20]
Max. depth of a tree	int	[0, 20]
Max. number of features to be used for a split	int	[1, 2]
Bootstrapping in the random forest	$\operatorname{cat}$	$\{yes, no\}$
Soft or hard voting when combining models	$\operatorname{cat}$	$\{soft, hard\}$
Error value scaling to compute dataset weights	$\operatorname{cat}$	see text

Table 3: configuration space of the model-based policy selector.

across different datasets (Bardenet et al., 2013), we scale the individual error values for each dataset. At test time (TE2), we query all pairwise models for the given meta-features, and use voting to choose a policy  $\pi$ . We will refer to this strategy as the *Policy Selector*.

To improve the performance of the model-based policy selector, we applied BO to optimize the model-based policy selector's hyperparameters to minimize the cross-validation error (Lindauer et al., 2015). We optimized in total seven hyperparameters, five of which are related to the random forest, one is how to combine the pairwise models to get a prediction and the last one is the strategy of how to scale error values to compute weights for comparing datasets, i.e. using the raw observations, scale with [min, max] / [min, 1] across a pair or all policies or use the difference in ranks as the weight (see Table 3). Hyperparameters are shared between all pairwise models to avoid factorial growth of the number of hyperparameters with the number of new model selection strategies.

Meta-Features. To train our model-based policy selector and to select a policy, as well to use the backup strategy, we use meta-features (Brazdil et al., 2008; Vanschoren, 2019) describing all meta-train datasets (TR5) and new datasets (TE1). To avoid the problems discussed in Section 3.1 we only use very simple and robust meta-features, which can be reliably computed in linear time for every dataset: 1) the number of data points and 2) the number of features. In fact, these are most often already stored as meta-data for the data structure holding the dataset. In our experiments we will show that even with only these trivial and cheap meta-features we can substantially improve over a static policy.

Backup strategy. Since there is no guarantee that our model-based policy selector will extrapolate well to datasets outside of the meta-datasets, we implement a fallback measure to avoid failures. Such failures can be harmful if a new dataset is, e.g., much larger than any dataset in the meta-dataset and the model-based policy selector proposes to use a policy that would time out without any solution. More specifically, if there is no dataset in the meta-datasets that has higher or equal values for each meta-feature (i.e. dominates the dataset meta-features), our system falls back to use holdout with SH, which is the most aggressive and cheapest policy we consider.

### 4.2 Experimental Results

To study the performance of the policy selector, we compare it to *PoSH Auto-sklearn* as described in Section 4 and *Auto-sklearn* 1.0. From now on we refer to *PoSH Auto-sklearn* 

	10MIN	60MIN
	$\emptyset$ std	$\emptyset$ std
(1) Auto-sklearn $(2.0)$	<b>3.58</b> 0.23	<b>2.47</b> 0.18
(2) PoSH-Auto-sklearn	$4.11 \ 0.09$	$3.19 \ 0.12$
(3) Auto-sklearn $(1.0)$	$16.21 \ 0.27$	$7.17 \ 0.30$

Table 4: Final performance of Auto-sklearn 2.0, PoSH Auto-sklearn and Auto-sklearn 1.0. We report the normalized balanced error rate averaged across 10 repetitions on 39 datasets. We boldface the best mean value (per optimization budget) and underline results that are not statistically different according to a Wilcoxon-signed-rank Test ( $\alpha = 0.05$ ).

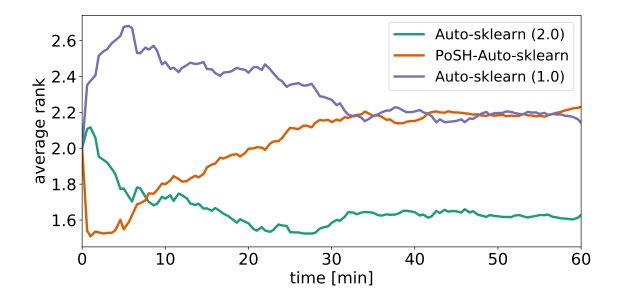


Figure 5: Performance over time. We report the normalized BER and the rank over time averaged across 10 repetitions and 39 datasets comparing our system to our previous AutoML systems.

equipped with a policy selector as *Auto-sklearn 2.0*. As before, we study two horizons, 10 minutes and 60 minutes, and use versions of *Auto-sklearn 2.0* tuned specifically for these time horizons.

Looking at Table 4, we see that Auto-sklearn 2.0 achieves the lowest error, being significantly better for both optimization budgets. Most notably, Auto-sklearn 2.0 reduces the relative error compared to Auto-sklearn 1.0 by 78% (10MIN) and 65%, respectively, which means a reduction by a factor of four and three.

It turns out that these results are skewed by several large datasets (task IDs 189873 and 75193) for both horizons; 189866, 189874, 168796 and 168797 only for the ten minutes

horizon) on which the KND initialization of *Auto-sklearn 1.0* only suggests ML pipelines that time out or hit the memory limit and thus exhaust the optimization budget for the full configuration space. Our new AutoML system does not suffer from this problem as it a) selects SH to avoid spending too much time on unpromising ML pipelines and b) can return predictions and results even if a ML pipeline was not evaluated for the full budget or converged early; and even after removing the datasets in question from the average, the performance of *Auto-sklearn 1.0* is substantially worse than that *Auto-sklearn 2.0*.

When looking at the intermediate system, i.e. *PoSH Auto-sklearn*, we find that it outperforms *Auto-sklearn 1.0* in terms of the normalized balanced error rate, but that the additional step of selecting the model selection and budget allocation strategy gives *Auto-sklearn 2.0* an edge. When not considering the large datasets *Auto-sklearn 1.0* failed on, their performance becomes very similar.

Figure 5 provides another view on the results, presenting average ranks (where failures obtain less weight compared to the averaged performance). Auto-sklearn 2.0 is still able to deliver best results, PoSH Auto-sklearn should be preferred to Auto-sklearn 1.0 for the first 30 minutes and then converges to the roughly the same ranking.

#### 4.3 Ablation

Now, we study the contribution of each of our improvements in an ablation study. We iteratively disable one component and compare the performance to the full system. These components are (1) using a per-dataset model-based policy selector to choose a policy, (2) using only a subset of the available policies, and (3) warmstarting BO with a portfolio.

#### 4.3.1 Do we need per-dataset selection?

We first examine how much performance we gain by having a model-based policy selector to decide between different AutoML strategies based on meta-features and how to construct this model-based policy selector, or whether it is sufficient to select a single strategy based on meta-training datasets. We compare the performance of the full system using a model-based policy selector to using (1) a single, static strategy (single best) and (2) both, the model-based policy selector and the single best, without the fallback mechanism for out-of-distribution datasets and give all results in Table 5. We also provide two further baselines. First, a random baseline, which randomly assigns a policy to a run. Second, an oracle baselines, which marks the lowest possible error that we practically achieved by a model.<sup>7</sup>

First, we compare the performance of the model-based policy selector with the single best. We can observe that for 10 minutes there is a slight improvement in terms of performance, while the performance for 60 minutes is almost equal. While there is no significant difference to the single best for 10 minutes, there is for 60 minutes. These numbers can be compared with Table 2 to see how we fare against picking a single policy by hand. We find that our proposed algorithm selection compares favorably, especially for the longer time horizon.

<sup>7.</sup> We would like to note that the oracle performance can be unequal to zero, because we normalize the results using the single best test loss found for a single model to normalize the results. When evaluating the best policy on a dataset this most likely results in selecting a model on the validation set that is not the single best model on the test set we use to normalize data.

	$10 \mathrm{\ Min}$			
oracle	2.33	1.22		
random	8.32	6.18		
trained on	P P+BO P+BO+E	P P+BO P+BO+E		
single best	$3.88 \ \ \underline{3.67} \ \ \ \ \underline{3.69}$	2.49 2.38 2.44		
single best w/o fallback	$5.18 \ \underline{6.38} \ \underline{6.40}$	5.10 5.01 5.07		
model-based policy selector	3.58 $3.56$ $3.58$	$2.53 \ \underline{2.32} \ \ \underline{2.47}$		
model-based policy selector w/o fallback	5.43  5.68  4.79	$4.98 \ 5.36 \ \underline{5.43}$		

Table 5: Final performance (averaged normalized balanced error rate) for 10 and 60 minutes. We report the theoretical best results (oracle) and results for choosing a random policy (random) as baselines. The second part of the table shows the performance for the  $single\ best$  policy and the model-based  $policy\ selector$  when trained on different data obtained on  $\mathbf{D}_{meta}$  (P = Portfolio,  $BO = Bayesian\ Optimization, <math>E = Ensemble$ ) as well as the model-based  $policy\ selector\ without\ the$  fallback. We boldface the best mean value (per optimization budget) and underline results that are not statistically different according to a Wilcoxon-signed-rank  $Test\ (\alpha = 0.05)$ .

Second, to study how much resources we need to spend on generating training data for our model-based policy selector, we consider three approaches: (P) only using the portfolio performance, (P+BO) actually running the portfolio and BO for 10 and 60 minutes, respectively, and (P+BO+E) additionally also constructing ensembles, which yield the most realistic meta-data. Running BO on all 208 datasets (P+BO) is by far more expensive than the table lookups (P); building an ensemble (P+BO+E) adds only several seconds to minutes on top compared to (P+BO).

For both optimization budgets using P+BO yields the best results using the model-based policy selector closely followed by P+BO+ENS, see Table 5. The cheapest method, P, yields the worst results showing that it is worth to invest resources into computing good meta-data. Looking at the single best, surprisingly, performance gets worse when using seemingly better meta-data. We investigated the reason why P+BO performs slightly better than P+BO+ENS. When using a model-based policy selector this can be explained by a single dataset for both time horizons for which the chosen policy is worse than the single best. When looking at the single best, there is no single datasets which stands out. To summarize, investing additional resources to compute realistic meta-data results in improved performance, but so far it appears that this is mostly to have the effect of BO in the meta-data, while the ensemble appears to lead to lower meta-data quality.

Finally, we also take a closer look at the impact of the fallback mechanism to verify that our improvements are not solely due to this component. We observe that the performance drops for all policy selection strategies when we not include this fallback mechanism. For the shorter 10 minutes setting we find that the model-based policy selector still outperforms the single best, while for the longer 60 minutes setting the single best leads to better

		10 Min		60 ]	Min
		Ø	$\operatorname{std}$	Ø	$\operatorname{std}$
	selector	3.58	0.23	2.47	0.18
All	$\operatorname{random}$	7.46	2.02	5.64	1.95
	oracle	2.33	0.06	1.22	0.08
	selector	4.03	0.14	3.18	0.15
Only Holdout	$\operatorname{random}$	3.78	0.23	3.13	0.12
	oracle	3.23	0.10	2.62	0.07
	selector	6.11	0.11	5.09	0.19
Only CV	$\operatorname{random}$	8.66	0.70	6.85	0.86
	oracle	5.28	0.06	3.94	0.10
	selector	3.50	0.20	2.39	0.18
Full budget	$\operatorname{random}$	7.64	2.00	5.46	1.52
	oracle	2.59	0.09	1.51	0.06
	selector	3.63	0.19	2.44	0.24
Only SH	$\operatorname{random}$	6.95	1.98	5.13	1.72
	oracle	2.75	0.07	1.68	0.12

Table 6: Final performance (averaged normalized balanced error rate) for the full system and when not considering all model selection strategies.

performance. The rather stark performance degradation compared to the regular model-based policy selector can mostly be explained by a few, huge datasets, to which the model-based policy selector cannot extrapolate (and which the single best does not account for). Based on these observations we suggest research into an adaptive fallback strategy which can change the model selection strategy during the execution of the AutoML system so that a policy selector can be used on out-of-distribution datasets. We conclude that using a model-based policy selector is very beneficial, and using a fallback strategy to cope with out-of-distribution datasets can substantially improve performance.

#### 4.3.2 Do we need different model selection strategies?

Next, we study whether we need the different model selection strategies. For this, we build model-based policy selectors on different subsets of the available eight combinations of model selection strategies and budget allocations:  $\{3\text{-fold CV}, 5\text{-fold CV}, 10\text{-fold CV}, \text{holdout}\} \times \{\text{SH, FB}\}$ . Only Holdout consists of holdout with SH or FB (2 combinations), Only CV comprises 3-fold CV, 5-fold CV and 10-fold CV, all of them with SH or FB (6 combinations), Full budget contains both holdout and cross-validation and assigns each pipeline evaluation the same budget (4 combinations) and Only SH uses SH to assign budgets (4 combinations).

In Table 4.3.2, we compare the performance of selecting a policy at random (random), the performance of selecting the best policy on the test set and thus giving a lower bound

		10min		60n	nin
		Ø	$\operatorname{std}$	Ø	$\operatorname{std}$
With Portfolio	Policy selector Single best			$\frac{2.47}{2.44}$	
Without Portfolio	Policy selector Single best	5.63 5.37	$0.89 \\ 0.58$		$0.32 \\ 0.61$

Table 7: Final performance (ADTM) after 10 and after 60 minutes with portfolios (top) and without (bottom). The row "with portfolio" and "policy selector" constitutes the full AutoML system including portfolios, BO and ensembles) and the row "without portfolios" and "policy selector" only removes the portfolios (both from the meta-data for model-based policy selector construction and at runtime). We boldface the best mean value (per optimization budget) and underline results that are not statistically different according to a Wilcoxon-signed-rank Test ( $\alpha = 0.05$ ).

on the ADTM (oracle) and our model-based policy selector. The oracle indicates the best possible performance with each of these subsets of model selection strategies. It turns out that both Only Holdout and Only CV have a much worse oracle performance than All, with the oracle performance of Only CV being even worse than the performance of the model-based policy selector for All. Looking at Full budget, it turns out that this subset would be slightly preferable in terms of performance with a policy selector, however, the oracle performance is worse than that of All which shows that there is some complementarity between the different policies which cannot yet be exploited by the policy selector. For Only Holdout surprisingly the random policy selector performs slightly better than the model-based policy selector. We attribute this to the fact that both holdout with SH and FB peform very similar and that the choice between these two cannot yet be learned, possibly also indicated by the close performance of the random selector.

These results show that a large variety of available model selection strategies to choose from increases best possible performances. However, they also show that a model-based policy selector cannot yet necessarily leverage this potential. This questions the usefulness of choosing from all model selection strategies, similar to a recent finding which proves that increasing the number of different policies a policy selector can choose from leads to reduced generalization (Balcan et al., 2020). However, we believe this points to the research question whether we can learn on the meta-datasets which model selection and budget allocation strategies to include in the set of strategies to choose from. Also, with an ever-growing availability of meta-datasets and continued research on robust policy selectors, we expect this flexibility to eventually yield improved performance.

#### 4.3.3 Do we still need to warm-start Bayesian optimization?

Last, we analyse the impact of the portfolio. Given the other improvements, we now discuss the question whether we still need to add the additional complexity and invest resources to warm-start BO (and can therefore save the time to build the performance matrices to

construct the portfolios). For this study, we completely remove the portfolio from our AutoML system, meaning that we directly start with BO and construct ensembles – both for creating the data we train our policy selector on and for reporting performance. We report the results in Table 7.

Comparing the performance of an AutoML system with a model-based policy selector with and without portfolios (Row 1 and 3), there is a clear drop in performance showing the benefit of using portfolios in our system. Comparing Rows 2 and 4 also demonstrates that a portfolio is necessary when using the single best policy. This ablation demonstrates the importance of initializing the search procedure of AutoML systems with well-performing pipelines.

# 5. Comparison to other AutoML systems

Having established that Auto-sklearn~2.0 does indeed improve over Auto-sklearn~1.0, we now compare our system to other well established AutoML systems. For this, we use the publicly available AutoML benchmark suite which defines a fixed benchmarking environment for AutoML systems (Gijsbers et al., 2019) comparisons. We use the original implementation of the benchmark and compare Auto-sklearn~1.0 and Auto-sklearn~2.0 to the provided implementations of Auto-WEKA (Thornton et al., 2013), TPOT (Olson et al., 2016a,b), H2O AutoML (LeDell and Poirier, 2020) and a TunedRF on 39 datasets as implemented by the benchmark suite. These 39 are the same datasets as in  $\mathbf{D}_{test}$  and we provide details in Table 18 in the appendix.

#### 5.1 Integration and setup

To avoid hardware-dependent performance differences, we (re-)ran all AutoML systems on our local hardware (see Section 3.4.3). We used the pre-defined 1h8c setting, which divides each dataset into ten folds and gives each framework one hour on eight CPU cores to produce a final model. We furthermore assigned each run 32GB of RAM which is controlled by the SLURM cluster manager. In addition, we conducted three repeats to account for randomness. The benchmark comes with Docker containers (Merkel, 2014). However, Docker requires super user access on the execution nodes, which is not available on our compute cluster. Therefore, we extended the AutoML benchmark with support for Singularity images (Kurtzer et al., 2017), and used them to isolate the framework installations from each other.

The default resource allocation of the AutoML benchmark is a highly parallel setting with eight cores. We chose the most straight-forward way of making use of these resources for Auto-sklearn and evaluate eight ML pipelines in parallel, assigning each total\_memory/num\_cores RAM, which are 4GB. This allows us to evaluate configurations obtained from the portfolio or KND in parallel, but also requires a parallel strategy for running BO afterwards. In preliminary experiments we found that the inherent randomness of the random forest used by SMAC combined with the interleaved random search of SMAC is sufficient to obtain results which perform a lot better than the previous parallelism implemented in Auto-sklearn (Ramage, 2015). Whenever a pipeline finishes training, Auto-sklearn checks whether there is an instance of the ensemble construction running, and if not, it uses one of the eight slots to conduct ensemble building, and otherwise continues to

fit a new pipeline. We implemented this version of parallel Auto-sklearn using Dask (Dask Development Team, 2016).

#### 5.2 Results

We give results for the AutoML benchmark in Table 8. For each dataset we give the average performance of the AutoML systems across all ten folds and three repetitions and boldface the one with the lowest error, but cannot give any information about whether these results are significant as we cannot compute significances on cross-validation folds as described by Bengio and Grandvalet (2004).

We report the log loss for multiclass datasets and 1-AUC for binary datasets (lower is better). We provide two variants of the rank over datasets as an aggregate measure: 1) Average: we first average the loss on the ten folds per dataset, compute the rank per dataset and average these 39 numbers. 2) Per fold: we compute the rank per fold and average these 390 numbers. Furthermore, we count how often each framework is the winner on a dataset (champion), and give the losses, wins and ties against Auto-sklearn 2.0. We then use these to perform a binomial sign test (Demšar, 2006) to compare the individual algorithms against Auto-sklearn 2.0.

The results in Table 8 show that none of the AutoML systems is best on all datasets and even the TunedRF performs best on a few datasets. However, we can also observe that the proposed Auto-sklearn 2.0 has the lowest average rank. It is followed by H2O AutoML and Auto-sklearn 1.0 which perform roughly en par wrt the ranking scores and the number of times they are the winner on a dataset. According to both aggregate metrics, the TunedRF, Auto-WEKA and TPOT cannot keep up and lead to substantially worse results. Finally, both version of Auto-sklearn appear more robust than TPOT and H2O as they reliably provide results on the largest dataset dionis with 355-classes.

### 6. Related Work

We now present related work on our individual contributions (portfolios, model-selection strategies, and algorithm selection) as well as on related AutoML systems.

#### 6.1 Related Work on Portfolios

Portfolios were introduced for hard combinatorial optimization problems, where the runtime between different algorithms varies drastically and allocating time shares to multiple algorithms instead of allocating all available time to a single one reduces the average cost for solving a problem (Huberman et al., 1997; Gomes and Selman, 2001), and had applications in different sub-fields of AI (Smith-Miles, 2008; Kotthoff, 2014; Kerschke et al., 2019). Algorithm portfolios were introduced to ML by the name of algorithm ranking with the goal of reducing the required time to perform model selection compared to running all algorithms under consideration (Brazdil and Soares, 2000; Soares and Brazdil, 2000), ignoring redundant ones (Brazdil et al., 2001). ML portfolios can be superior to hyperparameter optimization with Bayesian optimization (Wistuba et al., 2015b), Bayesian optimization with a model which takes past performance data into account (Wistuba et al., 2015a) or can be applied when there is simply no time to perform full hyperparameter optimization (Feurer

	AS 2.0	AS 1.0	AW	TPOT	H2O	TunedRI
adult	0.0692	0.0701	0.0926	0.0769	0.0691	0.0903
airlines	0.2725	0.2728	0.3216	0.2755	0.2682	
albert	0.2413	0.2388	-	0.2686	0.2530	0.2613
amazon	0.1226	0.1412	0.1846	0.1351	0.1219	0.1379
apsfailure	0.0086	0.0079	0.0344	0.0098	0.0082	0.0088
australian	0.0596	0.0702	0.0714	0.0674	0.0603	0.0613
bank-marketing	0.0607	0.0617	0.1440	0.0659	0.0611	0.0692
blood-transfusion	0.2431	0.2500	0.2622	0.2723	0.2449	0.3129
car	0.0014	0.0049	0.2010	2.6530	0.0033	0.0420
christine	0.1819	0.1708	0.2006	0.1824	0.1763	0.190'
cnae-9	0.1448	0.1784	0.6972	0.1510	0.1876	0.316
connect-4	0.3356	0.3536	2.0405	0.3936	0.3122	0.477'
covertype	0.1094	0.1455	3.3422	0.5348	0.1274	
credit-g	0.2038	0.2165	0.2520	0.2149	0.2077	0.1982
dilbert	0.0380	0.0331	2.1825	0.1155	0.0366	0.328
dionis	0.5735	0.6595	-	-	-	
fabert	0.7421	0.7466	5.9815	0.8412	0.7281	0.805
fashion-mnist	0.2518	0.2523	0.9592	0.4259	0.2768	0.361
guillermo	0.0945	0.0871	0.1254	0.1651	0.0909	0.097
helena	2.4975	2.5428	14.6759	2.8813	2.7759	
higgs	0.1824	0.1848	0.3315	0.1967	0.1846	0.196
jannis	0.6701	0.6642	2.4168	0.7247	0.6695	0.729
jasmine	0.1141	0.1211	0.1388	0.1125	0.1147	0.111
jungle_chess	0.2109	0.1891	1.7751	0.6018	0.1468	0.402
kc1	0.1615	0.1597	0.1782	0.1538	0.1787	0.158
kddcup09	0.1579	0.1638	-	0.1692	0.1635	0.206
kr-vs-kp	0.0001	0.0003	0.0183	0.0002	0.0002	0.000
mfeat-factors	0.0733	0.0913	0.5619	0.1038	0.0987	0.211
miniboone	0.0121	0.0128	0.0359	0.0178	0.0129	0.018
nomao	0.0035	0.0039	0.0160	0.0046	0.0036	0.004
numerai28.6	0.4696	0.4706	0.4724	0.4747	0.4693	0.479
phoneme	0.0298	0.0366	0.0441	0.0308	0.0322	0.034
riccardo	0.0002	0.0002	0.0021	0.0020	0.0003	0.000
robert	1.4176	1.3799	-	1.8832	1.4977	1.687
segment	0.1482	0.1767	1.1884	0.1638	0.1582	0.172
shuttle	$\boldsymbol{0.0002}$	0.0004	0.0097	0.0007	0.0005	0.000
sylvine	0.0106	0.0091	0.0292	0.0074	0.0107	0.015
vehicle	0.3359	0.3636	1.7837	0.4156	0.3077	0.483
volkert	0.7493	0.7872	3.9654	0.9888	0.8148	0.979
Rank	1.77	2.69	5.71	4.01	2.45	4.3
Best performance	19	8	0	2	8	
Wins/Losses/Ties of AS 2.0	-	28/11/0	39/0/0	36/3/0	26/13/0	36/3/
P-values (AS 2.0 vs. other methods),	-	0.009	0.000	0.000	0.053	0.00
based on a Binomial sign test						

Table 8: Results of the AutoML benchmark averaged across three repetitions. We report log loss for multiclass datasets and 1-AUC for binary classification datasets (lower is better). AS is short for Auto-sklearn and AW for Auto-WEKA. Auto-sklearn has the best overall ranks, the best performance in most datasets and, based on a Binomial sign test, statistically significantly outperforms four out of five of the other methods.

and Hutter, 2018). Furthermore, such a portfolio-based model-free optimization is both easier to implement than regular Bayesian optimization and meta-feature based solutions, and the portfolio can be shared easily across researchers and practitioners without the necessity of sharing meta-data (Wistuba et al., 2015a,b; Pfisterer et al., 2018) or additional hyperparameter optimization software. Here, our goal is to have strong hyperparameter settings when there is no time to do optimization with a typical blackbox algorithm.

The efficient creation of algorithm portfolios is an active area of research with the Greedy Algorithm being a popular choice (Xu et al., 2010, 2011; Seipp et al., 2015; Wistuba et al., 2015b; Lindauer et al., 2017b; Feurer et al., 2018; Feurer and Hutter, 2018) due to its simplicity. Wistuba et al. (2015b) first proposed the use of the Greedy Algorithm for pipelines of ML portfolios, minimizing the average rank on meta-datasets for a single ML algorithm. Later, they extended their work to update the members of a portfolio in a round-robin fashion, this time using the average normalized misclassification error as a loss function and relying on a Gaussian process model (Wistuba et al., 2015a). The loss function of the first method does not optimize the metric of interest, while the second method does not guarantee that well-performing algorithms are executed early on, which could be harmful under time constraints.

Research into the Greedy Algorithm continued after our submission to the second AutoML challenge and the publication of the employed methods (Feurer et al., 2018; Feurer and Hutter, 2018). Pfisterer et al. (2018) suggested using a set of default values to simplify hyperparameter optimization. They argued that constructing an optimal portfolio of hyperparameter settings is a generalization of the *Maximum coverage problem* and propose two solutions based on *Mixed Integer Programming* and the *Greedy Algorithm* which we also use as the base of our algorithm. The greedy algorithm recently also drew interest in deep learning research where it was applied in its basic form for the tuning the hyperparameters of the popular ADAM algorithm (Metz et al., 2020).

Extending these portfolio strategies which are learned offline, there are online portfolios which can select from a fixed set of machine learning pipelines, taking previous evaluations into account (Leite et al., 2013; Wistuba et al., 2015a,b; Fusi et al., 2018; Yang et al., 2019, 2020). However, such methods cannot be directly combined with all budget allocation strategies as they require the definition of a special model for extrapolating learning curves (Klein et al., 2017b; Falkner et al., 2018) and also introduce additional complexity into AutoML systems.

There exists other work on building portfolios without prior discretization (which we do for our work and was done for most work mentioned above), which directly optimizes the hyperparameters of ML pipelines to add next to the portfolio in a greedy fashion (Xu et al., 2010, 2011; Seipp et al., 2015), to jointly optimize all configurations of the portfolio with global optimization (Winkelmolen et al., 2020), and to also build parallel portfolios (Lindauer et al., 2017b). We consider these to be orthogonal to using portfolios in the first place and plan to study improved optimization strategies in future work.

### 6.2 Related Work on Successive Halving

Large datasets, expensive ML pipelines and tight resource limitations demand for sophisticated methods developed to speed up pipeline selection. One line of research, multi-fidelity

optimization methods, tackle this problem by using cheaper approximations of the objective of interest. Practical examples are evaluating a pipeline only on a subset of the dataset or for iterative algorithms limit the number of iterations. There exists a large body of research on optimization methods leveraging lower fidelities, for example working with a fixed set of auxiliary tasks (Forrester et al., 2007; Swersky et al., 2013; Poloczek et al., 2017; Moss et al., 2020), solutions for specific model classes (Swersky et al., 2014; Domhan et al., 2015; Chandrashekaran and Lane, 2017) and selecting a fidelity value from a continuous range (Klein et al., 2017a; Kandasamy et al., 2017; Wu et al., 2020; Takeno et al., 2020). Here, we focus on a methodologically simple bandit strategy, SH (Karnin et al., 2013; Jamieson and Talwalkar, 2016), which successively reduces the number of candidates and at the same time increases the allocated resources per run till only one candidate remains. Our use of SH in the 2nd AutoML challenge also inspired work on combining a genetic algorithm with SH (Parmentier et al., 2019). Another way of quickly discarding unpromising pipelines is the intensify procedure which was used by Auto-WEKA (Thornton et al., 2013) to speed up cross-validation. Instead of evaluating all folds at once, it evaluates the folds in an iterative fashion. After each evaluation, the average performance on the evaluated folds is compared to the performance of the so-far best pipeline on these folds, and the evaluation is only continued if the performance is equal or better. While this allows evaluating many configurations in a short time, it cannot be combined with post-hoc ensembling and reduces the cost of a pipeline to at most the cost of holdout, which might already be too high.

# 6.3 Related Work on Algorithm Selection

Automatically choosing a model selection strategy to assess the performance of an ML pipeline for hyperparameter optimization has not previously been tackled, and only Guyon et al. (2015) acknowledge the lack of such an approach. However, treating the choice of model selection strategy as an algorithm selection problem allows us to apply methods from the field of algorithm selection (Smith-Miles, 2008; Kotthoff, 2014; Kerschke et al., 2019) and we can in future work reuse existing techniques besides the pairwise classification we employ in this paper (Xu et al., 2011), such as the AutoAI system AutoFolio (Lindauer et al., 2015).

#### 6.4 Background on AutoML Systems and Their Components

AutoML systems have recently gained traction in the research community and there exists a multitude of approaches, often accompanied by open-source software. In the following we provide background on the main components of AutoML frameworks before describing several prominent instantiations in more depth.

#### 6.4.1 Components of AutoML systems

AutoML systems require a flexible pipeline configuration space and are driven by an efficient method to search this space. Furthermore, they rely on model selection and budget allocation strategies when evaluating different pipelines. Additionally, to speed up the search procedure, information gained on other datasets can be used to kick-start or guide the search procedure (i.e. meta-learning). Finally, one can also combine the models trained during the search phase in a post-hoc ensembling step.

Configuration Space and Search Mechanism While there are configuration space formulations that allow the application of multiple search mechanisms, not all formulations of a configuration space and a search mechanism can be mixed and matched, and we therefore describe the different formulations and the applicable search mechanisms in turn.

The most common description of the search space is the CASH formulation. There is a fixed amount of hyperparameters, each with a range of legal values or categorical choices, and some of them can be conditional, meaning that they are only active if other hyperparameters fulfill certain conditions. One such example is the choice of a classification algorithm and its hyperparameters. The hyperparameters of an SVM are only active if the categorical hyperparameter of the classification algorithm is set to SVM.

The CASH problem can be solved by standard blackbox optimization algorithms, and it was first proposed to use SMAC (Hutter et al., 2011) and TPE (Bergstra et al., 2011). Others proposed the use of evolutionary algorithms (Bürger and Pauli, 2015) and random search (LeDell and Poirier, 2020). It is also known as the full model selection problem (Escalante et al., 2009), and solutions in that strain of work proposed the use of particle swarm optimization (Escalante et al., 2009) and a combination of a genetic algorithm with particle swarm optimization (Sun et al., 2013). To improve performance one can prune the configuration space to reduce space to search through (Zhang et al., 2016), split the configuration space into smaller, more managable subspaces (Alaa and van der Schaar, 2018; Liu et al., 2020), or heavily employ expert knowledge (LeDell and Poirier, 2020).

Instead of a fixed configuration space, genetic programming can make use of a flexible, and possibly infinite space of components to be connected (Olson et al., 2016b,a). This approach can be formalized further by using grammar-based genetic programming (de Sa et al., 2017). Context-free grammars can also be searched by model-based reinforcement learning algorithms (Drori et al., 2019).

Formalizing the search problem as a search tree allows the application of a custom Monte-Carlo tree search (Rakotoarison et al., 2019) and hierarchical task networks with best-first search (Mohr et al., 2018). With discrete spaces it is also possible to use combinations of meta-learning and matrix factorization (Yang et al., 2019, 2020; Fusi et al., 2018). In the special case of using only neural networks in an AutoML system it is possible to stick with standard blackbox optimization (Mendoza et al., 2016, 2019; Zimmer et al., 2021), but one can also employ recent advances in neural architecture search (Elsken et al., 2019).

Meta-Learning. When there is knowledge about previous runs of the AutoML system on other datasets available, it is possible to employ meta-learning. One option is to define a dataset similarity measure, often by using hand-crafted meta-features which describe the datasets (Brazdil et al., 1994), to use the best solutions on the closest seen datasets to warmstart the search algorithm (Feurer et al., 2015a). While this way of meta-learning can be seen as an add-on to existing methods, other works use search strategies designed to take meta-learning into account, for example matrix factorization (Yang et al., 2019, 2020; Fusi et al., 2018) or reinforcement learning (Drori et al., 2019; Heffetz et al., 2020).

**Model Selection.** Given training data, the goal of an AutoML system is to find the best performing ML pipeline. Doing so, requires to best approximate the generalization error

to 1) provide a reliable and precise signal for the optimization procedure<sup>8</sup> and 2) select the model to be returned in the end. Typically, the generalization error is assessed via the trainvalidation-test protocol (Bishop, 1995; Raschka, 2018). This means that several models are trained on a  $training\ set$ , the best one is selected via holdout (using a single split) or the K-fold cross-validation, and the generalization error is then reported on the test set. The AutoML system then returns a single model in case of holdout, and a combination of K models in case of K-fold cross-validation (Caruana et al., 2006). One could also use model selection strategies aiming to reduce the effect of overfitting to the validation set (Dwork et al., 2015; Tsamardinos et al., 2018), but while such model selection strategies are an important area of research, houldout or K-fold cross-validation remain the most prominent choices (Henery, 1994; Kohavi and John, 1995; Hastie et al., 2001; Guyon et al., 2010; Bischl et al., 2012; Raschka, 2018).

The influence of the model selection strategy on the performance is well known (Kalousis and Hilario, 2003) and researchers have studied their impact (Kohavi, 1995). However, there is no single best strategy and since there is a tradeoff between approximation quality and time required to compute the validation loss.

**Post-hoc Ensembling.** AutoML systems evaluate dozens or hundreds of models during their optimization procedure. Thus, it is a natural next step to not only use a single model at the end, but to ensemble multiple for improved performance and reduced overfitting.

This was first proposed to combine the solutions found by particle swarm optimization (Escalante et al., 2010) and then by an evolutionary algorithm (Bürger and Pauli, 2015). While these works used heuristic methods to combine multiple models into a final ensemble, it is also possible to treat this as another optimization problem (Feurer et al., 2015a) and solve it with ensemble selection (Caruana et al., 2004) or stacking (LeDell and Poirier, 2020).

Instead of using a single layer of machine learning models, Automatic Frankensteining (Wistuba et al., 2017) proposed two-layer stacking, applying AutoML to the outputs of an AutoML system instead of a single layer of ML algorithms followed by an ensembling mechanism. Auto-Stacker went one step further, directly optimizing for a two-layer AutoML system (Chen et al., 2018).

#### 6.4.2 Automl systems

To the best of our knowledge, the first AutoML system which tunes both hyperparameters and chooses algorithms was an ensemble method (Caruana et al., 2004). This system randomly produces 2 000 classifiers from a wide range of ML algorithms and constructs a post-hoc ensemble. It was later robustified (Caruana et al., 2006) and employed in a winning submission to the KDD challenge (Niculescu-Mizil et al., 2009).

The first AutoML system to jointly optimize the whole pipeline was *Particle Swarm Model Selection* (Escalante et al., 2007, 2009). It used a fixed-length representation of

<sup>8.</sup> Different model selection strategies could be ignored from an optimization point of view, where the goal is to optimize performance given a loss function, as is often done in the research fields of meta-learning and hyperparameter optimization. However, for AutoML systems this is highly relevant as we are not interested in the optimization performance (of some subpart) of these systems, but the final estimated generalization performance when applied to new data.

the pipeline and contained feature selection, feature processing, classification and post-processing implemented in the CLOP package<sup>9</sup> and was developed for the IJCNN 2007 agnostic learning vs. prior knowledge challenge (Guyon et al., 2008). It placed 2nd among the solutions using the CLOP package provided by the organizers, only loosing to a sub-mission based on robust hyperparameter optimization and ensembling (Reunanen, 2007). Later systems started employing model-based global optimization algorithms, such as *Auto-WEKA* (Thornton et al., 2013; Kotthoff et al., 2019), which is built around the WEKA software (Hall et al., 2009) and SMAC (Hutter et al., 2011) and uses cross-validation with racing for model evaluation, and Hyperopt-sklearn (Komer et al., 2014), which was the first tool to use the now-popular scikit-learn (Pedregosa et al., 2011) and paired it with the TPE algorithm from the hyperopt package (Bergstra et al., 2011, 2013) and holdout.

We extended the approach of parametrizing a popular machine learning library and optimizing its hyperparameters with a blackbox optimization algorithm using meta-learning and post-hoc ensembles in *Auto-sklearn* (Feurer et al., 2015a, 2019). For classification, the space of possible ML pipelines currently spans 16 classifiers, 14 feature preprocessing methods and numerous data preprocessing methods, adding up to 122 hyperparameters for the latest release. *Auto-sklearn* uses holdout as a default model selection strategy, but allows for other strategies such as cross-validation. *Auto-sklearn* was the dominating solution of the first AutoML challenge (Guyon et al., 2019).

The tree-based pipeline optimization tool (TPOT (Olson et al., 2016b; Olson and Moore, 2019)) uses grammatical evolution to construct ML pipelines of arbitrary length. Currently, it uses scikit-leearn (Pedregosa et al., 2011) and XGBoost (Chen and Guestrin, 2016) for its ML building blocks and 5-fold cross-validation to evaluate individual solutions. TPOT-SH (Parmentier et al., 2019), inspired by our submission to the second AutoML challenge, uses successive halving to speed up TPOT on large datasets.

The *H2O AutoML* package takes a radically different approach, building on a manually designed set of defaults and random search combined with stacking. It uses building blocks from the H2O library(H2O.ai, 2020) and XGBoost (Chen and Guestrin, 2016) and cross-validation.

Finally, there is also work on creating AutoML systems that can leverage recent advancements in deep learning, using either blackbox optimization (Mendoza et al., 2016; Zimmer et al., 2021) or neural architecture search (Jin et al., 2019).

Of course, there are also many techniques related to AutoML which are not used in one of the AutoML systems discussed in this section and we refer to Hutter et al. (2019) for an overview of the field of Automated Machine Learning and to Brazdil et al. (2008) for an overview on meta-learning research which pre-dates the work on AutoML.

### 7. Discussion and Conclusion

In this paper we introduced our winning entry to the 2nd ChaLearn AutoML challenge, *PoSH Auto-sklearn*, and automated its internal settings further, resulting in the next generation of our AutoML system: *Auto-sklearn 2.0*. *Auto-sklearn 2.0* provides a truly handsfree solution, which, given a new task and resource limitations, automatically chooses the best setup. Specifically, we introduce three improvements for faster and more efficient Au-

<sup>9.</sup> http://clopinet.com/CLOP/

toML: (i) to get strong results quickly we propose to use portfolios, which can be built offline and thus reduce startup costs, (ii) to reduce time spent on poorly performing pipelines we propose to add successive halving (SH) as a budget allocation strategy to the configuration space of our AutoML system and (iii) to close the design space we opened up for AutoML we propose to automatically select the best configuration of our system.

We conducted a large-scale study based on 208 meta-datasets and 39 datasets for testing and obtained substantially improved performance compared to  $Auto-sklearn\ 1.0$ , reducing the ADTM by up to a factor of four and achieving a lower loss after 10 minutes than  $Auto-sklearn\ 1.0$  after 60 minutes. Our ablation study showed that using a model-based policy selector to choose the model selection strategy has the greatest impact on performance and allows  $Auto-sklearn\ 2.0$  to run robustly on new, unseen datasets. Furthermore, we showed that our method is highly competitive and outperforms other state-of-the-art AutoML systems in the OpenML AutoML benchmark.

However, our system also introduces some shortcomings since it optimizes performance towards a given optimization budget, performance metric and configuration space. Although all of these, along with the meta datasets, could be provided by a user to automatically build a customized version of *Auto-sklearn 2.0*, it would be interesting whether we can learn how to transfer a specific AutoML system to different optimization budgets and metrics. Also, there still remain several hand-picked hyperparameters on the level of the AutoML system, which we would like to automate in future work, too, for example by automatically learning the portfolio size, learning more hyper-hyperparameters of the different budget allocation strategies (for example of SH) and proposing suitable configuration spaces given a dataset and resources. Finally, building the training data is currently quite expensive. Even though this has to be done only once, it will be interesting to see whether we can take shortcuts here, for example by using a joint ranking model (Tornede et al., 2020) or non-linear collaborative filtering (Fusi et al., 2018).

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# Appendix A. Additional pseudo-code

We give pseudo-code for computing the estimated generalization error of  $\mathcal{P}$  across all metadatasets  $\mathbf{D}_{\text{meta}}$  for K-folds cross-validation in Algorithm 2 and successive halving in Algorithm 3.

**Algorithm 2:** Estimating the generalization error of a portfolio with K-Fold Cross-Validation

```
1: Input: Ordered set of ML pipelines \mathcal{P}, datasets \mathbf{D}_{\text{meta}}, number of folds K,
 2: L = 0
 3: for d \in (1, 2, ..., |\mathbf{D}_{meta}|) do
         l_d = \infty
         for p \in \mathcal{P} do
             l = 0
 6:
             for k \in (1, 2, ..., K) do
 7:
                l = l + \widehat{GE}(\mathcal{M}_{\boldsymbol{\lambda}}^{\mathcal{D}_{\mathrm{train}}^{(\mathrm{train},k)}}, \mathcal{D}_{\mathrm{train}}^{(\mathrm{val},k)})
 8:
             end for
 9:
             l = l/K
10:
             if l < l_d then
11:
                 l_d = l
12:
             end if
13:
         end for
14:
          L = L + l_d
15:
16: end for
17: return L/|\mathbf{D}_{\text{meta}}|
```

# Appendix B. Additional results and experiments

In this section we will give additional results backing up our findings. Concretely, we will give further details on the reduced search space and provide further experimental evidence, we will provide the main results from the main paper without post-hoc ensembles, and we will give the raw numbers before averaging.

## **B.1** Early Stopping and Retrieving Intermittent Results

Estimating the generalization error of a pipeline  $\mathcal{M}_{\lambda}$  practically requires to restrict the CPU-time per evaluation to prevent that one single, very long algorithm run stalls the optimization procedure (Thornton et al., 2013; Feurer et al., 2015a). If an algorithm does not return a result within the assigned time limit, it is terminated and the worst possible generalization error is assigned. If the time limit is set too low, a majority of the algorithms do not return a result and thus provide very scarce information for the optimization procedure. A too high time limit, however, might as well not return any meaningful results since all time may be spent on long-running, under-performing pipelines. Additionally, for iterative algorithms (e.g., gradient boosting and linear models trained with stochastic

**Algorithm 3:** Estimating the generalization error of a portfolio with Successive Halving

```
1: Input: Ordered set of ML pipelines \mathcal{P}, datasets \mathbf{D}_{\text{meta}}, minimal budget b_{min},
      maximal budget b_{max}, downsampling rate \eta
 2: L=\infty
 3: R = b_{max}/b_{min}
 4: s_{max} = \lfloor \log_n(R) \rfloor
 5: B = (s_{max} + 1)R

6: n = \lceil \frac{B}{R} \frac{\eta^{s_{max}}}{(s_{max} + 1)} \rceil

7: r = R\eta^{-s_{max}}
 8: for d \in (1, 2, \dots, |\mathbf{D}_{\text{meta}}|) do
         l_d = \infty
         \mathcal{P}_d = \mathcal{P}
10:
          while True do
11:
             \mathcal{P}' = \mathcal{P}.pop(r) \# Pop top r machine learning pipelines
12:
13:
14:
             for i \in (0, \ldots, s_{max}) do
                 n_i = |n\eta^{-i}|
15:
                 r_i = r\eta^i
16:
                 for p \in \mathcal{P}' do
17:
                    l = \widehat{GE}(\mathcal{M}_{\lambda}^{\mathcal{D}_{\mathrm{train}}^{\mathrm{train}}}, \mathcal{D}_{\mathrm{train}}^{\mathrm{val}})
18:
                     l = l \cup l
19:
                     if l < l_d then
20:
21:
                         l_d = l
                     end if
22:
23:
                 end for
                 \mathcal{P}' = top(\mathcal{P}', \mathbf{l}, |(n_i/eta)|), where top(\mathcal{P}, \mathbf{l}, k) returns the top k performing
24:
                 machine learning pipelines.
             end for
25:
             if |\mathcal{P}_d| == 0 then
26:
                 break
27:
             end if
28:
          end while
29:
          L = L + l_d
30:
31: end for
32: return L/|\mathbf{D}_{\text{meta}}|
```

		10	STD 10	60	STD 60
(1)	Auto-sklearn (1.0)	16.21	0.27	7.17	0.30
(2)	Auto-sklearn (1.0) ISS	18.10	0.13	9.57	0.22
(3)	Auto-sklearn $(1.0)$ ISS + IRR	5.29	0.13	3.98	0.21
(4)	Auto-sklearn (1.0) ISS + IRR + Port	3.70	0.14	3.08	0.13

Table 9: Comparison of *Auto-sklearn 1.0* (1) with using only the iterative search space (2), using the iterative search space and iterative results retrieval (3) and also using a portfolio (4).

gradient descent), it is important to set the number of iterations such that the training converges and does not overfit, but most importantly finishes within this timelimit. Setting this number too high (training exceeds time limit and/or overfit) or too low (training has not yet converged although there is time left) has detrimental effects to the final performance of the AutoML system. To mitigate this risk we implemented two measures for iterative algorithms. Firstly, we use the early stopping mechanisms implemented by scikit-learn. Specifically, training stops if the loss on the training or validation set (depending on the model and the configuration) increases or stalls, which prevents overfitting (i.e. early stopping). Secondly, we make use of intermittent results retrieval, e.g., saving the results at checkpoints spaced at geometrically increasing iteration numbers, thereby ensuring that every evaluation of an iterative algorithm returns a performance and thus yields information for the optimizer. With this, our AutoML tool can robustly tackle large datasets without the necessity to finetune the number of iterations dependent on the time limit.

To study the effect of using the iterative results retrieval we compare Auto-sklearn 1.0 we one by one make the following changes: 1) move to a configuration space which consists only of iterative algorithms 2) enable intermittent results retrieval and 3) replace the KND by the portfolio. We give results in Table 9 and note that the KND uses meta-data gathered specifically for use with the reduced configuration space. Only restricting the configuration space leads to decreased performance which we attribute to the reduced hypothesis space. Intermittently writing results to disk reduces the amount of failures, and using a portfolio instead of the KND results in the best overall performance.

Once again, we also view the results through the eyes of a ranking plot in Figure 6. These results demonstrate that the iterative search space combined with intermittent results retrieval and a portfolio is especially dominating in the short term, and it takes a total of 50 minutes for *Auto-sklearn 1.0* to catch up. We would like to note that the performance of *Auto-sklearn 2.0* is even better as can be seen in Table 4, but it would be interesting to see how a portfolio of the full configuration space would perform, which we note as a further research question.

#### **B.2** Performance Without Post-Hoc Ensembling

We first give numbers comparing only Bayesian optimization, k-nearest datasets (KND) and a greedy portfolio. These results are similar to Table 2, but do not show the results of post-hoc ensembling, but using the single best model. Overall, they are qualitatively very

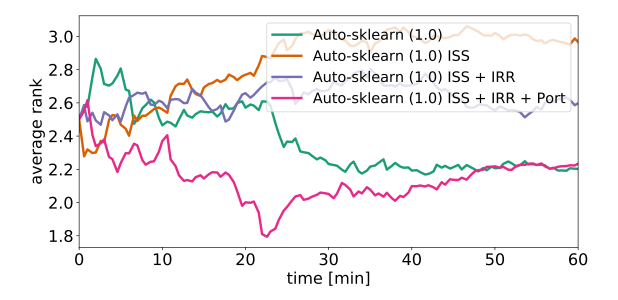


Figure 6: Ranking plot comparing *Auto-sklearn 1.0* (1) with using only the iterative search space (2), using the iterative search space and iterative results retrieval (3) and also using a portfolio (4).

	10	0 minute	es	60 minutes				
	ВО	KND	Port	ВО	KND	Port		
holdout	7.27	6.43	4.76	4.58	4.99	4.02		
SH; holdout	6.61	6.70	5.76	4.70	4.63	3.97		
3CV	9.58	8.95	7.88	7.10	7.12	5.98		
SH; 3CV	8.88	8.97	7.20	6.81	6.47	6.01		
5CV	10.48	15.24	13.77	7.34	7.47	5.66		
SH; 5CV	11.70	13.29	8.06	7.05	6.69	$\bf 5.93$		
10CV	23.20	27.45	18.73	17.59	17.47	16.17		
SH; 10CV	23.98	27.70	18.84	<u>16.94</u>	16.98	16.07		

Table 10: Results from Table 2 without post-hoc ensembles.

similar, but it can be observed that the ensemble improves the average normalized balanced error rate in every case.

Next, we compare Auto-sklearn 2.0 with PoSH Auto-sklearn and Auto-sklearn 1.0, but again only show the performance of the single best model and not of an ensemble as in the main paper. Again, the ensemble result in uniform performance improvements with Auto-sklearn 2.0 still leading in terms of performance.

	10M	IIN	60MIN		
	Ø	$\operatorname{std}$	Ø	$\operatorname{std}$	
Auto-sklearn (2.0)	5.01	0.18	3.18	0.31	
PoSH-Auto-sklearn	5.76	0.12	3.97	0.22	
Auto-sklearn $(1.0)$	23.24	0.29	8.68	0.21	

Table 11: Results from Table 4 without post-hoc ensembles.

# **B.3** Unaggregated results

To allow the readers to asses the performance of the individual methods on the individual datasets we present the balanced error rates before normalizing and averaging them. We give the raw results for portfolios from Table 2 in Tables 12 and 13. Additionally, we give the raw results for *Auto-sklearn 2.0*, *PoSH Auto-sklearn* and *Auto-sklearn 1.0* in Tables 14 and 15.

## Appendix C. Theoretical properties of the greedy algorithm

#### C.1 Definitions

**Definition 1** (Discrete derivative, from Krause & Golovin (Krause and Golovin, 2014)) For a set function  $f: 2^{\mathcal{V}} \to \mathbb{R}, \mathcal{S} \subseteq \mathcal{V}$  and  $e \in \mathcal{V}$  let  $\Delta_f(e|\mathcal{S}) = f(\mathcal{S} \cup \{e\}) - f(\mathcal{S})$  be the discrete derivative of f at  $\mathcal{S}$  with respect to e.

**Definition 2** (Submodularity, from Krause & Golovin (Krause and Golovin, 2014)): A function  $f: 2^{\mathcal{V}} \to \mathbb{R}$  is submodular if for every  $\mathcal{A} \subseteq \mathcal{B} \subseteq \mathcal{V}$  and  $e \in \mathcal{V} \setminus \mathcal{B}$  it holds that  $\Delta_f(e|\mathcal{A}) \geq \Delta_f(e|\mathcal{B})$ .

**Definition 3** (Monotonicity, from Krause & Golovin (Krause and Golovin, 2014)): A function  $f: 2^{\mathcal{V}} \to \mathbb{R}$  is monotone if for every  $A \subseteq B \subseteq V$ ,  $f(A) \leq f(B)$ .

### C.2 Choosing on the test set

In this section we give a proof of Proposition 1 from the main paper:

**Proposition 2** Minimizing the test loss of a portfolio  $\mathcal{P}$  on a set of datasets  $\mathcal{D}_1, \ldots, \mathcal{D}_{|\mathbf{D}_{meta}|}$ , when choosing a ML pipeline from  $\mathcal{P}$  for  $\mathcal{D}_d$  based on performance on  $\mathcal{D}_{d,test}$ , is equivalent to the sensor placement problem for minimizing detection time (Krause et al., 2008).

Following Krause et al. (Krause et al., 2008), sensor set placement aims at maximizing a so-called penalty reduction  $R(\mathcal{A}) = \sum_{i \in \mathcal{I}} P(i)R(\mathcal{A},i)$ , where  $\mathcal{I}$  are intrusion scenarios following a probability distribution P with i being a specific intrusion.  $\mathcal{A} \subset \mathcal{C}$  is a sensor placement, a subset of all possible locations  $\mathcal{C}$  where sensors are actually placed. Penalty reduction R is defined as the reduction of the penalty when choosing  $\mathcal{A}$  compared to the maximum penalty possible on scenario i:  $R(\mathcal{A},i) = \text{penalty}_i(\infty) - \text{penalty}_i(T(\mathcal{A},i))$ . In the simplest case where action is taken upon intrusion detection, the penalty is equal to the

Task ID	Name	holdout	SH; holdout	3CV	SH; 3CV	5CV	SH; 5CV	10CV	SH; 10CV
167104	Australian	0.1721	0.1569	0.1622	0.1617	0.1583	0.1602	0.1556	0.1559
167184	blood-transfusion	0.3641	0.3610	0.3725	0.3666	0.3689	0.3722	0.3674	0.3689
167168	vehicle	0.2211	0.2267	0.2017	0.2093	0.2172	0.2052	0.2310	0.1870
167161	credit-g	0.2942	0.2841	0.2939	0.2955	0.2942	0.2911	0.2939	0.2934
167185	cnae-9	0.0658	0.0680	0.0651	0.0616	0.0550	0.0629	0.0626	0.0553
189905	car	0.0049	0.0049	0.0097	0.0029	0.0047	0.0017	0.0023	0.0009
167152	mfeat-factors	0.0152	0.0164	0.0141	0.0107	0.0150	0.0117	0.0153	0.0149
167181	kc1	0.2735	0.2688	0.2720	0.2713	0.2547	0.2660	0.2477	0.2719
189906	segment	0.0666	0.0687	0.0681	0.0620	0.0664	0.0621	0.0643	0.0671
189862	jasmine	0.2044	0.2051	0.1982	0.1986	0.2010	0.2027	0.2043	0.2027
167149	kr-vs-kp	0.0067	0.0077	0.0093	0.0085	0.0079	0.0078	0.0071	0.0080
189865	sylvine	0.0592	0.0594	0.0600	0.0608	0.0582	0.0582	0.0560	0.0578
167190	phoneme	0.1231	0.1245	0.1168	0.1160	0.1152	0.1136	0.1129	0.1144
189861	christine	0.2670	0.2621	0.2608	0.2556	0.2517	0.2567	0.2587	0.2645
189872	fabert	0.3387	0.3399	0.3140	0.3120	0.3096	0.3204	0.3180	0.3172
189871	dilbert	0.0241	0.0248	0.0258	0.0220	0.0191	0.0211	0.0303	0.0647
168794	robert	0.5489	0.5861	0.5762	0.5583	0.5854	0.5873	0.6230	0.6230
168797	riccardo	0.0035	0.0052	0.0067	0.0054	0.0027	0.0027	0.5000	0.5000
168796	guillermo	0.2186	0.2102	0.2311	0.2228	0.2165	0.2837	0.5000	0.5000
75097	Amazon	0.2361	0.2431	0.2526	0.2526	0.2379	0.2385	0.2448	0.2443
126026	nomao	0.0353	0.0381	0.0360	0.0345	0.0312	0.0331	0.0403	0.0401
189909	$jungle\_chess$	0.1212	0.1251	0.1232	0.1156	0.1280	0.1180	0.1134	0.1141
126029	bank-marketing	0.1397	0.1436	0.1402	0.1407	0.1352	0.1435	0.1378	0.1362
126025	adult	0.1579	0.1575	0.1553	0.1540	0.1591	0.1562	0.1545	0.1585
75105	KDDCup09	0.2450	0.2495	0.2449	0.2512	0.2525	0.2487	0.2497	0.2456
168795	shuttle	0.0093	0.0086	0.0085	0.0084	0.0085	0.0087	0.0088	0.0084
168793	volkert	0.3735	0.3724	0.3939	0.3703	0.3720	0.3775	0.3867	0.3957
189874	helena	0.7483	0.7624	0.7475	0.7478	0.7483	0.7534	0.7476	0.7575
167201	connect-4	0.2629	0.2721	0.2653	0.2630	0.2537	0.2565	0.3003	0.2771
189908	Fashion-MNIST	0.1050	0.1098	0.1217	0.1195	0.1181	0.1153	0.1437	0.1437
189860	APSFailure	0.0384	0.0402	0.0355	0.0364	0.0355	0.0354	0.0410	0.0455
168792	jannis	0.3654	0.3685	0.3648	0.3655	0.3651	0.3567	0.3912	0.3881
167083	numerai28.6	0.4776	0.4765	0.4752	0.4749	0.4747	0.4775	0.4789	0.4788
167200	higgs	0.2736	0.2764	0.2730	0.2742	0.2724	0.2744	0.2832	0.2844
168798	MiniBooNE	0.0581	0.0589	0.0691	0.0633	0.0585	0.0644	0.0691	0.0685
189873	dionis	0.1172	0.1205	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000
189866	albert	0.3135	0.3171	0.3469	0.3277	0.5000	0.3354	0.3714	0.3703
75127	airlines	0.3423	0.3424	0.3450	0.3384	0.3419	0.3429	0.3408	0.3456
75193	covertype	0.0568	0.0564	0.0683	0.0600	0.0548	0.0556	0.2519	0.2527

Table 12: Results from Table 2 for 10 minutes using portfolios. Numbers are the balanced error rate. We boldface the lowest error.

Task ID	Name	holdout	SH; holdout	3CV	SH; 3CV	5CV	SH; 5CV	10CV	SH; 10CV
167104	Australian	0.1742	0.1674	0.1623	0.1626	0.1598	0.1608	0.1625	0.1557
167184	blood-transfusion	0.3648	0.3618	0.3631	0.3641	0.3689	0.3692	0.3684	0.3692
167168	vehicle	0.2125	0.2344	0.1702	0.1944	0.1657	0.1960	0.1959	0.2151
167161	credit-g	0.2922	0.2895	0.3035	0.2957	0.3056	0.2978	0.3008	0.2931
167185	cnae-9	0.0733	0.0761	0.0560	0.0616	0.0537	0.0536	0.0675	0.0518
189905	car	0.0036	0.0013	0.0037	0.0098	0.0007	0.0012	0.0008	0.0010
167152	mfeat-factors	0.0169	0.0186	0.0130	0.0117	0.0139	0.0132	0.0151	0.0122
167181	kc1	0.2728	0.2739	0.2680	0.2724	0.2678	0.2804	0.2546	0.2576
189906	segment	0.0708	0.0692	0.0647	0.0635	0.0588	0.0596	0.0621	0.0613
189862	jasmine	0.2048	0.2049	0.1995	0.1989	0.1995	0.1976	0.1995	0.1980
167149	kr-vs-kp	0.0060	0.0080	0.0081	0.0068	0.0064	0.0068	0.0055	0.0053
189865	sylvine	0.0590	0.0591	0.0584	0.0587	0.0577	0.0578	0.0573	0.0573
167190	phoneme	0.1222	0.1237	0.1152	0.1155	0.1111	0.1130	0.1117	0.1105
189861	christine	0.2673	0.2666	0.2575	0.2584	0.2532	0.2575	0.2549	0.2588
189872	fabert	0.3381	0.3319	0.3099	0.3097	0.3119	0.3080	0.3027	0.3071
189871	dilbert	0.0200	0.0200	0.0132	0.0146	0.0185	0.0209	0.0212	0.0149
168794	robert	0.5273	0.5199	0.5238	0.5183	0.5456	0.5605	0.5652	0.5407
168797	riccardo	0.0029	0.0016	0.0018	0.0019	0.0025	0.0076	0.5000	0.5000
168796	guillermo	0.2012	0.2025	0.2057	0.2081	0.2100	0.2039	0.5000	0.5000
75097	Amazon	0.2376	0.2394	0.2338	0.2381	0.2431	0.2384	0.2312	0.2324
126026	nomao	0.0352	0.0353	0.0334	0.0331	0.0320	0.0327	0.0313	0.0319
189909	jungle_chess	0.1214	0.1221	0.1154	0.1172	0.1171	0.1153	0.1108	0.1141
126029	bank-marketing	0.1388	0.1398	0.1380	0.1392	0.1382	0.1382	0.1370	0.1380
126025	adult	0.1546	0.1541	0.1550	0.1540	0.1550	0.1550	0.1539	0.1538
75105	KDDCup09	0.2492	0.2461	0.2477	0.2532	0.2466	0.2488	0.2617	0.2485
168795	shuttle	0.0136	0.0107	0.0125	0.0093	0.0084	0.0063	0.0127	0.0087
168793	volkert	0.3600	0.3673	0.3449	0.3551	0.3496	0.3487	0.3581	0.3563
189874	helena	0.7449	0.7494	0.7331	0.7369	0.7407	0.7404	0.7562	0.7452
167201	connect-4	0.2539	0.2556	0.2382	0.2428	0.2370	0.2373	0.2416	0.2369
189908	Fashion-MNIST	0.1010	0.0971	0.1046	0.1066	0.1102	0.1105	0.1191	0.1075
189860	APSFailure	0.0362	0.0374	0.0345	0.0364	0.0372	0.0347	0.0343	0.0334
168792	jannis	0.3670	0.3638	0.3589	0.3576	0.3584	0.3565	0.3473	0.3572
167083	numerai28.6	0.4765	0.4763	0.4774	0.4770	0.4750	0.4767	0.4755	0.4743
167200	higgs	0.2712	0.2734	0.2718	0.2680	0.2696	0.2680	0.2701	0.2683
168798	MiniBooNE	0.0576	0.0583	0.0560	0.0536	0.0571	0.0565	0.0560	0.0608
189873	dionis	0.0961	0.1068	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000
189866	albert	0.3116	0.3168	0.3170	0.3172	0.3094	0.3199	0.3183	0.3186
75127	airlines	0.3403	0.3410	0.3375	0.3401	0.3388	0.3390	0.3398	0.3399
75193	covertype	0.0537	0.0519	0.0496	0.0496	0.0454	0.0461	0.0458	0.0459

Table 13: Results from Table 2 for 60 minutes using portfolios. Numbers are the balanced error rate. We boldface the lowest error.

Task ID	Name	Auto-sklearn (2.0)	PoSH-Auto-sklearn	Auto-sklearn (1.0)
167104	Australian	0.1617	0.1569	0.1628
167184	blood-transfusion	0.3694	0.3610	0.3534
167168	vehicle	0.2030	0.2267	0.1654
167161	credit-g	0.2903	0.2841	0.2951
167185	cnae-9	0.0635	0.0680	0.0674
189905	car	0.0015	0.0049	0.0057
167152	mfeat-factors	0.0123	0.0164	0.0185
167181	kc1	0.2707	0.2688	0.2301
189906	segment	0.0646	0.0687	0.0624
189862	jasmine	0.2020	0.2051	0.1989
167149	kr-vs-kp	0.0086	0.0077	0.0089
189865	sylvine	0.0597	0.0594	0.0583
167190	phoneme	0.1158	0.1245	0.1257
189861	christine	0.2562	0.2621	0.2666
189872	fabert	0.3250	0.3399	0.3323
189871	dilbert	0.0240	0.0248	0.0066
168794	robert	0.5861	0.5861	0.6545
168797	riccardo	0.0052	0.0052	0.5000
168796	guillermo	0.2102	0.2102	0.5000
75097	Amazon	0.2435	0.2431	0.2610
126026	nomao	0.0336	0.0381	0.0383
189909	$jungle\_chess$	0.1205	0.1251	0.1231
126029	bank-marketing	0.1402	0.1436	0.1412
126025	adult	0.1547	0.1575	0.1608
75105	KDDCup09	0.2460	0.2495	0.2863
168795	shuttle	0.0084	0.0086	0.0111
168793	volkert	0.3717	0.3724	0.4233
189874	helena	0.7493	0.7624	0.9157
167201	connect-4	0.2642	0.2721	0.2809
189908	Fashion-MNIST	0.1070	0.1098	0.1383
189860	APSFailure	0.0372	0.0402	0.0370
168792	jannis	0.3654	0.3685	0.3637
167083	numerai28.6	0.4753	0.4765	0.4774
167200	higgs	0.2746	0.2764	0.2777
168798	MiniBooNE	0.0603	0.0589	0.0622
189873	dionis	0.1205	0.1205	0.6731
189866	albert	0.3171	0.3171	0.4407
75127	airlines	0.3404	0.3424	0.3536
75193	covertype	0.0564	0.0564	0.8571

Table 14: Results from Table 4 for 10 minutes. Numbers are the balanced error rate. We boldface the lowest error.

Task ID	Name	Auto-sklearn (2.0)	PoSH-Auto-sklearn	Auto-sklearn (1.0)
167104	Australian	0.1562	0.1674	0.1658
167184	blood-transfusion	0.3669	0.3618	0.3572
167168	vehicle	0.2187	0.2344	0.1822
167161	credit-g	0.2980	0.2895	0.3004
167185	cnae-9	0.0566	0.0761	0.0620
189905	car	0.0038	0.0013	0.0043
167152	mfeat-factors	0.0126	0.0186	0.0136
167181	kc1	0.2600	0.2739	0.2250
189906	segment	0.0609	0.0692	0.0697
189862	jasmine	0.1971	0.2049	0.1985
167149	kr-vs-kp	0.0060	0.0080	0.0085
189865	sylvine	0.0572	0.0591	0.0555
167190	phoneme	0.1140	0.1237	0.1235
189861	christine	0.2592	0.2666	0.2619
189872	fabert	0.3120	0.3319	0.3185
189871	dilbert	0.0163	0.0200	0.0090
168794	robert	0.5199	0.5199	0.5327
168797	riccardo	0.0016	0.0016	0.0016
168796	guillermo	0.2025	0.2025	0.1964
75097	Amazon	0.2371	0.2394	0.2481
126026	nomao	0.0323	0.0353	0.0361
189909	$jungle\_chess$	0.1145	0.1221	0.1136
126029	bank-marketing	0.1387	0.1398	0.1428
126025	adult	0.1544	0.1541	0.1574
75105	KDDCup09	0.2504	0.2461	0.2549
168795	shuttle	0.0093	0.0107	0.0109
168793	volkert	0.3563	0.3673	0.3440
189874	helena	0.7399	0.7494	0.7693
167201	connect-4	0.2408	0.2556	0.2709
189908	Fashion-MNIST	0.1023	0.0971	0.0984
189860	APSFailure	0.0343	0.0374	0.0375
168792	jannis	0.3591	0.3638	0.3641
167083	numerai28.6	0.4759	0.4763	0.4760
167200	higgs	0.2690	0.2734	0.2738
168798	MiniBooNE	0.0561	0.0583	0.0620
189873	dionis	0.1068	0.1068	0.6731
189866	albert	0.3168	0.3168	0.3143
75127	airlines	0.3394	0.3410	0.3449
75193	covertype	0.0519	0.0519	0.8571

Table 15: Results from Table 4 for 60 minutes. Numbers are the balanced error rate. We boldface the lowest error.

detection time (penalty<sub>i</sub>(t) = t). The detection time of a sensor placement T(A, i) is simply defined as the minimum of the detection times of its individual members:  $\min_{s \in A} T(s, i)$ .

In our setting, we need to do the following replacements to find that the problems are equivalent:

- 1. Intrusion scenarios  $\mathcal{I}$ : datasets  $\{\mathcal{D}_1, \dots, \mathcal{D}_{|\mathbf{D}_{meta}|}\}$ ,
- 2. Possible sensor locations C: set of candidate ML pipelines of our algorithm C, Detection time  $T(s \in A, i)$  on intrusion scenario i: test performance  $\mathcal{L}(\mathcal{M}_C, \mathcal{D}_{d, \text{test}})$  on dataset  $\mathcal{D}_d$ ,
- 3. Detection time of a sensor placement  $T(\mathcal{A}, i)$ : test loss of applying portfolio  $\mathcal{P}$  on dataset  $\mathcal{D}_d$ :  $\min_{p \in \mathcal{P}} \mathcal{L}(p, \mathcal{D}_{d, \text{test}})$
- 4. Penalty function penalty<sub>i</sub>(t): loss function  $\mathcal{L}$ , in our case, the penalty is equal to the loss.
- 5. Penalty reduction for an intrusion scenario  $R(\mathcal{A}, i)$ : the penalty reduction for successfully applying a portfolio  $\mathcal{P}$  to dataset d:  $R(\mathcal{P}, d) = \text{penalty}_d(\infty) \min_{p \in \mathcal{P}} \mathcal{L}(p, \mathcal{D}_{d, \text{test}})$ . 10

# C.3 Choosing on the validation set

We demonstrate that choosing an ML pipeline from the portfolio via holdout (i.e. a validation set) and reporting its test performance is neither submodular nor monotone by a simple example. To simplify notation we argue in terms of performance instead of penalty reduction, which is equivalent.

Let  $\mathcal{B} = \{(5,5), (7,7), (10,10)\}$  and  $\mathcal{A} = \{(5,5), (7,7)\}$ , where each tuple represents the validation and test performance. For e = (8,6) we obtain the discrete derivatives  $\Delta_f(e|\mathcal{A}) = -1$  and  $\Delta_f(e|B) = 0$  which violates Definition 2. The fact that the discrete derivative is negative violates Definition 3 because  $f(\mathcal{A}) > f(\mathcal{A} \cup \{e\})$ .

#### C.4 Successive Halving

As in the previous subsection, we use a simple example to demonstrate that selecting an algorithm via the successive halving model selection strategy is neither submodular nor monotone. To simplify notation we argue in terms of performance instead of penalty reduction, which is equivalent.

Let  $\mathcal{B} = \{((5,5),(8,8)),((5,5),(6,6)),((4,4),(5,5))\}$  and  $\mathcal{A} = \{((5,5),(7,7))\}$ , where each tuple is a learning curve of validation-, test performance tuples. For e = ((6,5),(6,5)), we eliminate entries 2 and 3 from  $\mathcal{B}$  in the first iteration of successive halving (while we advance entries 1 and 4), and we eliminate entry 1 from  $\mathcal{A}$ . After the second stage, the performances are  $f(\mathcal{B}) = 8$  and  $f(\mathcal{A}) = 5$ , and the discrete derivatives  $\Delta_f(e|\mathcal{A}) = -1$  and  $\Delta_f(e|\mathcal{B}) = 0$  which violates Definition 2. The fact that the discrete derivative is negative violates Definition 3 because  $f(\mathcal{A}) > f(\mathcal{A} \cup \{e\})$ .

<sup>10.</sup> This would be the general case for a metric with no upper bound. In case of metrics such as the misclassification error, the maximal penalty would be 1.

### C.5 Further equalities

In addition, our problem can also be phrased as a *facility location* problem (Krarup and Pruzan, 1983) and statements about the facility location problem can be applied to our problem setup as well.

# Appendix D. Implementation Details

### D.1 Software

We implemented the AutoML systems and experiments in the Python3 programming language, using numpy (Harris et al., 2020), scipy (Virtanen et al., 2020), scikit-learn (Pedregosa et al., 2011), pandas (McKinney, 2010), and matplotlib (Hunter, 2007).

### **D.2** Configuration Space

We give the configuration space we use in Auto-sklearn 2.0 in Table 16.

## D.3 Successive Halving hyperparameters

We used the same hyperparameters for all experiments. First, we set to eta = 4. Next, we had to choose the minimal and maximal budgets assigned to each algorithm. For the tree-based methods we chose to go from 32 to 512, while for the linear models (SGD and passive aggressive) we chose 64 as the minimal budget and 1024 as the maximal budget. Further tuning these hyperparameters would be an interesting, but an expensive way forward.

# Appendix E. Datasets

We give the name, OpenML dataset ID, OpenML task ID and the size of all datasets we used in Table 17 and 18.

Name	Domain	Default	Log
Classifier	(Extra Trees, Gradient Boosting, MLP,	Random Forest	-
	Passive Aggressive, Random Forest, SGD)		
Extra Trees: Bootstrap	(True, False)	False	-
Extra Trees: Criterion	(gini, entropy)	gini	-
Extra Trees: Max Features	[0.0, 1.0]	0.5	No
Extra Trees: Min Samples Leaf	[1, 20]	1	No
Extra Trees: Min Samples Split	[2, 20]	2	No
Gradient Boosting: Early Stopping	(off, valid, train)	off	-
Gradient Boosting: L2 Regularization	[1e-10, 1.0]	0.0	Yes
Gradient Boosting: Learning Rate	[0.01, 1.0]	0.1	Yes
Gradient Boosting: Max Leaf Nodes	[3, 2047]	31	Yes
Gradient Boosting: Min Samples Leaf	[1, 200]	20	Yes
Gradient Boosting: N Iter No Change	[1, 20]	10	No
Gradient Boosting: Validation Fraction	[0.01, 0.4]	0.1	No
MLP: Activation	(tanh, relu)	relu	_
MLP: Alpha	[1e - 07, 0.1]	0.0001	Yes
MLP: Early Stopping	(valid, train)	valid	_
MLP: Hidden Layer Depth	[1, 3]	1	No
MLP: Learning Rate Init	[0.0001, 0.5]	0.001	Yes
MLP: Num Nodes Per Layer	[16, 264]	32	Yes
Passive Aggressive: C	[1e - 05, 10.0]	1.0	Yes
Passive Aggressive: Average	(False, True)	False	-
Passive Aggressive: Loss	(hinge, squared_hinge)	hinge	
Passive Aggressive: Tol	[ $1e - 05, 0.1$ ]	0.0001	Yes
Random Forest: Bootstrap	(True, False)	True	168
Random Forest: Criterion	(gini, entropy)	gini	_
Random Forest: Max Features	[0.0, 1.0]	0.5	No
Random Forest: Min Samples Leaf	[0.0, 1.0] $[1, 20]$	0.5	No
1	[1, 20] $[2, 20]$	2	No
Random Forest: Min Samples Split Sgd: Alpha	[2, 20] $[1e - 07, 0.1]$	0.0001	Yes
Sgd: Aipha Sgd: Average	[1e - 07, 0.1] (False, True)	False	ies
9 9	, , ,	0.0001	Yes
Sgd: Epsilon Sgd: Eta0	$egin{array}{l} [1e-05,0.1] \ [1e-07,0.1] \end{array}$	0.0001	Yes
g	. , ,		Yes
Sgd: L1 Ratio	[1e - 09, 1.0]	0.15	res
Sgd: Learning Rate	(optimal, invscaling, constant)	invscaling	-
Sgd: Loss	(hinge, log, modified Huber,	log	-
C. I. D. Iv	squared hinge, perceptron)	10	
Sgd: Penalty	(l1, l2, elasticnet)	12	- NT
Sgd: Power T	[1e - 05, 1.0]	0.5	No
Sgd: Tol	[1e - 05, 0.1]	0.0001	Yes
Balancing: Strategy	(none, weighting)	none	_
Categorical Encoding: Choice	(no encoding, one hot encoding)	one hot encoding	-
Category Coalescence: Choice	(minority coalescer, no coalescense)	minority coalescer	-
Category Coalescence: Minimum Fraction	[0.0001, 0.5]	0.01	Yes
Imputation of missing values	(mean, median, most frequent)	mean	_
Rescaling: Choice	(Min/Max, none, normalize, Power,	standardize	_
5	Quantile, Robust, standardize)		
Quantile Transformer: N Quantiles	[10, 2000]	1000	No
Quantile Transformer: Output Distribution	(uniform, normal)	uniform	_
Robust Scaler: Q Max	[0.7, 0.999]	0.75	No
Robust Scaler: Q Min	[0.001, 0.3]	0.25	No

Table 16: Configuration space for Auto-sklearn 2.0 using only iterative models and only preprocessing to transform data into a format that can be usefully employed by the different classification algorithms. The final column (log) states whether we actually search  $log_{10}(\lambda)$ .

name	tid	#obs	#feat	#cls	name	id # ob	s # feat	# class	name	tid	#obs	#feat	#cls
OVA_O	75126	1545	10937	2	satim 21	20 643	37	6	fri_c	166950	500	11	2
OVA_C	75125		10937	2	Satel 1898				page	260	5473	11	5
OVA_P	75121	1545	10937	2	soybe 2	71 68	3 36	19	ilpd	146593	583	11	2
$OVA\_E$	75120	1545	10937	2	cardi 752	17 212	36	10	2dpla	75142	40768	11	2
OVA_K	75116	1545	10937	2	cjs 1466	01 - 279	35		fried	75161	40768	11	2
$OVA_L$	75115		10937	2	colle 752				rmfts	166859	508	11	2
OVA_B	75114		10937	2	puma3 751				stock	166915	950	10	2
UMIST			10305	20	Gestu 751				tic-t	279	958	10	2
amazo	189878		10001	50	kick 1898				breas	245	699	10	2
eatin	189786	945 60000	6374	7 10	bank3 751 wdbc 1465				xd6	167096 253	$973 \\ 1473$	10 10	2 3
GTSRB			3073						cmc		672	10	2
Biore	75156	51839 $3751$	$\frac{2917}{1777}$	43 2	Phish 752 fars 1898				profb diabe	146578 $267$	768	9	2
hiva	166996	4229	1618	2	hypot 30				abalo	2121	4177	9	28
GTSRB		51839	1569	43	steel 1687				bank8	75141	8192	9	2
GTSRB		51839	1569	43	eye_m 1897				elect	336	45312	9	2
Inter	168791	3279	1559	2	fri_c 751				kdd_e	166913	782	9	2
micro	146597	571	1301	20	fri_c 751				house	75176	20640	9	2
Devna	167203	92000	1025	46	wall 752	35 545	3 25	4	nurse	256	12960	9	5
GAMET	. 167085	1600	1001	2	led24 1898		) 25	10	kin8n	75166	8192	9	2
Kuzus	190154	270912	785	49	colli 1898	45 100	24	30	yeast	2119	1484	9	10
mnist	75098	70000	785	10	rl 1898	69 3140			puma8	75171	8192	9	2
Kuzus	190159	70000	785	10		54 812			analc	75143	4052	8	2
isole	75169	7797	618	26	meta 1668				ldpa		164860	8	11
har	126030	10299	562	6	jm1 750				pm10	166872	500	8	2
madel	146594	2600	501	2	pc1 751				no2	166932	500	8	2
KDD98		82318	478	2	kc2 1465				LED-d		500	8	10
phili	189864	5832	309	2	cpu_a 752				artif	126028	10218	8	10
madel	189863	3140	260	2	autoU 750				monks	3055	554	7	2
USPS	189858	9298	257	10	GAMET1670				space	75148	3107	7	2
semei	75236	1593	257	10	GAMET 1670				kr-vs	75223	28056	7	18
GTSRB India	211720	51839 9144	$\frac{257}{221}$	43 8	bosto 1669 GAMET1670				monks Run_o	3054 $167103$	601 88588	7 7	$\frac{2}{2}$
dna	167202	3186	181	3	GAMET 1670				delta	75173	9517	7	2
musk	75108	6598	170	2	churn 1670				strik	166882	625	7	2
Speed	146679	8378	123	2	clima 1671				mammo		11183	7	2
hill	146592	1212	101	2	micro 1898				monks	3053	556	7	2
fri_c	166866	500	101	2	GAMET 1670				kropt	2122	28056	7	18
MiceP	167205	1080	82	8	Traff 2117	24 7034	21	. 3	delta	75163	7129	6	2
meta	2356	45164	75	11	ringn 752				wilt	167105	4839	6	2
ozone	75225	2534	73	2	twono 751				fri_c	75131	1000	6	2
analc	146576	841	71	4	eucal 21				mozil	126024	15545	6	2
kdd_i	166970	10108	69	2	eleva 751				polle	75192	3848	6	2
optdi	258	5620	65	10	pbcse 1668				socmo	75213	1156	6	2
one-h	75154	1600	65	100	baseb 21				irish	146575	500	6	2
synth	146574	600	62	6	house 751				fri_c	166931	500	6	2
splic spamb	$\frac{275}{273}$	$\frac{3190}{4601}$	61 58	3 2	colle 751 BachC 1898				arsen	166957 $166956$	559 559	5 5	$\frac{2}{2}$
first	75221	6118	52	6		62 1099			walki	75250	149332	5	22
fri_c	75180	1000	51	2		36 2000			analc	146577	797	5	6
fri_c	166944	500	51	2		78 26325			bankn	146586	1372	5	2
fri_c	166951	500	51	2	eeg-e 752				arsen	166959	559	5	2
Diabe	189828	101766	50	3	wind 751		4 15	2	visua	75210	8641	5	2
oil_s	3049	937	50	2	Japan 1260	21 996	1 15	9	balan	241	625	5	3
pol	75139	15000	49	2	compa 2117	21 - 527	8 14	. 2	arsen	166958	559	5	2
tokyo	167100	959	45	2	vowel 30	47 99	) 13	11	volca	189902	10130	4	5
qsar	75232	1055	42	2	cpu_s 751				skin		245057	4	2
textu	126031	5500	41	11	autoU 1899	00 70	) 13	3	tamil	189846	45781	4	20
autoU	189899	750	41	8	autoU 751				quake	75157	2178	4	2
ailer	75146	13750	41	2	dress 1466				volca	189893	8654	4	5
wavef	288	5000	41	3	senso 1669				volca	189890	8753	4	5
cylin	146600	540	40	2	wine 1898				volca	189887	9989	4	5
water	166953	527	39	2	wine 1898				volca	189884	10668	4	5
annea	232	898	39	5	Magic 751				volca	189883	10176	4	5
mc1	75133 75092	$9466 \\ 1458$	39 38	2 2	mv 751 parit 1671				volca volca	189882 189881	$1515 \\ 1521$	$\frac{4}{4}$	5 5
pc4 pc3	75129	1563	38 38	2	parit 1671 mofn 1670				voica	189881	1623	4	5
pcs porto		595212	38	2	fri_c 751				Titan	167099	2201	4	2
pc2	75100	5589	37	2		40 82920			volca	189894	1183	4	5
r	. 5100	2000			F	, , , , , ,					-100		

Table 17: Characteristics of the 208 datasets in  $\mathbf{D}_{\text{meta}}$  (first part) sorted by number of features. We report for each dataset the name, the dataset id (as a link) and the task id as used on OpenML.org, and furthermore the number of observations, the number of features and the number of classes.

name	$\operatorname{tid}$	# obs	$\# {\rm feat}$	$\#\mathrm{cls}$	name	$\operatorname{tid}$	# obs	$\# {\rm feat}$	$\#\mathrm{cls}$
rober	168794	10000	7201	10	kr-vs	167149	3196	37	2
ricca	168797	20000	4297	2	higgs	167200	98050	29	2
guill	168796	20000	4297	2	helen	189874	65196	28	100
dilbe	189871	10000	2001	5	kc1	167181	2109	22	2
chris	189861	5418	1637	2	numer	167083	96320	22	2
cnae	167185	1080	857	9	credi	167161	1000	21	2
faber	189872	8237	801	7	sylvi	189865	5124	21	2
Fashi	189908	70000	785	10	segme	189906	2310	20	7
KDDCu	75105	50000	231	2	vehic	167168	846	19	4
mfeat	167152	2000	217	10	bank	126029	45211	17	2
volke	168793	58310	181	10	Austr	167104	690	15	2
APSFa	189860	76000	171	2	adult	126025	48842	15	2
jasmi	189862	2984	145	2	Amazo	75097	32769	10	2
nomao	126026	34465	119	2	shutt	168795	58000	10	7
alber	189866	425240	79	2	airli	75127	539383	8	2
dioni	189873	416188	61	355	car	189905	1728	7	4
janni	168792	83733	55	4	jungl	189909	44819	7	3
cover	75193	581012	55	7	phone	167190	5404	6	2
MiniB	168798	130064	51	2	blood	167184	748	5	2
conne	167201	67557	43	3					

Table 18: Characteristics of the 39 datasets in  $\mathbf{D}_{\text{test}}$  sorted by number of features. We report for each dataset the name, the dataset id (as a link) and the task id as used on OpenML.org, and furthermore the number of observations, the number of features and the number of classes.